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Ribozymes and Riboswitches: Modulation of RNA Function by Small Molecules[†]

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Abstract

Diverse small molecules interact with catalytic RNAs (ribozymes) as substrates and cofactors, and their intracellular concentrations are sensed by gene-regulatory mRNA domains (riboswitches) that modulate transcription, splicing, translation, or RNA stability. Although recognition mechanisms vary from RNA to RNA, structural analyses reveal recurring strategies that arise from the intrinsic properties of RNA such as base pairing and stacking with conjugated heterocycles, and cation-dependent recognition of anionic functional groups. These studies also suggest that, to a first approximation, the magnitude of ligand-induced reorganization of an RNA is inversely proportional to the complexity of the riboswitch or ribozyme. How these small molecule binding-induced changes in RNA lead to alteration in gene expression is less well understood. While different riboswitches have been proposed to be under either kinetic or thermodynamic control, the biochemical and structural mechanisms that give rise to regulatory consequences downstream of small molecule recognition by RNAs mostly remain to be elucidated.

Ribozymes and riboswitches starkly demonstrate the ability of RNA to fold into complex structures that position functional groups with exquisite precision. The former are catalytic RNAs; the latter, *cis*-acting regulatory mRNA domains that respond to the intracellular concentration of small molecule metabolites and second messengers [the first example of a *trans*-acting riboswitch RNA was recently described (1)]. *In vitro*, both ribozymes and riboswitches can function in the absence of protein cofactors, although some catalytic RNAs are known to require chaperones [reviewed in (2)] for *in vivo* activity, and riboswitches ultimately need to interface with the rest of the gene expression (transcription, splicing, translation, or RNA degradation) machinery for their small molecule-dependent regulatory activity to become manifest. Over the past decade, structural analyses have shed light on the mechanism of small molecule recognition by ribozymes (as substrates and coenzymes) and riboswitches (as regulatory signals). We review the state of knowledge of small molecule recognition by RNA, and how small molecule binding gives rise to genetic regulation.

1. Ligand recognition by natural and artificial aptamers

Structures of riboswitch aptamer domains

At the time of writing, structures of the ligand-binding [or "aptamer" (3)] domains of representatives of eleven different classes of naturally-occurring riboswitches have been

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reported: cyclic diguanylate (c-di-GMP) (4,5), flavin mononucleotide (FMN) (6), *glmS* (7,8), lysine (9,10), preQ₁ (11–13), magnesium ion (14), purine (guanine and adenine) (15,16), S-adenosylmethionine Class I (SAM-I) (17), SAM-II (18), SAM-III (19), and thiamine pyrophosphate (TPP) (20–22). The aptamer domains of these riboswitches all adopt distinctly different structures. There are no discernible evolutionary relationships between different classes of riboswitches, making it uncertain whether each class descends from an independent evolutionary ancestor, or if sequence and structure divergence have proceeded too far for lineage relationships to be apparent.

Riboswitch aptamer domains can be classified on the basis of structural features. For instance, and analogous to ribozyme classification (23,24), riboswitch aptamers can be divided into those that are organized around multi-helical junctions, and those whose architectures result from pseudoknot (25) formation. The c-di-GMP, purine, SAM-III and TPP aptamer domains are all 3-helix junctions, and except for SAM-III, form Y-shaped structures (26) stabilized by distal interactions between two of the helices (Figure 1A). The Mg²⁺ riboswitch contains two 3-helix junctions. The lysine riboswitch structure is organized around a 5-helix junction, and that of the FMN riboswitch, around a 6-helix junction. The latter is notable for a series of loop-loop interactions that result in a pseudo-symmetric molecule that recognizes its ligand asymmetrically. The SAM-II and preQ₁ riboswitch aptamer domains are simple H-type pseudoknots (Figure 1B). The SAM-I riboswitch aptamer domain is organized around a 4-way helical junction, but its folded structure also contains a pseudoknot (Figure 1C). Finally, the *glmS* riboswitch-ribozyme is comprised of three pseudoknots. Its minimal core, sufficient *in vitro* for ligand recognition and catalysis, comprises a nested double pseudoknot. A peripheral domain folds as an H-type pseudoknot that itself functions as a 3-helix junction (Figure 1D).

Ligand recognition strategies of riboswitches

Given the diversity of riboswitch aptamer domain architectures, it is not surprising that the specific mechanisms employed by these RNAs to recognize their cognate ligands vary greatly. Nonetheless, the characteristic chemical properties of RNA give rise to several recurrent molecular recognition strategies. Recognition of TPP by its cognate riboswitch illustrates three of these: base stacking, base pairing, and metal ion-mediated binding (Figure 2). TPP is bound in an extended conformation and spans two of the three helices of the aptamer domain. The pyrimidine ring of TPP is recognized through stacking and formation of a base pair-like interaction (27) with the sugar edge of G40. The central thiazole ring of TPP makes only van der Waals interactions with the RNA. The pyrophosphate of TPP is essential for recognition by the riboswitch. The RNA does not recognize the pyrophosphate directly; rather, it binds it as a chelate with two partially hydrated divalent cations. Indeed, most of the interactions between RNA and pyrophosphate atoms are mediated by the two divalent cations or by their hydration sphere (20,21) (Figure 2). Crystal structures of the riboswitch bound to thiamine monophosphate (TMP), against which it discriminates at least 30-fold (28), revealed that the riboswitch binds to the shorter ligand using essentially the same interactions. This is made possible by a slight compaction of the riboswitch, which brings the two helices that recognize the opposite ends of the ligand closer, and by coordination of the single phosphate of TMP with two divalent cations that in turn make interactions with the RNA similar to those made by the two cations chelated by TPP (20).

