A functional genetic screen reveals sequence preferences within a key tertiary interaction in cobalamin riboswitches required for ligand selectivity

Jacob T. Polaski, Otto A. Kletzien, Lea K. Drogalis and Robert T. Batey*
Department of Chemistry and Biochemistry, University of Colorado, Boulder, CO 80309, USA

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ABSTRACT
Riboswitches are structured mRNA sequences that regulate gene expression by directly binding intracellular metabolites. Generating the appropriate regulatory response requires the RNA rapidly and stably acquire higher-order structure to form the binding pocket, bind the appropriate effector molecule and undergo a structural transition to inform the expression machinery. These requirements place riboswitches under strong kinetic constraints, likely restricting the sequence space accessible by recurrent structural modules such as the kink turn and the T-loop. Class-II cobalamin riboswitches contain two T-loop modules: one directing global folding of the RNA and another buttressing the ligand binding pocket. While the T-loop module directing folding is highly conserved, the T-loop associated with binding is substantially less so, with no clear consensus sequence. To further understand the functional role of the binding-associated module, a functional genetic screen of a library of riboswitches with the T-loop and its interacting nucleotides was used to build an experimental phylogeny comprised of sequences that possess a wide range of cobalamin-dependent regulatory activity. Our results reveal conservation patterns of the T-loop and its interaction with the binding core that allow for rapid tertiary structure formation and demonstrate its importance for generating strong ligand-dependent repression of mRNA expression.

INTRODUCTION
Riboswitches are regulatory RNA elements found in the 5′-leader of bacterial mRNAs that modulate gene expression via their interaction with a small molecule metabolite. Regulatory activity is achieved through communication between two functional domains: a receptor (aptamer) domain that interrogates the local environment for the cognate metabolite and a regulatory (expression platform) domain that undergoes a structural transition based on the occupancy status of the receptor to direct the gene expression machinery (1,2). Currently, much of the information about the relationship between folding and regulatory activity of riboswitches has been inferred from in vitro studies of isolated receptor domains, resulting in a limited understanding of how conformational alterations drive regulatory activity in a cellular environment. Developing a more complete understanding of this relationship is important, as gene regulation under biological conditions places riboswitches under strong kinetic constraints requiring the aptamer domain to rapidly fold into a functional structure with high fidelity, limiting the temporal window available to generate the appropriate regulatory response (3,4). While these kinetic considerations were initially thought to be restricted to riboswitches regulating transcription, where an obligate decision must be made at the poly-uridinetract of the intrinsic terminator, it is becoming increasingly appreciated that translational riboswitches are also kinetically constrained (5–8). This is because these riboswitches also employ regulatory mechanisms such as rho-dependent transcriptional termination and RNA degradation in addition to occlusion or exposure of the ribosome binding site.

Rapid acquisition of a productive ligand-binding site within the aptamer domain is generally assisted by recurrent modules and structural motifs such as kink turns, T-loops, loop E motifs, and kissing loops to establish helical packing and long-range tertiary interactions (9–12). These motifs are typically identified as part of a bioinformatic analysis of a riboswitch’s aptamer domain that yields a consensus secondary structure and nucleotide conservation pattern. For most riboswitch aptamer families, these modules are some of the most conserved sequence elements, underscoring their importance in folding and regulatory function (13,14). The consensus sequence for each of these recurrent modules is derived from a set of RNAs that vary sig-

*To whom correspondence should be addressed. Tel: +1 303 735 2159; Email: robert.batey@colorado.edu
Present address: Jacob T. Polaski, Computational Biology Program, Public Health Sciences Division, Basic Sciences Division, Fred Hutchinson Cancer Research Center, Seattle, WA 98109, USA.

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significantly in their overall structure, interaction with other RNAs or proteins, and biological function (15). However, each distinct family of RNAs likely place different demands on the structural module, such that within the consensus are subsets of sequences that are suitable for a given RNA's function. For example, the kink turn motif exhibits different preferences for sequences interacting with proteins versus those found in RNA-only structures (9). The kinetic constraints placed upon regulatory activity of riboswitches may favor a subset of variants within the consensus able to fold rapidly or form more stable long-range interactions in the absence of proteins. Differences in folding properties of the kink turn modules of ribosomes, riboswitches, and protein-binding snoRNAs have been observed (16–18). For modules found in highly represented riboswitches such as class-I cobalamin and S-adenosylmethionine-I (SAM-I), sequence alignment and comparative analysis with other families of RNAs readily identify variation in the module contributing to altered folding and functional properties that can be subsequently validated using biochemical approaches. For RNAs having a limited number of representatives or for modules that display substantial variation within a phylogenetic alignment, sequence analysis approaches are of limited utility.

