Structure-seq2 probing of RNA structure upon amino acid starvation reveals both known and novel RNA switches in *Bacillus subtilis*

LAURA E. RITCHEY,1,5 DAVID C. TACK,2,6 HELEN YAKHNIN,3,4 ELIZABETH A. JOLLEY,1,4 SARAH M. ASSMANN,2 PHILIP C. BEVILACQUA,1,3,4 and PAUL BABITZKE3,4

1Department of Chemistry, 2Department of Biology, 3Department of Biochemistry and Molecular Biology, Pennsylvania State University, University Park, Pennsylvania 16802, USA
4Center for RNA Molecular Biology, Pennsylvania State University, University Park, Pennsylvania 16802, USA

ABSTRACT

RNA structure influences numerous processes in all organisms. In bacteria, these processes include transcription termination and attenuation, small RNA and protein binding, translation initiation, and mRNA stability, and can be regulated via metabolite availability and other stresses. Here we use Structure-seq2 to probe the in vivo RNA structurome of *Bacillus subtilis* grown in the presence and absence of amino acids. Our results reveal that amino acid starvation results in lower overall dimethyl sulfate (DMS) reactivity of the transcriptome, indicating enhanced protection owing to protein binding or RNA structure. Starvation-induced changes in DMS reactivity correlated inversely with transcript abundance changes. This correlation was particularly pronounced in genes associated with the stringent response and CodY regulons, which are involved in adaptation to nutritional stress, suggesting that RNA structure contributes to transcript abundance change in regulons involved in amino acid metabolism. Structure-seq2 accurately reported on four known amino acid-responsive riboswitches: T-box, SAM, glycine, and lysine riboswitches. Additionally, we discovered a transcription attenuation mechanism that reduces *yfmG* expression when amino acids are added to the growth medium. We also found that translation of a leader peptide (*YfmH*) encoded just upstream of *yfmG* encoded just upstream of *yfmG* expression. Our results are consistent with a model in which a slow rate of *yfmH* translation caused by limitation of the amino acids encoded in *YfmH* prevents transcription termination in the *yfmG* leader region by favoring formation of an overlapping antiterminator structure. This novel RNA switch offers a way to simultaneously monitor the levels of multiple amino acids.

Keywords: transcription attenuation; translation control; riboswitch; RNA structure; stress response; amino acid starvation

INTRODUCTION

Bacteria have the capacity to adapt to changing nutritional conditions via sweeping changes in gene expression. These regulatory events are tightly controlled such that various metabolic products are produced in sufficient quantities to sustain growth. Although regulation of transcription initiation plays a critical role in controlling gene expression, a variety of regulatory events that occur after transcription initiation play pivotal roles in modulating gene expression. Several of these posttranscriptional mechanisms rely on RNA structural features to regulate the level of transcription and translation. Some of these mechanisms involve binding of metabolites, proteins, and small RNAs (sRNAs) to the mRNA, leading to regulated transcription termination (attenuation and antitermination), translation initiation, and mRNA stability (Gollnick and Babitzke 2002; Grundy and Henkin 2004; Babitzke et al. 2009; Waters and Storz 2009). RNA aptamers bind to a ligand (e.g., metabolite, RNA, or protein) to regulate gene expression through structural switching. These RNA switches change structure upon ligand binding and cause the refolding of neighboring nucleotides to alter transcription termination or ribosome binding. For instance, intrinsic transcription termination requires an RNA hairpin followed by a U-rich sequence. In several transcription
attenuation mechanisms, formation of an antiterminator (AT) structure allows transcription readthrough by preventing formation of an overlapping terminator hairpin. Formation of the AT can in turn be prevented by an upstream anti-antiterminator (AAT) structure that overlaps the AT, and thus promotes formation of the terminator to cause transcription termination. An RNA aptamer involving these alternative structures allows control of transcription termination by altering the availability of the sequences involved and demonstrates the importance of RNA structure in bacterial gene regulation (Gollnick and Babitzke 2002).

Recent advances in genome-wide RNA structure probing methods allow investigation of these essential features of a transcriptome in vivo. We developed Structure-seq2 (Ritchey et al. 2017), an improved version of the Structure-seq method (Ding et al. 2014) that probes all RNAs directly in cells such that the effects of the native physico-chemical environment are taken into account. While the readout of in vivo chemical reactivity is reflective of a variety of cellular mechanisms including protein binding, we describe the reactivity patterns as indicative of RNA structure. The global set of RNA structures derived from this method is referred to as the “RNA structurome.” Other genome-wide RNA structure-probing methods have been used in bacteria, some of which focus on in vitro analysis of RNA structure with different conditions to identify RNA aptamers (Del Campo et al. 2015; Righetti et al. 2016; Tapsin et al. 2018). Recent in vivo studies on the model gram-negative bacterium Escherichia coli compared RNA structure in vivo to that in vitro and to that with stalled ribosomes to relate RNA structure to translation efficiency, revealing that transcripts with more stable structures have a lower translation efficiency, although the causality is not clear (Burkhardt et al. 2017; Mustoe et al. 2018). Additionally, several highly abundant static RNA structural features were identified upstream of ribosomal protein coding genes that could play important biological roles (Mustoe et al. 2018). These studies made important contributions to global structural trends in E. coli and highlighted some important RNA structures. In the present study, we examined global RNA structural changes that occur in vivo due to amino acid starvation in the model gram-positive organism Bacillus subtilis. These changes provide insight into how bacteria use RNA structure to respond to metabolite availability and nutritional stress.

The stringent response in B. subtilis is a global regulatory system that allows bacteria to respond to nutritional stress, including amino acid starvation, through the use of the alarmone (p)pGpp (Eymann et al. 2001). The stringent response diverts cellular resources away from cellular growth and division and into the production of amino acids, as compared to the general stress response system that responds in a more nonspecific protective manner (Voelker et al. 1995; Hecker and Volker 2001). The global proteome and transcriptome changes of the stringent response in B. subtilis have been studied (Eymann et al. 2002), but global changes to the RNA structurome have yet to be considered. Since RNA structure can change rapidly in response to the availability of metabolites, we examined RNA structurome changes that occur in response to amino acid starvation. We also compared the changes in the RNA structurome under this stress condition to known regulons in B. subtilis, as well as to known amino acid responsive RNA switches. After demonstrating the ability of Structure-seq2 to report on these features within B. subtilis, we used our data to predict unknown regulatory features. We identified several localized regions in which DMS reactivity of residues changed in response to amino acid limitation. One of these regions upstream of yfmg was explored in more detail and we found that yfmg is regulated by a transcription attenuation mechanism that responds to amino acid availability. This mechanism requires yfmh, a 23 amino acid leader peptide coding sequence just upstream of yfmg. Our results are consistent with a model in which a reduced rate of yfmh translation caused by amino acid limitation promotes transcription readthrough into the yfmg coding region.