Stacking and base pairing are predictably common in riboswitches that recognize nucleobases or other planar heterocycles (27). The purine, c-di-GMP and preQ₁ riboswitches all recognize guanine or guanine-containing ligands using these strategies, yet the specific nucleobase interactions employed are different in each case (4,5,11,16) (Figure 3). In all cases, stacking of the ligand between nucleobases is important, but it has been noted that the

stacking interactions employed by the purine riboswitch are sub-optimal. This may have the consequence of increasing the contribution to specificity of base pairing (29). Indeed, substitution of C74 (Figure 3A) to U is sufficient to switch the ligand specificity of the riboswitch from G to A (30). Consistent with their architectural diversity, the three structurally characterized classes of SAM riboswitches employ different base-pairing schemes to recognize the adenine moiety of their ligand. Their strategies for binding the other elements of SAM are also distinctly different (18,19). The FMN riboswitch recognizes the edge of the isoalloxazine ring of FMN that resembles a uracil through formation of a base pair-like interaction with an adenosine residue of the RNA (6).

Reminiscent of the TPP riboswitch, the FMN riboswitch employs a tightly bound metal ion to recognize the phosphate moiety of FMN. Like the TPP riboswitch, the FMN riboswitch can employ a variety of divalent cations *in vitro* to mediate recognition of the phosphate of its ligand (6). The phosphate groups of c-di-GMP and glucosamine-6-phosphate (GlcN6P) are also recognized by their respective cognate riboswitch as divalent metal ion chelates (5,7). Metal ion-mediated ligand recognition by riboswitches is not limited to phosphate-containing ligands. The lysine riboswitch has a strict dependence on K^+ for ligand recognition. The structure shows that K^+ coordinates the carboxylate of the bound lysine, and the cation in turn makes inner-sphere coordination with RNA functional groups (9). Lysine binding is 50–100 fold weaker if K^+ is replaced with Na^+ or Mg^{2+} (9). The preQ₁ riboswitch discriminates ~25-fold against guanine, from which its cognate ligand differs by the exocyclic aminomethyl group (and the deaza substitution at position 7) (Figures 3A, 3B) (31). The cocrystal structure shows that the exocyclic amine participates in tetrahedral hydrogen bonding with three ligands: a non-bridging phosphate oxygen, the carbonyl oxygen of G5, and a hydration water of a metal ion (11).

Comparison with artificial aptamers

Structural studies of many *in vitro* selected aptamer RNAs [reviewed in (32)] predate the discovery of small molecule-responsive natural riboswitches. These studies demonstrated the presence of structural motifs previously identified in natural RNAs in the artificial aptamers. For instance, structures of the ATP aptamer in complex with AMP (the aptamer does not recognize the phosphate groups) revealed that the adenosine binds to an internal loop of the aptamer that adopts the fold of the GNRA tetraloop (33), taking the place of the last A of the motif and base pairing with the first G of the motif in the manner of the natural loop (34). Several commonalities were suggested by analyses of a number of artificial aptamers including the use of stacking between base pairs and triples to recognize planar ligands and the coupling of aptamer folding with ligand binding (32). It was proposed that the latter is a defining characteristic of artificial aptamers, which evolved to satisfy a single function, ligand binding, unlike natural RNAs, which evolved as part of biological networks that require structural optimization. Comparison with structures of the aptamer domains of riboswitches confirms the wide-spread use of stacking and base pairing interactions in ligand recognition [compare, for instance, the structures of the FMN aptamer (35) and the FMN riboswitch (6)], and demonstrates a comparable degree of binding site-ligand shape complementarity in natural and artificial RNAs (36), but refutes the notion that ligand binding-induced folding (global induced fit) is a defining characteristic of *in vitro* evolved aptamers.

To a first approximation, it appears that the degree of ligand binding-induced folding of RNAs correlates inversely with their complexity. The preQ₁ riboswitch aptamer domain, the smallest known from nature (34 nt) folds only in the presence of ligand (37). *In vitro* selection experiments for high-affinity GTP aptamers resulted in a number of isolates that are more informationally complex than the lower-affinity ATP aptamer of similar global fold, suggesting that a structure capable of 10-fold tighter binding requires specification of