Recently, a tertiary interaction mediated by a structure similar to the classic T-loop module (10) was found to be important for ligand binding in the class-II cobalamin riboswitch (19). Members of this class of cobalamin riboswitches, which contain two distinct T-loops (Figure 1A), exhibit a broad spectrum of selectivity for the two biological forms of cobalamin—5′-deoxyadenosylcobalamin (AdoCbl) and methylcobalamin (MeCbl)—such that some tagenic analysis of an AdoCbl-selective (env8Cbl-IIa) and its long-range interaction with a region of the binding module that contains a base pair between nucleotides 9 and 13, where nucleotide 13 is in the syn orientation as dictated by the antiparallel arrangement of the two nucleotides, whereas the closing pair in the T-loop is most often a Watson–Crick/hoogsteen U9–A13 pair (Figure 1C and D), although other pairings are accepted. The lone pair triloop sequence in the Gutell classification scheme that the J1/3 element of env8Cbl-IIa most closely conforms to is ‘R1’, which this scheme describes as having the closing pair in a reversed Hoogsteen conformation (Figure 2A) (21). Consideration of long range tertiary interactions with J6/3 would suggest that this module is better classified as an ‘R2’ triloop, and while J1/3 does contain a pyrimidine at position 3 of the triloop like R2 representatives, J1/3 does not conform to the requirement that the second triloop nucleotide is a uridine and makes hydrogen bonding contacts with the backbone (Figure 2B). In the Major classification scheme, which is primarily grouped according to how the bases stack, J1/3 is best described as a class I, ‘SLSL’ subclass triloop, of which T-loops are a member (Figure 2C) (22). While several members of this subclass have trans Watson-Crick/Watson-Crick closing pairs, their backbone orientation is classified as parallel, rather than the antiparallel arrangement of the closing pair in J1/3. Thus, classification of this important module in the class-II cobalamin riboswitches is difficult, and unfortunately, limited sequence conservation in J1/3 does not help to clarify this issue.

To gain further insights into the role of the J1/3–J6/3 interaction in supporting cobalamin binding and regulatory activity, we implemented a cell-based functional screen in which a set of critical nucleotides in each element were fully randomized and active variants identified. In this screen we identified 91 unique sequences with a broad range of abilities to repress expression of a fluorescent reporter protein, ranging from variants as efficient or moderately more active than the wild type riboswitch, to sequences that repressed reporter expression with only 10% of wild type activity. Analysis of these data reveal that RNAs with the highest repressive activity conform to a consensus in which the most functionally important elements are the three nucleotides in the J1/3 triloop and the Watson-Crick base pair between position 11 of J1/3 and position 70 of J6/3. This consensus also fits within T-loops found in other genetic elements. In addition, these results reveal that the J1/3–J6/3 interaction may not be essential for function, as a number of low activity variants contain no identifiable consensus in J1/3 but with an alternative consensus in J6/3. Thus, sequence alignments based upon this screen yield novel insights into both cobalamin riboswitch structure and function.
MATERIALS AND METHODS

Library preparation and cloning

The riboswitch library was constructed using two single-stranded Ultramers® synthesized by Integrated DNA Technologies encompassing the entire riboswitch sequence with seven randomized positions (16 384 total variants), and NsiI and HindIII restriction sites at the 5′ and 3′ ends, respectively. Riboswitch inserts were amplified through 20 cycles of gap-filling PCR and were ligated into a reporter plasmid containing the fluorescent protein mNeon (23) and an ampicillin resistance cassette (Amp®). Ligations were performed using the manufacturer’s protocol for T4 DNA ligase (New England Biolabs) and 3 μl of the ligation reaction was transformed into 100 μl of chemically competent E. coli strain BW25113 (ΔbtuR) by heat shock at 42°C for 30 s, which was recovered by adding 900 μl of 2x YT medium and incubated at 37°C for 1 h. 100 μl aliquots of the transformation reaction were plated onto chemically defined salt broth (CSB) media (24) containing 1.2% agar supplemented with cyanocobalamin (5 μM final concentration) and 100 μg/ml carbenicillin and grown overnight at 37°C. Once visible colonies formed, plates were incubated for an additional 24 h at 4°C to arrest cell growth and enhance the fluorescent signal. To generate counting statistics, colonies were counted following incubation at 4°C using the OpenCFU software package (3.9.0) (25).

Fluorescent colony screening

Colonies were illuminated using a light source outfitted with 490 nm excitation and 510 nm emission filters, respectively, for screening. Colonies exhibiting a similar level of fluorescence as the parent riboswitch (low on CNCbl plates) were picked and grown on gridded replica plates containing CSB media containing 1.2% agar and 100 μg/ml carbenicillin with and without 5 μM cyanocobalamin. Gridded plates were incubated at 37°C overnight to allow for cell growth
and were subsequently incubated at 4°C for 24 h to arrest cell growth and enhance the fluorescent signal.