RESULTS

Structure-seq2 libraries

To obtain the in vivo genome-wide RNA structure landscape, B. subtilis was grown to mid-exponential phase in the presence of all amino acids (+AA), or in the absence of amino acids except for glutamine (−AA); glutamine is an important nitrogen source and was included to improve growth. DMS treatment in a single-hit kinetics range (Supplemental Fig. S1) was performed, which results in methylation of the Watson–Crick (WC) face of solvent-exposed A and C nucleobases. RNA was then extracted, depleted of rRNA, and subjected to Structure-seq2. In this method, reverse transcription with random hexamer primers results in cDNAs that terminate immediately before the site of DMS methylation. Additional library preparation steps allow for cDNA amplification and Illumina sequencing (Ritchey et al. 2017, 2019). After background subtraction using a library generated without DMS, solvent-exposed bases in reads that mapped to the transcriptome were identified. Comparison of DMS reactivity profiles ±AA was used to assess RNA structural changes that occurred in response to amino acid starvation. Each condition (+AA and ±DMS) was performed in biological triplicate. We generated a total of 287M Structure-Seq2 reads that passed initial filtering (Supplemental Table W.1). Aligning these high-quality reads against a custom transcriptome yielded a 77% average rate of mapping (Supplemental Table W.2). These mappings were then filtered to remove alignments with a mismatch on the first base pair or more than three total.
mismatches via StructureFold2 (Supplemental Table W.3; Tack et al. 2018). The efficacy of our DMS treatment was confirmed by high A and C specificity (Supplemental Table W.4; Supplemental Fig. S2). When +DMS replicates of each growth condition were pooled, coverage was sufficient (greater than 1 reverse transcriptase [RT] stop per A or C in the transcript) (Ding et al. 2014; Su et al. 2018; Ritchey et al. 2019; Waldron et al. 2019; Tack et al. 2020) to resolve 2602 (+AA) and 2600 (−AA) transcripts, with an overlap of 2333 transcripts. The RT stops of +DMS replicates of both +AA and −AA treatments were well-correlated for all resolvable transcripts within that condition (Supplemental Fig. S3).

Transcriptome-wide trends

The calculated DMS reactivity patterns for each transcript and its three regions (5’UTR, coding sequence [CDS], and 3’UTR) were analyzed to determine global effects of amino acid starvation. The overall trend was decreased DMS reactivity upon amino acid starvation for the whole transcript as well as the individual regions, although certain transcripts increased (Fig. 1A–D; Supplemental Tables T.2, D.1–D.4). These observations imply a more structured and/or protein-protected structurome under amino acid starvation conditions.

We also ascertained transcriptome-wide changes in the Gini index (Fig. 1E–H), as applied to DMS reactivity. Gini is a measure of the unevenness of the data, where a low Gini score has been found to be characteristic of unstructured regions of a transcript, as all nucleotides would have similar DMS reactivity, and a high Gini score reflects more structured regions (Rouskin et al. 2014). We found that the Gini score increased significantly in the 5’ and 3’UTRs in response to amino acid starvation (Fig. 1F,H; Supplemental Tables T.2, D.2, D.4). Decreasing DMS reactivity coupled with increasing Gini index in the 5’ and 3’UTRs supports greater RNA structure and is generally consistent with reported lowered translation efficiency (Burkhardt et al. 2017), as expected from amino acid starvation.

Change in DMS reactivity is inversely correlated with change in transcript abundance

In addition to structural information, Structure-seq2 provides transcript abundance, similar to RNA-seq (Su et al. 2018; Tack et al. 2018). Previously, we found that changes in transcript reactivity are inversely correlated with changes
in transcript abundance following 10 min of heat shock in rice seedlings (Su et al. 2018) and after persistent salinity stress in Arabidopsis (Su et al. 2018; Tack et al. 2020). Those transcripts with increased DMS reactivity following heat shock or salt stress are thus less innately structured or protein protected and may have increased exposure to degradation pathways. On the other hand, those transcripts with decreased DMS reactivity following heat shock or salt stress may bind proteins or have increased RNA folding (salt stress only) to gain protection from such mechanisms (Su et al. 2018; Tack et al. 2018). We investigated changes in transcript reactivity with changes in transcript abundance (Supplemental Tables D.5, D.6) for B. subtilis upon amino acid starvation, in part to see whether this trend is preserved in another domain of life.

We observed that globally, the change in mean transcript reactivity was inversely correlated with the change in abundance of that transcript (Fig. 2), just as was found in rice and Arabidopsis. Transcripts that exhibited increased DMS reactivity under amino acid starvation generally exhibited decreased abundance, whereas transcripts that exhibited decreased DMS reactivity in the absence of amino acids exhibited increased abundance (Fig. 2A; Supplemental Tables T.3, D.1–D.6). This inverse relationship was observed when comparing whole transcripts as well as the CDS, while the UTR regions did not show this same inverse correlation (Fig. 2B–D).

Previous studies in the model plant species Arabidopsis thaliana showed that more reactive transcripts were more abundant within a given condition (i.e., comparing absolute amounts rather than changes in amount) (Li et al. 2012), so we sought to confirm if this was also a trend in B. subtilis. Indeed, transcripts present in both conditions showed a positive correlation between average DMS reactivity and abundance when each condition was analyzed separately (Fig. 3). This is distinct from the inverse correlation between the change in DMS reactivity and the change in abundance. The reactivity and coverage for each individual structurome (+AA, −AA) was also calculated separately. This analysis yielded additional transcripts, as mutual coverage between conditions was not considered, as well as less dispersed reactivity values for −AA conditions, as there was no cross 2%–8% normalization (i.e., normalizing −AA by +AA) (Supplemental Fig. S4; Supplemental Tables D.7, D.8). Similar trends show that in each separate structurome, the same relationship holds, where more reactive transcripts are typically more abundant (Supplemental Fig. S4). The trend is slightly stronger in −AA conditions (r = 0.43 vs. r = 0.67, Fisher Z-test P = 0.0000), again pointing to more overall modulation of structure in stress conditions. Additional details can be found in Supplemental Table T.4. We also plotted each of the transcripts as a straight-line arrow to show the shift that occurred when changing from growth in the presence of amino acids to growth in their absence (Fig. 3B). The majority of changes reciprocated each other: An increase of DMS reactivity corresponded with a decrease in abundance (red, sloping downwards), while a decrease in DMS reactivity was associated with an increase in abundance (blue, sloping upwards). This was highlighted by plotting just the 5% largest increases and decreases in abundance (Fig. 3C).