five additional nucleotides (38). Biochemical and structural analyses, however, demonstrate that these additional nucleotides do not make direct RNA-ligand interactions (39,40). Rather, they engage in RNA-RNA interactions that pre-organize the RNA, reducing its conformational heterogeneity and increasing its propensity (41) to fold. Thus, most of the difference in affinity between a low affinity ATP aptamer and a high affinity GTP aptamer can be accounted for by a 45-fold faster k_{on} (40). The artificial RNA with the highest known affinity for its small molecule ligand ($K_d \sim 0.8$ nM) is the 70 nt tetracycline aptamer (42) (Table I). Biochemical and structural analysis of this Y-shaped RNA indicates that, unlike simple aptamers that are unfolded in the absence of ligand, this molecule is partially preorganized (42–44). Tetracycline is bound at a pocket formed between a 11-nt loop and the minor groove of an irregular duplex, and while the former appears to fold concomitantly with binding, the latter is structured in the absence of ligand (44). Such behavior is reminiscent of that of natural riboswitch aptamer domains of similar complexity. For instance, crystallographic (20), spectroscopic (45), calorimetric (28) and SAXS (46) analyses of the TPP riboswitch aptamer domain (83 nt) indicate that ligand binding induces local as well as global conformational rearrangement, and that divalent cations facilitate ligand binding by pre-organizing the RNA, in addition to forming part of the TPP-RNA interface. The *glmS* and lysine riboswitches, which have two of the largest riboswitch aptamer domains so far characterized (~150 and ~180 nt, respectively) have global conformations that are insensitive to ligand occupancy (7,9,10,47–49).

2. Small molecule recognition by natural and artificial ribozymes

Diversity of ribozyme catalyzed reactions

Thus far, natural ribozymes have been discovered that catalyze peptidyl transfer (the ribosome) and phosphoryl transfer reactions [reviewed in (50)]. The latter includes sequence-specific cleavage and ligation catalyzed by ribozymes through two types of reactions. The first is an internal transesterification involving the nucleophilic attack of a 2' oxygen of a ribose on the adjacent 3' phosphate group, resulting in two cleavage products, one containing a 2',3'-cyclic phosphate and the other a 5'-OH. Five classes of natural RNAs, known as the 'small self-cleaving ribozymes', are known to catalyze this reaction: the hammerhead, hairpin, hepatitis delta virus (HDV), Varkud satellite (VS), and *glmS* ribozymes [reviewed in (24)]. The second type of reaction is catalyzed by RNase P and the group I and II self-splicing introns (which are responsible for both phosphodiester cleavage and ligation), and related RNAs. These ribozymes yield either ligated exons or products with 5'-phosphate and 3'-OH termini, and can employ internal or external nucleophiles (50).

While the ability of RNAs to function as biochemical catalysts was first discovered in natural ribozymes, it has been through *in vitro* selections from randomized libraries by which most RNAs catalyzing different chemical transformations have been discovered. Artificial ribozymes include aminoacyl-RNA synthetases, RNA ligases and kinases ribozymes, nucleotide synthetases, thioester synthetases, and Diels-Alderase [reviewed in (51)]. Although few of these *in vitro* evolved ribozymes achieve rate enhancements comparable to those of natural ribozymes (52), their discovery has greatly broadened the known spectrum of ribozyme catalysis.

Small molecule recognition by natural ribozymes

Structural information is available on small molecule binding by two classes of natural ribozymes: the *glmS* riboswitch and group I introns. The *glmS* riboswitch is also a catalytic RNA. Binding of GlcN6P activates a latent self-cleavage activity of the ribozyme, increasing the rate of cleavage of a specific phosphodiester bond by $\sim 10^6$. GlcN6P functions as a coenzyme, and is absolutely required for activity of the ribozyme. Structures of *glmS*

riboswitches from two different bacterial species have been determined, each in multiple functional states [reviewed in (53)]. These studies suggest that neither binding of GlcN6P, catalytic activation, or product release is accompanied by RNA conformational changes. Indeed, the ribozyme is competent for GlcN6P binding and catalysis in the crystalline state. Thus, like other natural ribozymes [*e.g.* (54)], once the *glmS* ribozyme folds, catalysis proceeds with motions localized to the active site. Binding of GlcN6P to the rigid *glmS* ribozyme is made possible by its open, solvent-accessible ligand binding site, which contrasts with the partially or totally solvent occluded binding sites of non-catalytic riboswitches. Analog studies show that the amine group of GlcN6P is essential for catalysis, and that the anomeric hydroxyl and phosphate groups of the coenzyme play important roles in binding (55,56). The structures show that GlcN6P stacks underneath the nucleobase of the guanosine residue immediately following the scissile phosphate, with the phosphate of GlcN6P hydrogen bonding to N1 of the guanine. Outer-sphere coordinated divalent cations provide non-specific stabilization of GlcN6P binding. The amine of GlcN6P hydrogen bonds to the 5'-oxygen (the leaving group of the transesterification reaction), and the anomeric hydroxyl hydrogen bonds to the scissile phosphate. Experimental data are consistent with the amine of GlcN6P functioning as a general acid and electrostatic catalyst (57).