Colonies showing cobalamin-dependent regulation of mNeon upon visual inspection were picked, transferred to 10 ml of 2xYT media supplemented with 100 μg/ml ampicillin, and grown overnight to saturation at 37°C. Plasmids were isolated from overnight cultures via miniprep (E.Z.N.A.® Plasmid Mini Kit, OMEGA bio-tek) the following day and analyzed by Sanger sequencing. Plasmids harboring unique riboswitch sequences were transformed into chemically competent E. coli BW25113 (ΔΔΔbtuR) cells, plated onto CSB medium containing 1.2% agarose supplemented with 100 μg/ml carbenicillin and grown overnight at 37°C.

**Riboswitch activity assays**

To determine levels of mNeon expression, three colonies for each riboswitch were picked and grown overnight to saturation in 5 ml of CSB media supplemented with 100 μg/ml ampicillin. The following day, 5 μl of each overnight culture was transferred to tubes containing 5 ml of CSB medium with and without 5 μM cyanocobalamin supplemented with 100 μg/ml ampicillin. Cultures were grown to mid-log phase at 37°C in a roller drum, at which time 300 μl of each culture was added to wells in a Costar® 96-well half area microplate. Expression of mNeon was measured at an excitation wavelength of 490 nm and a 517 nm emission wavelength using a Tecan Infinite M200® PRO plate reader.

Levels of mNeon expression were determined from a single technical replicate of three biological replicates, which were normalized to the cell density (OD600) in each well and background corrected using cell density normalized fluorescence from wells containing a pBR327 empty vector. Fold repression was calculated by dividing the average normalized background corrected fluorescence of the unpressed condition (absence of cyanocobalamin) by the average normalized background corrected fluorescence of the repressed condition (presence of cyanocobalamin).

**Data processing and phylogenetic tree construction**

Sequence alignments were performed using the Multiple Sequence Comparison by Log-Expectation (MUSCLE) alignment tool in Jalview and the phylogenetic tree was generated using the Geneious software package (version 11.0.2, https://www.geneious.com) (26). Tree distances and sequence relatedness was determined by the method of Tamura and Nei (27) using a neighbor-joining model with no outgroup, which was converted to a sunburst format to aid visualization.

**RESULTS**

**Design and implementation of the screen**

To obtain a functionally active library of J1/3–J6/3 variants of the env8Cbl-IIa riboswitch, nucleotides 9–13 of J1/3 and nucleotides 69 and 70 of J6/3 were fully randomized and yielded 384 theoretically possible variants (red dots, Figure 3). This complete variant library was cloned into a reporter plasmid with the riboswitch in the leader sequence of an mRNA encoding the fluorescent protein mNeon (23), whose transcription is driven by a moderately strong synthetic insulated constitutive promoter ‘proD’ (28). This reporter system has been previously validated and used to investigate the mechanism of cobalamin-dependent gene regulation and ligand selectivity of the env8Cbl-IIa cobalamin riboswitch (5,20). To increase the dynamic range of the regulatory response, an env8Cbl-IIa variant was used that features an extension of a linker sequence (J1/13) connecting the receptor and regulatory domains from 7 nucleotides in
Library variants

J1/3
J6/3

(1)

(2)

(3)

+ CNCbl

-CNCbl

- Activity assays
- Sequencing

Figure 3. General workflow for the cell-based screening strategy. A simplified schematic of the parental env/Cbl-Ha riboswitch is featured at the top with the randomized nucleotides in J1/3 and J6/3 colored red. In step (1), library variants are transformed into E. coli and plated on CSB agar containing CNCbl. In step (2), colonies are selected that exhibit similar levels of mNeon expression as the parental sequence and transferred to gridded plates where they are grown either in the presence or absence of CNCbl. In step (3), colonies that exhibit CNCbl-dependent repression of mNeon expression upon visual inspection are harvested and used for activity assays in liquid culture and sequenced.

The wild type riboswitch to 19 nucleotides (5). Lengthening this region of the RNA increases regulatory activity by increasing the amount of fluorescent protein expression in the absence of cobalamin while retaining the ability to repress gene expression to wild type levels in the presence of ligand. This alteration does not affect ligand binding to the aptamer but does improve the ability to visually sort riboswitch variants that demonstrate cobalamin-dependent gene regulation. The library of plasmids was transformed into E. coli strain BW25113 (ΔbtuR), which is defective in cob(II)yrinic acid a,c-diamide adenosyltransferase, an enzyme that adenosylates cobalamin during import (29,30), ensuring that cyanocobalamin (CNCbl) used in the growth medium is not converted to AdoCbl.