The stringent response and CodY regulons show coordinated changes in abundance and DMS reactivity in response to amino acid availability

Bacterial survival depends upon the capacity to rapidly alter metabolism, physiology, and behavior in response to changing environmental conditions. These responses involve changes in the expression of numerous genes. To coordinate these responses, bacteria utilize global regulatory networks through the action of a limited repertoire of regulatory factors (Eymann et al. 2002; Sonenshein 2005; Fujita 2009; Romeo et al. 2013; Cavanagh and Wassarman 2014; Geiger and Wolz 2014; Kovacs 2016). As RNA abundance is a common way in which cells regulate metabolic pathways in response to certain signals, we sought to determine if a change in RNA accessibility
also related to metabolic control. To determine how these RNA structural trends may affect physiology, we considered regulons, which are defined as sets of genes that are controlled by the same signal or regulatory molecule. We sought to uncover whether RNA structural features may contribute to coordination of these regulons. After assessing 174 known regulons in *B. subtilis* for a relationship between the changes in DMS reactivity and abundance as obtained through Structure-seq2 (Supplemental Item 1), we found that transcripts encoded by genes in both the stringent response and the CodY regulons experienced pronounced coordinated changes in abundance and DMS reactivity in response to amino acid availability (Fig. 4).

The stringent response regulon is the set of genes that are positively or negatively regulated in response to the alarmone (p)ppGpp following nutritional stress, including amino acid starvation (Eymann et al. 2002). Transcripts corresponding to genes that are known to be positively regulated by the stringent response showed a decrease in DMS reactivity and we confirmed their previously reported increased abundance (Eymann et al. 2002) when grown in the absence of amino acids, while genes known to be negatively regulated showed an increase in DMS reactivity and we confirmed their previously reported decreased abundance (Eymann et al. 2002) when grown in the absence of amino acids. We repeated our reactivity-to-abundance correlations excluding either or both of these regulons and found little change in the overall pattern (Supplemental Table T.4). The other 172 regulons analyzed did not show this strong correlation between known levels of regulation and DMS reactivity changes, which underscores the specificity of the response to the stringent response and CodY regulons.

**Structure-seq2 identifies known riboswitches during changes to amino acid availability**

The global RNA structural changes that we observed indicate that several mechanisms occur within the cell in response to amino acid availability. RNA switches are common mechanisms in which RNA structural features switch conformation following binding of another molecule. This switch then affects downstream gene expression. There have been numerous biochemical and molecular genetic studies characterizing various classes of RNA switches (Sherwood and Henkin 2016; McCown et al. 2017). Each of these experimental studies has focused predominantly on one class of RNA switch at a time. Currently, there are three known classes of direct amino acid-binding RNA switches: the lysine (Grundy et al. 2016)
2003; Sudarsan et al. 2003), glycine (Mandal et al. 2004), and glutamine (Ames and Breaker 2011) riboswitches. Additionally, amino acids are sensed indirectly through the SAM riboswitches that sense methionine levels (McDaniel et al. 2003), the T-box elements that sense the amount of uncharged tRNAs (Green et al. 2010), and protein-responsive elements such as the tryptophan-responsive TRAP binding elements (Babitzke and Gollnick 2001). These well-characterized RNA switches provide a starting point to benchmark Structure-seq2 and thereby hint that there may be additional amino acid-responsive RNA switches, especially given that there are 20 common amino acids and a large number of intermediates in the cell.

We initially focused on 31 RNA switches that are known to respond to amino acids in B. subtilis (Supplemental Table L.3). The 30 RNA switches able to be resolved in StructureFold included the lysine and glycine riboswitches that directly bind amino acids (Grundy et al. 2003; Sudarsan et al. 2003; Mandal et al. 2004), T-box elements that monitor the charged state of tRNAs (Green et al. 2010), S-box riboswitches that bind S-adenosylmethionine (McDaniel et al. 2003), and the protein-mediated RNA switches for tryptophan (Gollnick et al. 2005). The overall change in mean DMS reactivity of the 5′ UTRs of the 30 targeted transcripts is not significant (Fig. 5A, P = 0.71). This is expected since ligand binding to riboswitches generally involves a trading of base pairs rather than any large net loss or gain. This issue is examined in detail below. While these 5′ UTRs do change mean DMS reactivity between conditions, this overall change may not mirror or be indicative of changes in the switches and functional elements. The average DMS reactivity decreased in the CDS for these 30 targeted transcripts upon amino acid starvation, as seen with all transcripts (Figs. 1C, 5B, P = 3.25 × 10⁻³).

We plotted the change in DMS reactivity after amino acid starvation onto known secondary structures and crystal structures of some of the characterized RNA switches, including the T-box, S-box, glycine, and lysine riboswitches (Fig. 6; Supplemental Fig. S5; Supplemental Table L.4). Structure-seq2 faithfully reported on nucleotides known to be protected by the ligand, and the data aligned well with previously reported in vitro probing data in which changes in RNA structure were observed depending on the absence or presence of bound ligand.