Group I introns employ the 3'-OH of an exogenous guanosine (exoG) as the nucleophile in the first transesterification reaction, resulting in release of the 5' exon and addition of exoG to the 5' terminus of the intron. This is followed by local rearrangements of the ribozyme and replacement of exoG in the binding site with the conserved guanosine residue at the 3' terminus of the intron (ω G), which provides the nucleophile for the second step of splicing. Structures have been determined of group I introns from three species in multiple functional states. Although none of these carry exoG, its mode of binding can be inferred from the manner in which ω G is bound. The guanosine forms part of a base triple, and this triple is in turn sandwiched by two additional triples (Figure 4A). Although the participation of divalent cations in formation of the active site (including bridging ribose moieties of the guanosine with the substrate portion of the RNA) and in the catalytic steps is well established, the precise location of these ions in the transition state is a subject of active research. Different lines of evidence indicate that exoG binding, release, and replacement with ω G are accompanied by conformational rearrangements of the intron, but the structural details of these remain to be established because none of the currently available structures are determined at better than 3 Å resolution [reviewed in (58)].

Small molecule recognition by artificial ribozymes

Structural information is available on small molecule binding by two classes of artificial ribozymes: a Diels-Alderase and flexizyme. The Diels-Alder reaction is a [4+2] cycloaddition of great importance in organic chemistry. Natural protein enzymes are known that catalyze this reaction, and Diels-Alderase catalytic antibodies have been selected by using transition state-mimics as haptens. Several *in vitro* selection experiments have been carried out to isolate Diels-Alderase ribozymes [reviewed in (59)]. The structure of a 49-nt Diels-Alderase selected to catalyze the cycloaddition of a biotinylated maleimide and a tethered anthracene derivative (60) was determined both free and bound to the reaction product (61). The RNA folds into a compact double pseudoknot that cradles the cycloaddition product in a solvent-exposed groove, interacting with it primarily through aromatic-aromatic stacking and non-polar van der Waals contacts, using very limited hydrogen bonding. Shape complementarity of the ribozyme to the product is good, but calculations suggest that it is even better to the transition state. Thus, catalysis appears to result from proximity effects (binding and positioning of the reactants) and preferential

transition state binding. The rigidity of the folded ribozyme ensures a high degree of regioselectivity [see also (62)].

Flexizyme catalyzes the regioselective 3'-aminoacylation of tRNAs using pre-activated (*e.g.* adenylated) amino acids (63). The 45-nt flexizyme recognizes its tRNA substrate by base pairing to its NCCA 3'-terminus (64), and positions the acceptor helix approximately perpendicular to an irregular, underwound active site helix. The minor groove of this helix buckles open to expose the planar faces of two nucleotides to provide a binding site for the amino acid side chain. The nature of this pocket is consistent with flexizyme's strong preference for aromatic amino acids. Indeed, cocrystallization of flexizyme with phenylalanine-ethylester (an inhibitor) suggests that the benzyl moiety stacks on G24 so as to maximize polar-aromatic interactions (65). This positions the carbonyl of the amino acid in van der Waals contact with the nucleophile 3'-OH of the tRNA (Figure 4B). Structural and biochemical analyses suggest that flexizyme can achieve considerable specificity despite this simple, open active site architecture by coupling amino acid binding to productive positioning of the acceptor terminus, that is, employing an induced-fit mechanism.

Small molecule recognition by natural and artificial ribozymes

Although the basic small-molecule recognition strategies employed by ribozymes (such as stacking, base pairing and cation-mediated binding) are the same as those of riboswitches, ribozyme active sites appear to be qualitatively simpler than riboswitch ligand binding pockets. This in part must reflect the fact that crystal structures are of ground-state complexes, while ribozyme active sites have evolved to be complementary to the transition state (66). The *glmS* riboswitch and the Diels-Alderase appear to be rigid structures that permit catalysis to proceed with minimal additional loss of entropy once the substrates (cofactor, for *glmS*) are bound. On the other hand, group I intron active sites necessarily must remain flexible so that the same active site can catalyze both steps of splicing. It appears that flexizyme has evolved to exploit its flexibility to couple amino acid binding site structure formation to productive tRNA docking, and in this way overcomes the limitations of its rather featureless amino acid binding pocket.

3. Two paths to the same outcome: kinetic and thermodynamic control of riboswitches

Diverse mechanisms of genetic control

How do the conformational and energetic changes undergone by natural or artificial RNAs upon ligand binding result in alteration of gene expression? The tetracycline aptamer has been employed as an artificial riboswitch to modulate alternative pre-mRNA splicing (67) or translation initiation (68). In Gram-positive bacteria the GlcN6P-dependent self-cleaving activity of the *glmS* ribozyme controls the stability of the mRNA it is part of (69). The only riboswitch class thus far discovered in eukaryotes is the TPP riboswitch. In plants, fungi and algae, these riboswitches control alternative splicing. In all characterized cases, segments of the aptamer domain of the ligand-free TPP riboswitch (such as the pyrophosphate binding nucleotides) can base pair with nearby splice sites, occluding them. Upon TPP binding, the riboswitch aptamer domain folds, and exposes the previously occluded splice site, leading to changes in the pre-mRNA splicing pattern [reviewed in (70)]. In bacteria, most riboswitches appear to function by transcription attenuation (termination *versus* readthrough) or translation initiation control. For these mechanisms, sequence elements in addition to the riboswitch aptamer domain are needed. These have been termed the 'expression platform' (3). Transcription attenuation, particularly effective in Gram-positive bacteria due to increased operon size and complexity, primarily makes use of an overlapping RNA segment,

which chooses between mutually exclusive RNA structures leading to either premature transcription termination or readthrough.