To ensure that colonies were sufficiently separated for visual screening while maximizing the degree of library coverage per round of screening, transformation reactions were optimized to yield ~150–200 colonies per 60 mm diameter plate. To obtain CNCbl-responsive riboswitches, the transformed library was screened using a two-step approach. In the first step, the transformed library was plated on a rich, chemically defined medium (24) containing 5 μM CNCbl (step (1)), Figure 3. Colonies were visually screened for expression of the mNeon reporter and manually picked if they exhibited a fluorescent signal similar to the parent riboswitch (dark colonies, Supplementary Figure S1). Selected colonies were then grown on gridded plates (step (2), Figure 3) in the absence or presence of 5 μM CNCbl and cells that showed visual differences in their brightness were used for downstream analysis and characterization (step (3), Figure 3).

In four rounds of screening, 74 300 colonies were surveyed, representing ~4.5-fold redundancy (Supplementary Table S2). Theoretical calculation suggests this level of redundancy results in ~99% probability of observing each unique sequence in the original pool of variants (31), but in practice, this is likely lower. In total, four rounds of screening yielded ~2578 colonies that were selected for secondary analysis using gridded plates, which resulted in 146 variants that exhibited visually detectable CNCbl-dependent repression of mNeon. Each positive switch was sequenced, resulting in 91 unique variants (Supplementary Table S3).

Riboswitch variants cluster into well-defined classes based on sequence

Analysis of the functionally active variant library was initiated by sequence alignment of the 91 variants using the Multiple Sequence Comparison by Log Expectation (MUSCLE) software package (Supplementary Figure S2) (32), and a cladogram calculated to visualize related sequence clusters using the neighbor joining method of Tamura and Nei (Figure 4A) (27). At the topmost level, the cladogram is readily divided into three major classes (yellow, cyan and magenta sectors) with two outlying sequences that do not group with the three major clusters (green sector). The cladogram can be further divided into eight subclasses (SC-I to SC-VIII) to provide more detailed information about conservation patterns within smaller related sub-clusters. Additionally, analysis of consensus sequences was informed using activity assays performed in liquid culture to determine the fold repression of all 91 unique variants (red circles, Figure 4A and Supplementary Figure S3 and Supplementary Table S4).

Highly active variants cluster into one major grouping

One major class of the cladogram is largely populated with sequences exhibiting strong repressive activity and includes the parental riboswitch (red star, Figure 4A). The consensus sequences of J1/3 and J6/3 show that members of this class generally preserve the Y11–G70 base pair observed in the parental riboswitch, have a nearly invariant guanosine at position 10, a near equal preference for an adenosine
Figure 4. Cladogram and consensus sequences for variants found in the screen. (A) Sunburst style cladogram depicting well-defined clusters of related sequences along with activity assay data obtained from liquid culture assays. Major sequence classes are separated using colors (yellow, cyan, and magenta), with two unrelated outlying sequences highlighted in green. Relative fold repression values from activity assays for each variant is represented with a red circle in the first shell, with darker red circles denoting riboswitches with higher dynamic ranges. Blue circles in the second shell represent relative values of cell density normalized fluorescence for cells grown in the absence of CNCbl, and blue circles in the outermost shell represent cell density normalized fluorescence of cells grown in the presence of CNCbl. In both cases, circles with a darker blue color indicate higher levels of mNeon expression. The three major classes are further divided into subclasses, which are shown using the designation ‘SC-X’, and the parental riboswitch used to design the library is denoted with a red star in the yellow sector. (B) Sequence logo representation (52) of consensus sequences for J1/3 (nucleotides 9–13) and J6/3 (nucleotides 69 and 70) for the subclasses shown in panel A. All sequence logos were generated using the WebLogo 3.5.0 software package (53). The secondary structural interactions in wild type J1/3–J6/3 are shown in the ‘yellow’ panel and use the same notation as Figure 1D. Note that these interactions may not be present in all sequences.

or guanosine at position 12, and a moderate preference for a uridine at position 69 as the intercalating base. Notably, members of this class show a high diversity of nucleotides comprising the closing base pair between positions 9 and 13 of J1/3, including non-Watson–Crick C–A, C–C, U–U, C–U and A–A pairs in sequences with high dynamic range, indicating this portion of the loop is highly tolerant of diverse potential base pairs.