FIGURE 4. Transcript DMS reactivity and abundance changes in the stringent response and CodY regulons are controlled by RNA accessibility. (A) Transcripts previously reported to be positively (teal) and negatively (red) regulated by the stringent response were subset from the data and plotted, comparing the change (Δ) in reactivity against the change (Δ) in abundance as per Figure 2. Abundance changes from our work clearly parallel reported abundance changes; the majority of positively regulated transcripts increased in abundance in −AA media (quadrant II). Likewise, the majority of negatively regulated transcripts decreased in abundance (quadrant IV). (B) Transcript names of positively regulated transcripts from quadrant II and (C) negatively regulated transcripts from quadrant IV. (D–F) Same analysis but for the CodY regulon, where repressed transcripts are also found in quadrant II and activated transcripts in quadrant IV. A full table summarizing these data is included (Supplemental Tables L.1, L.2).
The T-box element is a structurally responsive element that binds uncharged tRNAs and is often found in the 5′ UTR of genes encoding tRNA synthetases that charge the tRNA with the cognate amino acid. When uncharged tRNA binds to the nascent transcript it stabilizes the anti-terminator conformation in the 5′ UTR such that transcription continues into the downstream coding sequence, and the resultant increase in tRNA synthetase promotes the charging of tRNA molecules (Grundy and Henkin 2004; Sherwood and Henkin 2016). In our system, the absence of amino acids would drive tRNA to bind to the T-box riboswitch, presumably leading to protection from DMS reactivity. While there are multiple T-box riboswitches found within B. subtilis, the published secondary structure of the glyQS T-box was used to demonstrate structural changes associated with T-box functionality (Caserta et al. 2015). Although most of the nucleotides participating in the structure of the glyQS T-box did not show altered DMS reactivity values (Fig. 6A; Caserta et al. 2015), there are key residues that were altered in the absence of amino acids. For instance, when plotting the change in Structure-seq2 DMS reactivity on the part of the Geobacillus kaustophilus glyQS T-box that has a solved crystal structure (Grigg and Ke 2013), one nucleotide in the specifier loop, A93, is less reactive in the absence of amino acids, likely because it is protected by bound tRNA (Fig. 6B). When bound with tRNA, this nucleotide stacks on the wobble position of the anticodon and forms a hydrogen bond with a separate residue (Zhang and Ferre-D’Amare 2013). Additionally, A89 is known to form a sheared AA base pair that is near the tRNA that could account for the lower reactivity when tRNA binds. This binding promotes the formation of a pocket to accommodate tRNA modifications and could also explain the lower reactivity of nearby A87 when tRNA binds in the absence of amino acids (Zhang and Ferre-D’Amare 2013). Our observed reactivity changes imply that charged tRNA does not bind tightly to the T-box leader.

The S-box, or SAM riboswitch, binds S-adenosylmethionine, which contains methionine, and promotes termination (Grundy and Henkin 1998). There were a substantial number of nucleotides within the SAM riboswitch of the yitJ mRNA in B. subtilis that changed DMS reactivity in the absence of amino acids (Fig. 6C; McDaniel et al. 2003). When aligning published in vitro probing data with our in vivo Structure-seq2 data, nucleotides that had data in both data sets were in good agreement (Winkler et al. 2003). For instance, A24, A34, A37, A52, A111, and A113, which are ligand-protected in vitro, had increased DMS reactivity in the absence of amino acids in vivo (Fig. 6C; note overlap of red lettering with yellow shading). This increased DMS reactivity suggests that the RNA is bound by SAM in the presence of amino acids and unbound in their absence. According to the crystal structure of the B. subtilis yitJ SAM riboswitch (Lu et al. 2010), two key residues that participate in recognizing the SAM moiety, C73 and A74, were exposed in the absence of amino acids in our in vivo Structure-seq2 data (Fig. 6D). Other SAM-protected nucleotides in the in vitro data set showed no significant change in DMS reactivity in vivo, with the exception of A120, which showed decreased DMS reactivity without amino acids in vivo; these discrepancies could reflect the influence of full flanking sequence present in vivo, cellular factors, or differences in probing reactivity (OH− for in-line probing vs. DMS in Structure-seq2). While the T-box and S-box riboswitches monitor the availability of amino acids indirectly, the glycine and lysine riboswitches bind amino acids directly. Similar analyses were performed on these riboswitches, which supported our Structure-seq2 data by faithfully monitoring loss of amino acid binding upon starvation (Supplemental Fig. S5).

To identify regions of significant structure change more broadly, we used StructureFold2 (Tack et al. 2018) to perform a sliding window analysis throughout the 5′ UTRs of the 30 transcripts known to contain amino acid responsive RNA switches, looking at regions with increased DMS reactivity in the absence of amino acids (Supplemental Table L.5). The top 25% of windows with increased DMS reactivity in the absence of amino acids revealed overrepresented sequences corresponding closely to conserved regions within the S-box riboswitches (Fig. 6E,F). One region of the RNA, which is in a dashed boxed in Figure
FIGURE 6. Structure-seq2 data align well with known riboswitches and reveal functionally significant regions. (A) Change in DMS reactivity displayed on the secondary structure of the B. subtilis glyQS T-box element. Blue shows nucleotides that had decreased DMS reactivity without amino acids (darker shade shows more extensive change of DMS reactivity), green shows nucleotides that did not change in DMS reactivity, and yellow shows nucleotides that had increased DMS reactivity without amino acids. Secondary structure model adapted from Caserta et al. (2015). (B) Change in DMS reactivity displayed on the crystal structure of the G. kaustophilus glyQS T-box (Grigg and Ke 2013). Blue and yellow are nucleotides that had decreased or increased DMS reactivity without amino acids, respectively, and the tRNA ligand is displayed in forest green. PDBID: 4MGN. (C) Change in DMS reactivity displayed on the secondary structure of the B. subtilis yitJ SAM riboswitch. Color scheme is the same as in A, and nucleotides reported to have decreased DMS reactivity with bound SAM in a previous in vitro study are in red font (Winkler et al. 2003). (D) Change in DMS reactivity displayed on the crystal structure of the yitJ SAM riboswitch (Lu et al. 2010). Blue and yellow are nucleotides that had decreased or increased DMS reactivity without amino acids, respectively, and the SAM ligand is displayed in forest green. PDBID: 4KQY. (E,F) Highly overrepresented sequences that increased in DMS reactivity without amino acids from sliding windows of the 5′UTRs of the 30 resolved transcripts. (E) This region corresponds to the anti-antiterminator which would open up upon removal of SAM (dashed box in C). (F) This region corresponds to the 3′-end of the terminator, which would become exposed upon removal of SAM (black box in C).
6C with MEME (Bailey et al. 2009) in Figure 6E, is known to become partially sequestered in an anti-antiterminator structure when SAM binds and would be released in the absence of SAM binding. In addition, a second region of the RNA, which is in a solid boxed region in Figure 6C with MEME in Figure 6F, makes up the extreme 3’ end of the terminator hairpin that would become released in the absence of SAM binding. These sliding windows thus revealed important functional regions within RNA switches.