Transduction of ligand binding into a genetic decision can occur under kinetic or thermodynamic control. The relevant parameters can include the rate of aptamer folding, ligand concentration, k_{on} , k_{off} , and K_d for aptamer, speed of transcription, delay time at transcriptional pauses, competition between alternative RNA structures, height of activation barrier between bistable RNA structures, etc. A kinetic control scheme precisely gears the kinetic parameters of transcription to those of ligand binding and co-transcriptional folding of aptamers, whereas a thermodynamic scheme depends on an equilibrium between ligand association and dissociation, or between bistable, inter-converting RNA structures whose relative populations are shifted by the concentration of ligands.

A large discrepancy between the concentration of FMN required to elicit a half maximal shift in termination efficiency *in vitro* (T_{50}) and the dissociation constant prompted two studies of the kinetic basis of FMN and adenine riboswitch regulation (71,72). By monitoring the binding- induced quenching of FMN and 2-amino-purine fluorescence, k_{on} and k_{off} values were directly measured in these studies. [Interestingly, most riboswitch aptamers examined to date exhibit similar k_{on} values on the order of 10^4 to $10^5 \text{ M}^{-1}\text{s}^{-1}$, whereas k_{off} are found to range from 10^{-3} to 10^{-1} s^{-1} (Table 1).] These quantities, combined with commonly observed micromolar levels of metabolite concentrations, place the time constant τ ($\sim 1/k_{\text{off}}$) leading to association and dissociation equilibria in the range of seconds. This is comparable to that of the elongation of RNA polymerase (Δt_{RNAP}) from the end of the aptamer domain to the termination decision point. Δt_{RNAP} depends on the length of the expression platform, the elongation speed of the cognate RNAP, and importantly, the recognition efficiency and dwell times at any long-lived transcriptional pause sites (73).

The hallmark of kinetic control is $\Delta t_{\text{RNAP}} < \tau$, which means that RNAP arrives at the termination decision point before ligand and aptamer reach binding equilibrium (Figure 5, τ_2 , green dot). Since equilibrium is not reached, a ligand concentration equivalent to K_d does not lead to 50% ligand binding, producing significant discrepancies between T_{50} and K_d . In contrast, a thermodynamic mechanism would need to satisfy $\Delta t_{\text{RNAP}} > \tau$, which allows ligand and aptamer to reach equilibrium before RNAP reaches the transcription terminator (Figure 5, τ_1 , orange dot). Under these conditions, T_{50} would approximate K_d . Conceivably, the choice of kinetic or thermodynamic control may have evolved to match the range of K_d and the intracellular level of ligand in question. It is also possible for a riboswitch to operate under either kinetic or thermodynamic control depending on environmental cues such as nutritional availability that would alter intracellular NTP levels, and therefore elongation rates and pause durations.

Kinetic control: the FMN riboswitch

FMN binding to the B. subtilis ribD FMN riboswitch aptamer occurs with $k_{\text{on}} = 2.0 \times 10^5 \text{ M}^{-1}\text{s}^{-1}$, $k_{\text{off}} = 0.002 \text{ s}^{-1}$, $K_d = 12 \text{ nM}$, and $T_{50} = 200\text{--}500 \text{ nM}$. A T_{50} much higher than K_d suggests kinetic control in which substantially more FMN is required rapidly to occupy 50% of the aptamers before the RNAP reaches the termination decision point. As expected for a kinetically regulated riboswitch, the termination efficiency is sensitive to NTP concentrations. Two prominent transcriptional pauses were identified that when disrupted led to decreased termination and another 4-fold increase in T_{50} . This suggests that these functionally important, long-lived pauses provide necessary delays for FMN to bind to the aptamer (71).

Thermodynamic control: the preQ₁ riboswitch

Fluorescence and NMR spectroscopic studies of the *F. nucleatum* preQ₁ riboswitch revealed the co-existence of two competing stem-loop structures in the expression platform, corresponding to the antiterminator and terminator hairpins. The relative populations of these two structures at equilibrium are rapidly (relative to elongation rates of some bacterial RNAPs) shifted by the presence of preQ₁, allowing for thermodynamic control. However, unavailability of cognate *F. nucleatum* RNAP precluded accurate estimates of Δt_{RNAP} , and thus definitive determination of the control mechanism for this riboswitch (37). Nonetheless, the bistable nature of the terminator and antiterminator hairpin loops and their rapid exchange suggests a paradigm for riboswitches under thermodynamic control.