Within this class, a subset of sequences (SC-I, Figure 4B) has a consensus in which three nucleotides of J1/3 (nucleotides 10–12) and position 70 of J6/3 are almost invariant, suggesting the most important feature of the tertiary interaction is maintaining a Watson–Crick C11–G70 base pair, which is invariant in the group. The structural role of nucleotide 10 is to stabilize the turn in J1/3 via interaction between the Watson–Crick face of guanosine and the phosphate moiety of nucleotide 13. It should be noted that within the T-loop and the lone triloop module, the nucleotide corresponding to nucleotide 10 is often a uridine, whose base can make an equivalent set of contacts with the backbone (10). Like the yellow class as a whole, the closing pair of J1/3 shows no conservation pattern or pairing preference–potential C–C, C–U, U–A, C–A, G–G and U–G pairs are observed, which can all potentially form a reverse Watson–Crick or a wobble base pair with two hydrogen bonds. However, the wild type A9–U13 pair was not ob-
served, which likely reflects that the screen did not capture all possible active variants. Position 69 shows a near equal preference for uridine or adenosine, with only one member containing a cytosine at this position; it is not clear why cytosine and guanosine should be so strongly selected against. The variant with the lowest dynamic range in this subclass (GGCGG/UG, 2.2-fold repression) is the only member that has a guanosine rather than an adenosine at position 12. However, it should be noted that in all members of SC-II discussed below, which is also enriched in highly active variants, this position is always a guanosine, indicating that this alone does not abrogate regulatory activity.

A second subset of sequences (SC-II, Figure 4B) is largely comprised of highly active variants, and not surprisingly, the consensus within this subclass is very similar to SC-I. What differentiates this class from SC-I is the nucleotide identity of position 12, which is exclusively a guanosine rather than an adenosine. Within this subclass, the least active variant (CGGC/C/G, 2.9-fold repression) is the only member with a potential C9–C13 pair in J1/3, a type that is represented in SC-I in two highly active sequences (CGCAC/UG, 14-fold repression and CGCAC/AG, 12-fold repression), suggestive of covariation between the identity of position 12 and the closing base pair in J1/3. Overall, SC-I and SC-II contain sequences most likely to exhibit robust gene regulation in a biological context, a result consistent with the presence of the parental sequence in SC-II (star, Figure 4A).

The third subclass of the yellow sector (SC-III, Figure 4B) contains variants with modestly lower activity (2.8–7.2-fold repression) than sequences in SC-I and SC-II. The major difference between SC-III compared to SC-I and SC-II is the presence of a U11–G70 base pair in all sequences except for one with low dynamic range (UGUGA/A, 3.0-fold repression). SC-III is also enriched in sequences that contain either a U9–A13 or U9–U13 closing base pair in J1/3, which is found in more highly active SC-I representatives, suggesting that a more optimal closing lone pair is required to compensate for a weaker 11–70 G–U pair. Additionally, SC-III has a strong preference for an intercalating uridine at position 69; one sequence has an adenosine at this position and represents the lowest activity member (UGUGA/AG, 2.8-fold repression). An adenosine at position 69 in SC-I and SC-II is well-tolerated, suggesting that the presence of the U11–G70 wobble pair imposes a requirement for the more optimal uridine at position 69.

Together, the sequences of SC-III reveal that deviations from the optimal triloop and Watson–Crick 11–70 pair can be compensated for by other features of this module.

A second major class largely includes sequences with moderate activity

The second major cluster of variants in the cladogram is populated by sequences exhibiting moderate dynamic range compared to the parental riboswitch (magenta sector, 5.4 median fold repression; Figure 4A). Like the parental sequence, this class contains many variants (11/30 sequences) that have the potential to form a closing Watson–Crick/Hoogsteen base pair between A9 and U13 in J1/3, as well as many variants (14/30 sequences) with a uridine as the intercalating base at position 69 and an adenosine at position 12 (26/30 sequences), which is also observed in members of the yellow class. However, unlike the parental riboswitch and RNAs in the yellow class, only five variants in the magenta class can form a C11–G70 Watson–Crick pair between J1/3 and J6/3 (most variants have a U11–A70 base pair at this position), further reinforcing that this tertiary contact plays a significant role in the assembly of the cobalamin-binding core and efficient gene regulation by the RNA. The loss of the optimal C11-G70 pair is likely to be partially compensated for by optimal closing 9–13 pairing.

Within this major class, a subset of variants (SC-VI, Figure 4B) with moderate dynamic ranges (3.9- to 5.7-fold repression) feature a conserved non-Watson-Crick closing A–A base pair in J1/3, an invariant adenosine at position 12, and either a C11–G70 or U11–G70 base pair between J1/3 and J6/3. Notably, although members of this subclass contain sequence features present in more active RNAs from SC-I and SC-II, the main difference in SC-VI representatives is a nearly invariant cytosine at position 10 (3/4 sequences), which is almost always a guanosine in SC-I and SC-II. While the nucleotide identity of position 10 could have a strong effect on regulatory function, the low number of sequences in this subclass (4) makes it difficult to draw strong conclusions using its consensus.