**Discovery of a transcription attenuation mechanism via Structure-seq2**

In addition to further characterization of known amino acid responsive RNA switches, we sought to use our Structure-seq2 data to discover new RNA switches. Computational analyses to discover classes of RNA switches have been based primarily on conservation of sequence, structure, and/or gene function of neighboring genes (Weinberg et al. 2017). This analysis pipeline has been modified to uncover increasingly rare RNA switches (Weinberg et al. 2017), but there are limitations to these methods. For instance, RNA switches belonging to just one type of bacteria or without a conserved sequence or known regulatory mechanism may be missed using this methodology. In addition, sequences conserved across bacteria may not function identically in all species. Thus, there is the potential that additional novel RNA switches exist, especially considering the vast number of molecules within a cell (Stav et al. 2019). Since known RNA switches have been found predominately within the 5’UTR of transcripts, we chose to first perform a search for new candidates within the 5’UTRs. We analyzed the 5’UTRs plus the first 50 nucleotides of the coding sequence from the Structure-seq2 reactivity files. We used the sliding window module from StructureFold2 (Tack et al. 2018) to analyze the top 5% of DMS reactivity changes within 20 nt windows, sliding by 10 nt intervals, in an effort to identify those that have the greatest change in DMS reactivity between our two growth conditions (i.e., ±amino acids). Many of the windows generated corresponded to regions of genes with unknown function (Supplemental Table I.6). One such gene, yfmG, contains a region with homology to the formylglycine-generating enzyme domain (Caspi et al. 2014). Members of this family convert cysteine to formylglycine posttranslationally (Fig. 7A), which plays a role in sulfate scavenging (Carlson et al. 2008). Within this 5’UTR a NusA-dependent intrinsic transcription terminator was identified previously by Term-seq when cultures were grown in minimal medium (Fig. 7B,C; Mondal et al. 2016), but not when grown in LB (Dar et al. 2016). There were large changes in DMS reactivity upon amino acid starvation (Fig. 7D), with the largest change upstream of this terminator. The structures predicted using in vivo constraints in RNAStructure (Reuter and Mathews 2010) were predicted to have more base-pairing interactions in the presence of amino acids (Fig. 7B,C).

The yfmG transcription start site was determined prior to conducting regulatory studies. Total cellular RNA was extracted from mid-exponential phase cultures grown under the same conditions used for Structure-seq2. Primer extension analysis led to the identification of a single yfmG transcription start site 201 nt upstream of the yfmG translation initiation codon (Fig. 8A,B), which allowed us to identify a $A^\delta$-dependent promoter with an extended −10 element (Fig. 8A). The long 201-nt leader suggested that yfmG expression could be controlled posttranscriptionally. Structural predictions based on DMS reactivity provide an average structure of the full-length leader and do not take the kinetics of transcription into consideration. Therefore, we also performed RNA structure predictions using Mfold (Zuker 2003) on 3’-shortened fragments that do not include the entire terminator to search for a potential competing structure. In addition to the previously identified terminator, a putative antiterminator (AT) structure was identified that overlapped the terminator hairpin. In addition, a potential anti-antiterminator (AAT) structure was identified even further upstream that overlapped with the AT structure. These structures suggested a model in which formation of the AAT structure would prevent formation of the AT structure, leading to termination upstream of the yfmG coding sequence, whereas formation of the AT would lead to transcription readthrough (Fig. 8C). Furthermore, inspection of the sequence between the transcription start site and the yfmG translation initiation codon led to the identification of a putative 23 amino acid coding sequence (Fig. 8A). This putative minigene was previously annotated as a protein coding sequence (yfmH) (Zuber 2001) and as a sRNA (Irnov et al. 2010). Note that the sRNA is derived from termination at the NusA-dependent terminator (Mondal et al. 2016).

A chromosomally integrated PyfmHG-yfmH-lacZ transcriptional fusion was generated such that the fusion junction was just downstream from the yfmH transcription termination site (+175 relative to the start of transcription) (Fusion 1, Fig. 9A). High-level expression was observed for this fusion when grown in the absence of amino acids, whereas expression was about eightfold lower when grown in the presence of amino acids (Fusion 1, Fig. 9B). These results indicate that yfmHG expression is greatly reduced when grown in the presence of amino acids. We next tested a PyfmHG-lacZ transcriptional fusion in which all but the first 5 nt of the yfmHG leader region were deleted (Fusion 2, Fig. 9A). In this case, expression was reduced just 1.5-fold when grown in the presence of amino acids (Fusion 2, Fig. 9B), indicating that amino acids had only a small effect on transcription initiation from the yfmHG promoter. A third fusion was generated in which the constitutive trp operon promoter was used to replace the yfmHG promoter (Fusion 3, Fig. 9A). Expression of this fusion
was still strongly reduced (fivefold) when grown in the presence of amino acids (Fusion 3, Fig. 9B). We conclude that amino acid starvation leads to a substantial increase in expression of the yfmHG operon and that the long yfmH leader but not the promoter is responsible. We further infer that amino acid starvation leads to readthrough past the terminator separating yfmH and yfmG (Fig. 8).

A PyfmHG-yfmH′-lacZ translational fusion was generated to determine whether yfmH was actually translated (Fusion 4, Fig. 9A). The junction of this fusion was just after the 23rd codon (i.e., the yfmH stop codon was replaced with the lacZ coding sequence). This fusion was expressed in the absence and presence of amino acids (Fusion 4, Fig. 9C), whereas replacing the AUG start codon of yfmH with a UAG stop codon eliminated expression (Fusion 4*, Fig. 9C). These results establish that yfmH is translated and likely encodes a leader peptide that regulates yfmG expression. We next determined the effect of yfmH translation on expression of PyfmHG-yfmH-lacZ transcriptional and PyfmHG-yfmH′-lacZ translational fusions (Fusions 1 and 5, Fig. 9A). Introduction of the yfmH start to stop codon mutation greatly reduced expression of the transcriptional fusion; however, expression of this fusion still responded to amino acids (Fusion 1*, Fig. 9C). Although expression of the PyfmHG-yfmH′-lacZ translational fusion was low, its expression was reduced 2.5-fold upon addition of amino acids (Fusion 5, Fig. 9C). Importantly, introduction of the yfmH start-to-stop codon mutation resulted in threefold lower expression and eliminated the effect of amino acids on expression (Fusion 5*, Fig. 9C).