4. From simplified models to realistic regulatory scenarios

Nascent RNA structures impact transcription

Riboswitch and ribozyme function *in vivo* can be much more complex than in these defined *in vitro* systems, due to the rugged kinetic landscape of RNAP elongation through these structured RNAs, co-transcriptional folding of aptamers, modulation of RNA folding rates by cognate and non-cognate small molecules, coupling of translation with transcription, and the action of RNA binding proteins. During the course of transcription, nascent RNA structures such as pause hairpins exert significant effects on the kinetics of transcription (74,75). Some nascent RNAs can bind the RNAP and confer to it robust anti-pausing and anti-termination capabilities. Examples are the *PutL* RNA in the lambdoid phage HK22 anti-termination (76), and the *B. subtilis* EAR (*eps*-associated RNA) element, which was proposed to associate with RNAP and confer processivity necessary to complete an usually long (16kb) *eps* (exopolysaccharide) transcript important in biofilm formation (77). The effects of ligand binding (or *glmS* riboswitch self cleavage or intron self-splicing) on nascent riboswitch transcription (including RNAP pausing) remains to be elucidated.

Kinetic coupling and regulator trafficking between translation and transcription machines

In bacteria, trailing ribosomes typically travel slower than leading RNAP, and thus also utilize transcriptional pauses to ensure coupling of translation to transcription, preventing the termination factor Rho from loading onto naked RNA and causing undesired termination. Ribosomes not only drive paused RNAPs back to elongation by disrupting pause hairpins but also directly modulate the rates of transcription by preventing spontaneous RNAP backtracking and providing assistance through DNA roadblocks (78). How ligand binding to riboswitches affects this coupling is unknown. Moreover, little kinetic characterization of riboswitches that function through translation has been reported.

RNA binding proteins and chaperones may affect riboswitch function

In addition to being restricted by the leading RNAP and trailing ribosome, the riboswitch mRNA emerging from the RNAP exit channel is subject to binding and potential control by additional protein factors. NusA, docked on the RNAP exit channel during elongation, may hand over partially folded nascent RNA structures onto other protein factors such as ribosomal protein L7Ae homologs (*e.g.* YbxF and Ylq in *B. subtilis*), which recognize K-turn motifs present in many structured RNAs (*e.g.*, SAM-I, Lysine, and T-box riboswitches) (79–81). While riboswitches have been shown to function without proteins *in vitro*, RNA-binding proteins may play architectural roles by assisting folding of complex aptamers, regulatory roles by selectively stabilizing one of mutually exclusive RNA structures, introducing post-transcriptional RNA modifications (82), or even directly acting as the signaling ligand, such as the case of L4 associating with the S10 operon 5' leader and causing termination of its own operon (83).

Riboswitches and ribozymes provide a biological context where small molecule recognition by RNA, RNA folding assisted by small molecules, and transcription, translation, and splicing kinetics influenced by nascent RNA structures synergistically operate genetic switches. In light of the rapid pace of structure determination of natural and artificial riboswitches and ribozymes, the interface between the RNA-ligand modules and their expression platforms warrants more research. A mechanistic understanding of such coupling would allow modular design and rational engineering of artificial bacterial and mammalian riboswitches and ribozymes as well as modulation of their natural counterparts.

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Abbreviations

c-di-GMP	cyclic diguanylate
EAR	<i>eps</i> -associated RNA
exoG	exogenous guanosine
FMN	flavin mononucleotide
GlcN6P	glucosamine-6-phosphate
GTP	guanosine triphosphate
HDV	hepatitis delta virus
NTP	nucleotide triphosphate
ωG	the conserved guanosine at the 3' terminus of Group I introns
RNAP	RNA polymerase
SAM	S-adenosylmethionine
SAXS	small angle X-ray scattering
TMP	thiamine monophosphate
TPP	thiamine pyrophosphate
VS	Varkud satellite