A second subclass within the magenta class (SC-VII; Figure 4B) contains variants with a broad spectrum of dynamic ranges (1.2- to 14.0-fold repression), but is largely populated with sequences exhibiting regulatory activity near the center of this distribution (5.3 median fold repression). SC-VII features three highly conserved positions within J1/3 (nucleotides G10, U11 and A12) and an invariant adenosine at position 70 in J6/3. This means that, unlike members of highly active subclasses SC-I and SC-II, SC-VII representatives almost invariably have a U11–A70 base pair rather than a C11–G70 pair. Notably, the variant of SC-VII with the highest dynamic range (CGUAA/UA; 14.0-fold repression) features a suboptimal U11–A70 base pair, which may be offset by the presence of G10 and an intercalating uridine that is found in highly active SC-I and SC-II representatives. Members of SC-VII also show some preference for a U–A as the closing pair of the J1/3 triloop, however this preference is not strong, as RNAs with A–A, U–U, C–A and U–G pairs are also found in this subclass.

Like SC-VII, SC-VIII is populated by variants with a broad range of regulatory activities (1.6- to 15.2-fold repression), but on average contains sequences with similar dynamic ranges as SC-VII (5.5 median fold repression). SC-VIII representatives show a strong preference for a closing A–U pair in the J1/3 triloop and have a lower degree of nucleotide conservation at positions 11 and 70; variants with U–A, C–A, U–G, A–G and C–G base pairs are found in this subclass. Similar to highly active RNAs found in SC-I and SC-II, the consensus for SC-VIII shows an almost universal conservation of a guanosine at position 10, but unlike all other subclasses shows a moderate preference (8/14 sequences) for a cytosine as the intercalating base at position 69.
A third major class contains variants with low levels of regulatory activity

The third major class (cyan sector; Figure 4A) is dominated by sequences with low repressive activity compared to the parental sequence (the highest is 5.0-fold repression with sequence CUUAA/GA) and significantly less nucleotide conservation throughout J1/3 compared to other subclasses. The most highly conserved nucleotide in this class is the intercalating base at position 69, which is predominantly a guanosine.

There are two subclasses within this class (SC-IV and SC-V; Figure 4B) that show low degrees of nucleotide conservation in J1/3 and J6/3 compared to all other subclasses. Like the cyan class as a whole, variants comprising SC-IV show nearly universal conservation of a guanosine at position 69, a moderate preference for a uridine at position 10, and either adenosine or guanosine at position 70. While representatives of the second subclass (SC-V) show universal conservation of a guanosine at position 69, these RNAs show a moderate preference for a U9–A13 closing pair of salconservation of aguanosine at position 69, these RNAs representives of the second subclass (SC-V) show univer-

sation throughout J1/3 and J6/3, and either adenosine or guanosine at position 70. While representatives of the second subclass (SC-V) show universal conservation of a guanosine at position 69, these RNAs show a moderate preference for a U9–A13 closing pair of the J1/3 triloop, which is the most highly represented base pair (4/8 sequences). Like highly active variants in SC-I and SC-II, some riboswitches in SC-V feature a C11–G70 base pair (3/8 sequences) between J1/3 and J6/3; however, other sequences in this subclass have non-Watson–Crick G–A or G–G pairs at this position. This suggests that while some members of the cyan class contain the optimal C11–G70 base pair, a combination of other non-optimal features in the motif contribute to the lower dynamic ranges observed in both subclasses.

The two outlying sequences (green sector; Figure 4A) have J1/3 and J6/3 sequences: GUCAU/GG and GUCGU/UG and exhibit low dynamic range compared to the parental sequences (3- and 2-fold repression, respectively). Although both sequences can form the optimal C11–G70 base pair observed in active variants comprising SC-I and SC-II, other sub-optimal features, including a guanosine at position 9, a uridine at position 10, and for one sequence a guanosine as the intercalating base, likely diminish the regulatory activity of these two variants.

Analysis of consensus sequences by activity-based clustering

An alternative way to group sequences obtained from the screen is by their dynamic range relative to the wild type sequence (14.0-fold repression of reporter gene expression). To achieve this, variants were categorized into four groups: >75%, >50–75%, >25–50% and >10–25% of wild type activity, as described below.