One possibility that could explain this expression data is that the rate of yfmH translation would dictate which of the alternative structures form upstream of yfmG. Since expression was high in the absence of amino acids, we postulated that a slow rate of yfmH translation would favor formation of the AT structure by interfering with formation of the AAT structure (Fig. 8C). Inspection of the yfmH coding sequence indicated that it contains codons specifying 13 different amino acids (Fig. 8A). We also noticed that there are three arginine-lysine (RK) motifs in the first half of YfmG, as well as one K further downstream. We found that growth in minimal medium plus just these two amino acids resulted in a 2.5-fold reduction in expression of the PyfmHG-yfmH-lacZ transcriptional fusion (Fig. 9D). In addition, growth in minimal medium plus isoleucine, leucine and valine (ILV), which corresponds to five YfmH codons,
resulted in a threefold decrease in expression. Finally, when we grew cultures in minimal medium containing all 13 amino acids encoded in YfmH we found that expression of the \textit{PyfmHG}–\textit{yfmH}–\textit{lacZ} transcriptional fusion was almost as low as when this strain was grown with all amino acids (Fig. 9D). These results are consistent with a model in which a slow rate of YfmH translation favors formation of the AT structure, leading to transcriptional readthrough into the \textit{yfmG} coding sequence. Conversely, a fast rate of YfmH translation would interfere with AT formation, leading to transcription termination upstream of \textit{yfmG}. This novel type of RNA switch offers a way to simultaneously monitor the levels of several amino acids at once.

**DISCUSSION**

RNA structure in bacterial cells plays important and diverse roles on both global and gene-specific levels. Using Structure-seq2 to probe the in vivo RNA structure genome-wide in \textit{B. subtilis} in the presence and absence of amino acids, we saw that there is transcriptome-wide reduced DMS reactivity of RNA in response to amino acid starvation. Similarly, changes in the RNA structureome were observed in eukaryotic species where both heat stress and salinity stress promoted reduced DMS reactivity of RNA (Su et al. 2018; Tack et al. 2020). These findings suggest that there may be a universal stress response mechanism throughout the different domains of life that is manifested in changes in RNA accessibility. One possible explanation for reduced DMS reactivity in the CDS is that translating ribosomes may move more slowly during amino acid starvation, which would passively shield the RNA from DMS. Another explanation is that reduced rates of translation may result in an increase in the distance between translating ribosomes, resulting in the formation of RNA structures that would not have the chance to form when translation is efficient under amino acid replete conditions.

**FIGURE 8.** \textit{yfmHG} promoter and leader region. (A) Sequence of the \textit{yfmHG} promoter and leader region. The −35 and extended (ext) −10 promoter elements and the transcription start site (+1) are in bold. The \textit{yfmH} Shine-Dalgarno (SD) sequence and the \textit{YfmH} protein-coding sequence including start (ATG) and stop (TAA) codons are shown. The \textit{yfmG} SD sequence and TTG start codon are also shown. The anti-antiterminator (AAT), antiterminator (AT), and terminator (T) hairpins are indicated by pairs of inverted arrows. The termination site (term) identified in vivo by Term-seq is also marked (Mondal et al. 2016). The overlap between the AAT and AT structures are shown in cyan, while the overlap between the AT and T structures are in red. Numbering is with respect to the start of transcription. (B) Primer extension mapping of the \textit{yfmHG} transcription start site. Primer extension analysis was performed on total cellular RNA extracted from a wild-type strain of \textit{B. subtilis} grown in minimal + glutamine medium (M) and minimal medium containing all amino acids (AA). Sequencing lanes (A, C, G, T) are shown. (C) Transcription attenuation model. Predicted structures of the anti-antiterminator and terminator structures (left) or the antiterminator structure (right) are shown. Overlaps and color coding between the competing structures are as described in A. The \textit{yfmH} SD sequence, start (met) and stop codons are indicated. Numbering is with respect to the start of transcription.
When relating DMS reactivity changes with abundance changes following amino acid starvation, we saw an inverse correlation among all transcripts in which higher average DMS reactivity in the absence of amino acids corresponded to a decrease in transcript abundance. This inverse correlation found in eukaryotes (rice, Arabidopsis) (Su et al. 2018; Tack et al. 2020) and now in bacteria (B. subtilis) suggests that the regulation of RNA abundance by adjusting transcript accessibility may be a deeply conserved characteristic of all life. Beyond this, the inverse relationship was observed when comparing whole transcripts as well as the CDS, but not when comparing the 5′ UTR and 3′ UTR regions (Fig. 2B, D). These data suggest that 5′ UTRs, which do not contain translating ribosomes nor sequence constraints due to coding, have a wider range of structural change, supporting the notion that most RNA structural changes that regulate gene expression occur within 5′ UTRs.

Within B. subtilis, the inverse relationship between DMS reactivity and abundance under amino acid starvation conditions is notably pronounced in the stringent response and CodY regulons, both of which are known to be involved in regulating gene expression in response to amino acid starvation. As other regulons did not show this dramatic correlation, these results suggest that only certain transcripts are susceptible to the inverse relationship under these specific conditions, which may provide an advantage to B. subtilis under amino acid starvation conditions. We hypothesize that a change in amino acid availability causes direct and indirect effects on the structure of the transcripts within the stringent response and CodY regulons that then causes a corresponding change in abundance. This pattern is likely due to a mixture of mechanisms. For example, during the stringent response, protection from DMS reactivity caused by increased translation would lead to increased transcript abundance by protecting the mRNAs from nucleolytic digestion (Fig. 4A, B, teal).

Importantly, we demonstrate that genome-wide RNA structural data are able to report on, as well as predict, specific RNA switches that control gene expression. We showed that under two different conditions of B. subtilis growth, Structure-seq2 can accurately characterize RNA switches known to respond to amino acids. We found that certain nucleotides show differences in reactivities between prior in vitro studies and this in vivo analysis (Fig. 6; Supplemental Fig. 5). These differences likely arise from the complex cellular environment that is absent from in vitro conditions. Our study presents changes in DMS reactivity between two conditions, which is interpreted as changes in RNA structure. Further work should be done to precisely identify the native in vivo folds and the other factors affecting these folds. While single-gene in vitro characterization can provide detailed mechanistic information on a single riboswitch, in vivo genome-wide studies provide a robust tool to analyze the structure and function of numerous RNA switches in a single study, to do so in the more relevant condition of the cell, and to identify new candidate RNA switches.

By using DMS reactivity values generated through Structure-seq2, we were able to discover novel RNA features through these high-throughput sequencing data. Particularly, we identified a transcription attenuation mechanism in the yfmG leader region by analyzing regions with high DMS reactivity changes in response to amino acids within the RNA structurome. Within this region, the mRNA encoding a short leader peptide overlaps AAT...
and AT hairpins, and the absence of amino acids found within this peptide promotes transcription readthrough into yfmG. These findings suggest that yfmG, a gene with unknown function, may play a role in survival under amino acid starvation conditions. Additional studies are required to elucidate the function of yfmHG, and how it relates to starvation for the 13 amino acids encoded in yfmH. Uncovering this mechanism has broad implications—it suggests that B. subtilis can use a combination of RNA structure and short peptides to sense multiple amino acids. In conclusion, our study demonstrates the utility of using the in vivo genome-wide data generated from Structure-seq2 to discover novel RNA-based regulatory mechanisms.