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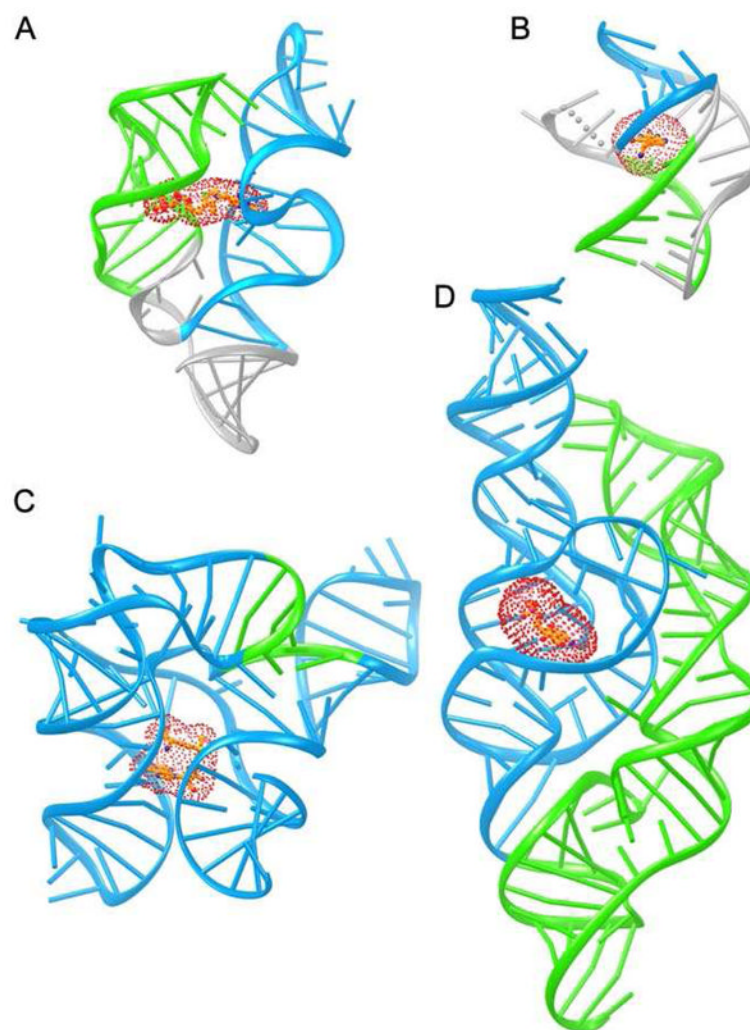
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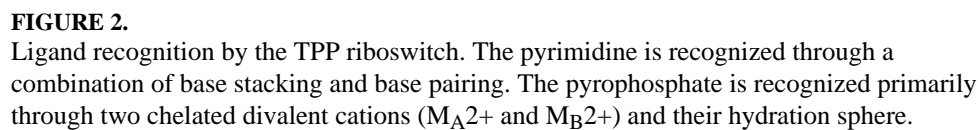
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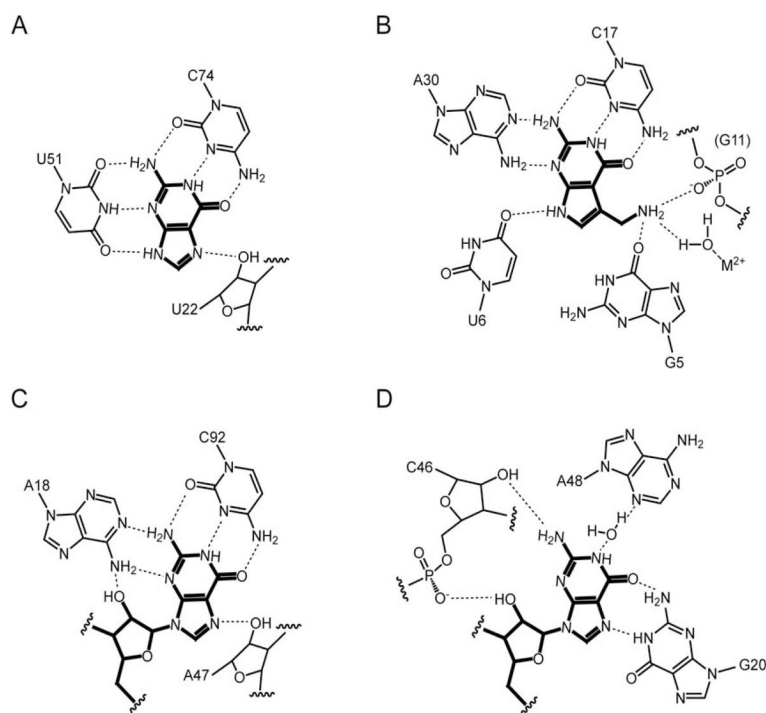
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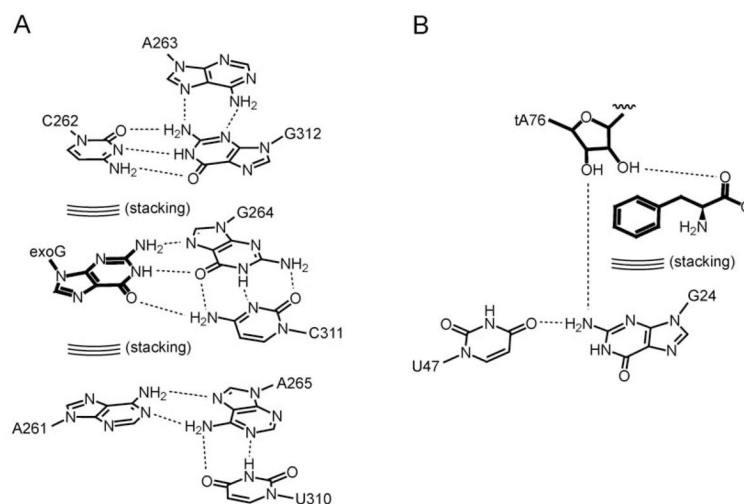
**FIGURE 1.**

Overall structure of representative riboswitch aptamer domains in cartoon form. (A) The TPP riboswitch (21) adopts a Y-shaped structure that recognizes its ligand (red) in an elongated conformation between the pyrimidine-binding helix (cyan) and the pyrophosphate binding helix (green). (B) The preQ₁ riboswitch (11) folds as an H-type pseudoknot, with its ligand continuing the stack between the two A-form helices (cyan and green). (C) The SAM-I riboswitch (17) is organized around a four helix junction (cyan) but also comprises a pseudoknot (green). (D) The *glmS* ribozyme-riboswitch is comprised of a double-pseudoknot core domain (cyan) and a peripheral domain (green) that consists of a pseudoknot that functions as a 3-helix junction.