Riboswitches with >75% wild type activity

This group contains all sequences exhibiting greater than or equal to 10.5-fold repression of mNeon expression in the presence of CNCbl. The consensus sequences of J1/3 and J6/3 in this cluster (Figure 5A) are highly similar to SC-I and SC-II, which is expected as the majority of these sequences originate from these subclasses. The only members not from SC-I and SC-II are AGCAU/CN (15.2-fold repression) and CGUAU/UA (14.0-fold repression), found within SC-VIII, and SC-VII, respectively. Overall, the consensus indicates a strong preference for guanosine and cyto-

sine at positions 10 and 11 in the J1/3 triloop, respectively, and a modest preference for either adenosine or guano-

sine at position 12. Additionally, like the parental sequence, members of this cluster show a preference for a uridine as the intercalating base at position 69; however, 4/15 RNAs tolerate an adenosine in this position and 2/15 feature a guanosine. Members of this cluster also have an invariant guanosine at position 10, which forms key contacts with the backbone to enforce the turn of the triloop, as well as a strong preference for a C11–G70 Watson–Crick base pair between J1/3 and J6/3 (only 1/15 sequences has an alter-

native U11–A70 base pair). The consensus for J1/3 shows a weak preference for the identity of the closing base pair between positions 9 and 13, which is noteworthy because if all the sequences in this cluster conform to having a reverse Watson–Crick or wobble (9-anti/13-syn) base pair scheme, seven types of pairs can support this arrangement (C–A, U–A, A–U, C–C, U–U, A–A and U–G).

Riboswitches with >50% to 75% wild type activity

The consensus sequences of J1/3 and J6/3 in this group are substantially relaxed compared to the most active RNAs (Figure 5B). In this cluster, G10 remains the most highly conserved position, although uridine is also infrequently observed at this position (2/9 sequences). The identity of the intercalating base at position 69 is more variable than the more active sequence cluster, and positions 11 and 70 show a lesser preference for a C11–G70 Watson–Crick pair, with a U11–A70 base pair occurring in 2/9 and a U11–G70 pair in 2/9 representatives. There remains a strong prefer-
ence for a purine at position 12, although there is a single example of a uridine at this position, which is also the sequence that contains a non-Watson–Crick A11–G70 base pair (UGAUU/CG). Finally, the identity of nucleotides comprising the closing pair between positions 9 and 13 are similar to the most active group, with the exception of a single sequence with a C9–U13 pair that is also capable of forming a reverse Watson–Crick pair with two hydrogen bonds. This arrangement would place the C(O2) and U(O4) in close proximity, suggesting that an unstable pair or even no pairing at this position still supports the tertiary interaction between J1/3 and J6/3.

**Riboswitches with >25% to 50% of wild type activity**

This large group of variants (34 sequences) has a further relaxed consensus sequence compared to the more active clusters (Figure 5C). The preference for a guanosine at position 10 is significantly reduced, particularly among members with the lowest repressive activity, which typically have a cytosine or adenosine. Within this group there appears to be no preference for a particular nucleotide at position 69, suggesting that the weak interaction between nucleotides 10 and 69 is not critical for establishing the interaction between J1/3 and J6/3. Furthermore, while many members have a Watson–Crick base pair between positions 11 and 70, potential non-Watson–Crick pairs at this position are increasingly observed, including U11–G70 (6/34 sequences), C11–A70 (2/34 sequences), G11–G70 (1/34 sequences) and A11–G70 (1/34 sequences). However, despite the increased variation in pairing schemes between positions 11 and 70, the strong preference for Y11–R70 remains. Similar to the more active clusters, there is only weak conservation of the identity of the closing base pair between positions 9 and 13, with G9–G13 and G9–A13 pairs observed along with the previously noted base pairs.

**Riboswitches with >10% to 25% of wild type activity**

This group represents a set of 32 RNAs that exhibit weak regulatory activity that ranges from 3.4- to 1.6-fold repression. The consensus of the J1/3 element of this cluster does not show a conservation pattern consistent with the lone pair triloop motif or T-loop module (Figure 5D). Instead, the only well-defined consensus occurs in J6/3 at position 70, which is always a purine, but less than half (14/32) have a nucleotide at position 11 that can form a Watson–Crick base pair. Representatives of this cluster also show a modest preference (19/32 sequences) for a guanosine as the intercalating base at position 69, which deviates from the preference for a uridine in more highly active sequences. Overall, the consensus sequences of members from this group suggest the interaction between J1/3 and J6/3 in these RNAs is weak or potentially absent. In this case, the lack of this tertiary interaction weakens the affinity of the binding pocket for cobalamin (19), which impairs the ability of the RNA to inhibit translation initiation.