MATERIALS AND METHODS

Bacterial strains, plasmids, and oligonucleotides

B. subtilis strains used in this study are listed in Table 1. Plasmids used in this study are described in Supplemental Table M.1 (Grandoni et al. 1993; Merino et al. 1995) and DNA oligonucleotides in Supplemental Table M.2.

Primer extension assay

Total RNA was isolated from late-exponential phase cultures of B. subtilis PLBS338 grown in minimal + glutamine (−AA) medium (1× Spizizen salts, 0.5% glucose, 0.04% L-glutamine) and ACH + tryptophan (+AA) medium (1× Spizizen salts, 0.5% glucose, 0.2% acid casein hydrolysate, 0.01% L-tryptophan) using the RNeasy Mini kit (Qiagen). Ten µg of total RNA was hybridized to 150 nM 32P-5′-end-labeled DNA oligonucleotide complementary to nucleotides +94 to +109 (relative to yfmHG transcription) for 3 min at 80°C. Reaction mixtures (6 µL) containing 2 µL of hybridization mixture, 375 µM of each dNTP, 10 mM dithiothreitol (DTT), 1× SuperScript III buffer, and 0.25 µL SuperScript III Reverse Transcriptase (Life Technologies) were incubated for 30 min at 37°C in a shaking water bath. These conditions provide appropriate single-hit kinetics, which is generally defined as a 10%–20% decrease in the full-length transcript-specific reverse transcription band (Supplemental Fig. S1). After 5 min, 1 g DTT was added to the culture to quench the reaction for 2 min in

β-galactosidase assay

B. subtilis cultures containing various transcriptional or translational fusions were grown at 37°C in −AA medium supplemented with 5 µg/mL chloramphenicol or in +AA medium supplemented with 5 µg/mL chloramphenicol. When indicated, −AA growth media also contained 0.004% alanine, 0.02% arginine, 0.004% asparagine, 0.004% aspartic acid, 0.002% cysteine, 0.002% glutamic acid, 0.01% glycine, 0.01% histidine, 0.01% isoleucine, 0.01% leucine, 0.008% lysine, 0.007% methionine, 0.003% phenylalanine, 0.02% proline, 0.01% serine, 0.01% threonine, 0.003% tyrosine, 0.02% valine and/or 0.01% L-tryptophan. Cells were grown to mid-exponential phase and β-galactosidase activity was determined as described previously (Du and Babitzke 1998).  

Bacterial growth and DMS treatment

DMS treatments were performed in a chemical fume hood. B. subtilis strain PLBS338 (Yakhnin et al. 2004) was grown at 37°C in +AA or −AA media. For each replicate (three per growth media), cultures were grown to mid-exponential phase prior to DMS treatment. For the −AA control, 6 mL of culture was transferred to a tube containing 0.45 g DTT and slightly agitated for 2 min. The solution was then poured into an equal volume of frozen slurry killing buffer (10 mM Tris-HCl, pH 7.2, 5 mM MgCl2, 25 mM NaN3, 1.5 mM chloramphenicol, and 12.5% ethanol). DMS was added to the remaining 14 mL of culture to a final concentration of 50 mM and the reaction was allowed to proceed for 5 min in a 37°C in a shaking water bath. These conditions provide appropriate single-hit kinetics, which is generally defined as a 10%–20% decrease in the full-length transcript-specific reverse transcription band (Supplemental Fig. S1). After 5 min, 1 g DTT was added to the culture to quench the reaction for 2 min in

### Table 1. B. subtilis strains used in this study

| Strain     | Genotype* | Source                |
|------------|-----------|-----------------------|
| PLBS338    | Prototroph| Yakhnin et al. 2004   |
| PLBS864    | amyE::PyfmHG-yfmH-lacZ Cmr’ (−253 to +180)*i | This study |
| PLBS979    | amyE::PyfmHG-yfmH’-lacZ Cmr’ (−253 to +132) | This study |
| PLBS984    | amyE::PyfmHG-lacZ Cmr’ (−253 to +5) | This study |
| PLBS985    | amyE::Ptryp-yfmH-lacZ Cmr’ (+1 to +180) | This study |
| PLBS986    | amyE::PyfmHG-yfmHG’-lacZ Cmr’ (−253 to +216) | This study |
| PLBS995    | amyE::PyfmHG-yfmHG’-lacZ Cmr’ (−253 to +216, A64T:T65A) | This study |
| PLBS996    | amyE::PyfmHG-yfmH’-lacZ Cmr’ (−253 to +180, A64T:T65A) | This study |
| PLBS1000   | amyE::PyfmHG-yfmH-lacZ Cmr’ (−253 to +180) | This study |

*aNumbers in parentheses indicate the cloned yf8 region relative to the start of transcription, as well as y8 leader mutations. Cmr, chloramphenicol.

*iNumbers in parentheses indicate yfmHG DNA in the fusion relative to the start of transcription, as well as nucleotide substitutions converting the yfmH start codon to a stop codon.
the shaking water bath. An amount of 6 mL of the quenched culture was then added to frozen slurry killing buffer. Culture tubes were centrifuged for 10 min at 10,000 rpm. Cell pellets were suspended in 1 mL ice cold killing buffer, and then cells were pelletted in a microfuge tube. The wash procedure was repeated and then cell pellets were stored at −20°C.

**Structure-seq2 library preparation**

To obtain RNA for Structure-seq2 library preparation, total RNA from the cell pellets was extracted using the RNeasy Mini Kit (Qiagen). After extraction, the quality of RNA was assessed by observing rRNA bands following fractionation through a 1% agarose gel, as well as by a Prokaryote Total RNA Nano Bioanalyzer scan to identify rRNA peaks. rRNA was then depleted using the Ribo-Zero rRNA Removal Kit for Gram-positive bacteria (Illumina), and the yield and quality were assessed via spectrophotometry and by an mRNA Pico Bioanalyzer scan to confirm depletion of the rRNA peaks.