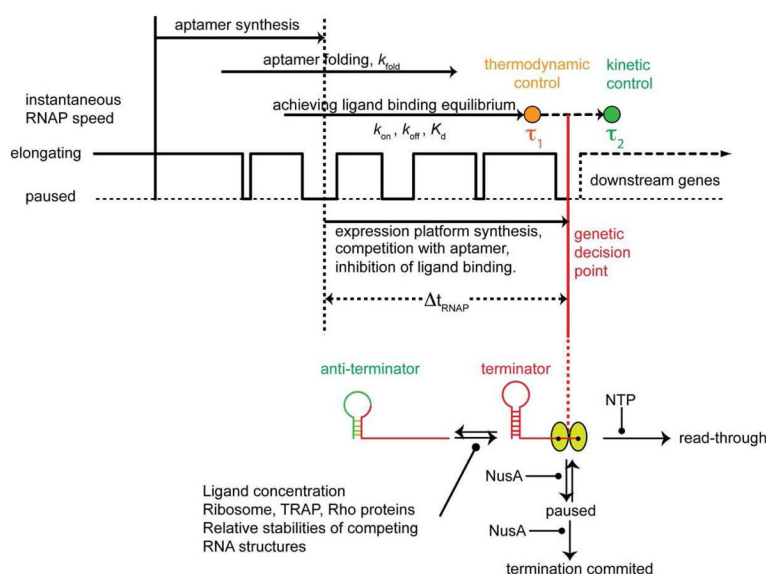


**FIGURE 3.**

Comparison of base-pairing schemes employed by three riboswitches that recognize guanine and related ligands. The four examples are the (A) purine, (B) preQ₁, and (C) first (g_I) and (D) second guanine (g_{II}) residues of c-di-GMP bound to its cognate riboswitch.

**FIGURE 4.**

Ligand recognition strategies employed by (A) the *Tetrahymena* group I intron in recognizing the base of the exogenous guanosine substrate, and (B) flexizyme in positioning the acceptor end (residue 76) of tRNA and phenylalanine.

**FIGURE 5.**

Kinetic vs. thermodynamic control of riboswitches and ribozymes. Transcriptionally controlled riboswitches can operate under kinetic control (green dot) or thermodynamic control (orange dot). Upper panel depicts events relating to riboswitch control as horizontal lines with arrowheads (not to scale). Δt_{RNAP} represents the time from the completion of aptamer synthesis to the genetic decision point (*e.g.* RNAP reaching a transcription terminator), shown as a vertical red line; τ represents the time constant for reaching ligand binding and dissociation equilibrium. When $\Delta t_{\text{RNAP}} < \tau$ (τ_2 , green), RNAP arrives at the terminator prior to ligand and aptamer reaching binding equilibrium, the riboswitch operates under kinetic control. When $\Delta t_{\text{RNAP}} > \tau$ (τ_1 , orange), RNAP arrives at the terminator after ligand and aptamer reach equilibrium and the riboswitch exhibits thermodynamic control. Lower panel depicts a simplified kinetic scheme for competing events occurring at the terminator, depicted as a strong RNA hairpin followed by a slippery U-track. Two yellow ovals depict the RNAP active site. At the terminator, RNAPs are kinetically partitioned into species that read through the terminator or that enter a reversible pause before committing to termination. Nucleotide triphosphates (NTPs) and NusA modulate rates of these events. Additionally, the relative populations of terminator and antiterminator RNAs can be modulated by ligand concentration, ribosomes etc.

Table 1

Thermodynamic and kinetic parameters for small molecule-RNA interactions

Ligand/substrate	Riboswitch or ribozyme	ΔH (kcal mol ⁻¹)	$-T\Delta S$ (kcal mol ⁻¹)	k_{on} (10 ⁴ M ⁻¹ s ⁻¹)	k_{off} (s ⁻¹)	K_d (nM)	ref.
Adenine	<i>B. subtilis</i> pbuE	n/a*	n/a	26	0.15	581	(72)
Hypoxanthine	<i>B. subtilis</i> xpt-pbuX	-35	28 ^a	n/a	n/a	3000	(15)
FMN	<i>B. subtilis</i> ribD	-15	6 ^b	20	0.002	12	(71) (47)
preQ _i	<i>F. nucleatum</i>	n/a	n/a	60	0.17	283	(37)
c-di-GMP	<i>V. cholerae</i> tfoX	n/a	n/a	1.7	1.8×10 ⁻⁷	0.011	(5)
TPP	<i>E. coli</i> thiM	-18	6 ^a	8.7	0.043	495	(28) (37)
SAM	<i>T. tengcongensis</i> metF-meth2	-20	12 ^b	n/a	n/a	1350	(47)
Lysine	<i>C. acetobutylicum</i>	-23	15 ^b	n/a	n/a	1800	(47)
Tetracycline	<i>In vitro</i> selected aptamer	-23	11 ^c	n/a	n/a	0.8	(42)
Theophylline	<i>In vitro</i> selected aptamer	n/a	n/a	17	0.07	300	(84)
Guanosine	<i>Tetrahymena</i> group I ribozyme	n/a	n/a	0.7	0.77	1.1×10 ⁵	(85)

* n/a, not available.

^a *T* = 303 K,

^b *T* = 310 K,

^c *T* = 298 K.