**DISCUSSION**

In this work, the sequence preferences of a key tertiary interaction in a class-II cobalamin riboswitch was investigated using a cell-based functional screen to generate an experimental phylogeny of an RNA structural module critical for cobalamin selectivity and regulatory function. This approach allows for the unbiased generation of a diverse set of unique riboswitch variants, where differences in the sequence composition and their impact on regulatory activity are assessed in the same overall structural context. A screening-based approach with manual colony screening was chosen because this ensured we would be able to access variants that span a broad spectrum of repressive activities, rather than only the most active variants that would likely emerge from selection approaches. These low activity variants have the potential to reveal aspects of the module or RNA of interest that the most active sequences cannot. Colony screening also enables functional testing of each active variant in an integrated workflow to yield a dataset that reveals correlations between sequence and regulatory activity. While this result can be achieved using higher-throughput approaches, the limited size of this library and nature of the activity screen made colony screening an optimal strategy despite being labor-intensive. Furthermore, this approach is generalizable to study myriad RNA structural motifs, which can be used to understand evolutionary trajectories and fitness landscapes of a wide range of functional RNAs, as has been done previously for proteins (33,34) and catalytic RNA (35).

The screening strategy employed in this study yielded 91 unique riboswitch variants with a broad range of biological activity. The most active variants cluster into three sub-classes (SC-I, SC-II and SC-III) that together share a highly similar consensus. These RNAs do not have a strong preference for the identity of the closing pair of the lone pair triloop in J1/3, which is consistent with previous large-scale surveys of this motif in various rRNA and tRNA crystal structures that revealed the presence of diverse non-Watson–Crick base pairs at this position (21). Instead, the most important features are the sequence of the triloop itself and the Watson–Crick pair between nucleotides 11 and 70. SC-I and SC-II representatives have a universally conserved C11–G70 Watson–Crick base pair between J1/3 and J6/3, while the consensus of members from SC-III, which are generally less active than SC-I and SC-II variants, feature either U11–G70 or U11–A70 base pairs at this position. Overall, the nucleotide conservation patterns of sequences supporting >50% of wild type activity strongly correlate with the T-loop conservation pattern in genetic elements (Figure 6) (10). Thus, this element should be classified as the second T-loop motif in the Cbl-II riboswitch family—the first being L4 (Figure 1A). It should be noted that the T-loop is part of the larger lonespair triloop motif as defined by Gutell (21) and Lisi and Major (22).

The crystal structure of the env8Cbl-IIa aptamer in complex with hydroxocobalamin provides structural insights into the observed sequence preferences (19). A strong base pairing interaction between J1/3 and J6/3 likely serves to prevent alternative pairing between J3/4 and J6/3 that would collapse the cobalamin binding pocket, thereby preventing translational repression. A rapidly acquired and stable J1/3 structure is thus likely to be important in establishing the cobalamin binding pocket on a timescale conducive to regulatory function. Our data suggest that a
gene expression in the presence of ligand, reflecting a lower repressive activity typically allow for higher levels of activity. This hypothesis is supported by our data. In fact, the strong preference for a Y11–R70 base pair. Furthermore, cytosine, which is not tolerated across all structurally defined T-loops, is observed in our data. In contrast, the T-loop in the riboswitch motif is less accommodating of diversity in the closing pair. This likely reflects the functional role of the T-loop in the riboswitch. Another constraint for the sequence of the J1/3 T-loop is the kinetic pressure to fold both rapidly and reliably, as the coupling of transcription and translation in bacteria places strong temporal requirements on sensing the occupancy status of the receptor to produce the appropriate regulatory outcome. Furthermore, many RNAs fold under the influence of proteins and chaperonins, which places a different set of evolutionary constraints on their sequence composition, as the RNA bases must make productive contacts with amino acids. Thus, it is not surprising that there are clear differences in sequence preferences for T-loops found in tRNAs, rRNAs and between different genetic elements.

Ultimately, for riboswitches and other functional RNAs, the cellular environment places constraints on the allowable sequence space to achieve proper folding and the desired cellular function. The screening strategy in this work uncovered dozens of variants with dynamic ranges spanning a broad range of repressive activity, suggesting the sequence composition of the J1/3-J6/3 interaction plays an important role in the folding of this riboswitch and the subsequent regulatory response. While additional work is required to determine the effect of different J1/3 T-loop sequences on folding kinetics, in the case of cobalamin riboswitches, this element likely serves a central role in rapidly establishing a productive binding pocket for cobalamin to ensure repression of gene expression on the timescales required for a co-transcriptional/translational regulatory mechanism. Recurrent motifs in many other functional RNAs may act in a similar fashion as critical nodes along folding pathways, and developing a more comprehensive understanding of allowable and optimal sequence space for these modules will have broad implications for RNA structure prediction and the design of synthetic RNA-based devices.
SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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