Following rRNA depletion, RNA was either sent for RNA-seq analysis or libraries were prepared following the biotin variation of Structure-seq2, as described previously (Ritchey et al. 2019). Briefly for Structure-seq2 preparation, using ~300 ng RNA-depleted RNA, cDNA was synthesized using a random hexamer to prime reverse transcription in the presence of 500 µM each dNTP, 62.5 µM biotinylated-dCTP, and 62.5 µM biotinylated-dUTP. RT stops immediately before the site of DMS modification, allowing these modifications to be read out transcriptome-wide. The reactions were purified via RNA Clean & Concentrator-5 (Zymo Research) and streptavidin pull-down. A hairpin adaptor was then ligated to the 3′ end of the cDNA fragments, and products were purified via RNA Clean & Concentrator-5 and streptavidin pull down. A polymerase chain reaction (PCR) cycle test was performed to determine the appropriate number of cycles for amplification and 18 cycles was chosen. PCR was performed with sequences complementary to Illumina sequencing primers incorporated into the primers. The resulting libraries were fractionated through a denaturing 8.3 M urea 10% polyacrylamide gel, purified via a crush and soak method to select for sizes 200–600 nt, and submitted for sequencing. Prior to sequencing, the quality of libraries was assessed via Bioanalyzer scans.

**Transcriptome assembly**

To maximize the amount of resolvable sequence space containing potential structural regulatory elements (i.e., 5′- and 3′-UTRs), we generated our own reference transcriptome. Standard RNA-seq libraries were generated from both +AA and −AA conditions, yielding 39,310,106 and 42,864,834 reads, respectively. These libraries were trimmed with trimmomatic, version 0.36 (Bolger et al. 2014) yielding 39,280,081 +AA and 41,830,095 −AA reads for assembly. Trimmed reads were used as input for Rockhopper, version 2.03 (Tjaden 2015), which aligned 38,256,615 and 40,582,875 reads on the 8. subtilis strain 168 genome for a combined assembly containing 2369 5′UTR regions and 2119 3′UTR regions. The identified boundaries of these transcripts were used to parse out the putative transcripts (https://github.com/StructureFold2/NotToosubtilis) into fasta format. The fasta file of this whole transcriptome is included in Supplemental Item S2.

To confirm the efficacy of our Structure-Seq2/StructureFold2 method in finding RNA switches, we manually generated a panel of the complete transcripts of known riboswitches, thus relying on annotation where it was available, rather than relying on a reference-based assembly as with the whole transcriptome. Sequences were found with BsubCyc (Caspi et al. 2014), and compiled manually into a fasta file. Where transcription start sites were not noted in BsubCyc, they were estimated based on the beginning of the annotated regulatory features (e.g. T-box upstream of the gene. The fasta file of this targeted transcriptome is included in Supplemental Item S2.

**Reactivity analyses**

Structure-Seq2 libraries were first trimmed via Cutadapt, version 1.16 (Martin 2011) via the fastq_trimmer module of StructureFold2 (Supplemental Table W.1; Tack et al. 2018). These trimmed reads were aligned to both the whole transcriptome (for all genome-wide analyses) and to the panel of known riboswitches (for targeted analyses) via Bowtie2, version 2.3.4.1 (Langmead and Salzberg 2012) via the fastq_mapper module of StructureFold2 (Tack et al. 2018) (Supplemental Table W.2.a,b). Both sets of mappings were filtered to remove reads with a mismatch on the first base or reads with more than three mismatches with the StructureFold2 sam_filter module (Supplemental Table W.3.a,b; Tack et al. 2018) which utilizes SAMtools, version 0.1.19 (Li et al. 2009).

Filtered mappings were converted to .rtsc (reverse transcriptase stop count) files, replicates were merged, and per transcript coverage was calculated via StructureFold2 (Tack et al. 2018). For transcripts above the coverage threshold of 1 in the merged file (amounting to ≥1 RT stop per A or C residue in the transcript) (Ding et al. 2014; Su et al. 2018; Tack et al. 2020), we correlated raw RT stop patterns between component replicates (Supplemental Fig. S3; Supplemental Table T.1) with R (R Core Team 2019) using the plyr package (Wickham 2011) via the rtsc_correlation module of StructureFold2 (Tack et al. 2018); this follows the same process for evaluating library replicate consistency as was done previously (Su et al. 2018; Waldron et al. 2019; Tack et al. 2020). Merged rtsc files were then used to calculate per-base reactivity via the rtsc_to_react module of StructureFold2 (Tack et al. 2018), yielding react files.

React files for the whole transcriptome analyses were analyzed with the StructureFold2 (Tack et al. 2018) react_statistics module to obtain overall transcript reactivity statistics. To estimate transcript abundance, the corresponding +AA/−AA rtsc files were used to calculate transcripts per million reads (TPM) (Wagner et al. 2012) via the StructureFold2 (Tack et al. 2018) rtsc_abundance module. The DMS reactivity statistics and abundances were analyzed in R (R Core Team 2019) and plotted using the ggplot2 package (Wickham 2016). Fisher Z-tests between correlations were calculated with the cocor package (Diedenhofen and Musch 2015). For batch analysis comparing our data to known regulons, we integrated a list of regulons from SubtiWiki (Zhu and Stulke 2018) (http://www.subtiwiki.uni-goettingen.de/v3/exports). For the sliding window analyses, we used the react_windows module (Tack et al. 2018) to find areas of reactivity change between conditions. We queried by net change (delta) and searched for the top 5% of most changed 20 nt windows sliding by 10 nt within the 5′UTRs of all transcripts (Supplemental Table L.6).
The RT stop counts of the targeted switch transcript 5’UTRs were first subset from the entire targeted switch transcript counts, creating separate 5’UTR exclusive rtsc files. These rtsc files were then used to calculate per base reactivity on the 5’UTR in both +AA and −AA conditions, applying a 5’UTR specific 2%–8% scale from +AA to the −AA data, thus generating react files. These react files containing the final calculated reactivity values were then converted to .csv files (included in Supplemental Item S3) and manually analyzed for changes, or used as restraints to guide folding using RNAstructure (Reuter and Mathews 2010) via StructureFold2 (Tack et al. 2018). For the sliding window analysis on the targeted windows, we followed a similar procedure as above and queried for gains, losses, and net change. The fasta version of these results were submitted for MEME analysis using the online webserver with default parameters (Bailey et al. 2009). Only the top 25% of windows that gained reactivity produced significant motif enrichment (Fig. 6E, F; Supplemental Table L5). Structure-Seq2 and RNA-seq libraries were deposited in Gene Expression Omnibus (GEO) (accession number GSE148936).

Library quality analysis

We used StructureFold2 (Tack et al. 2018) to process and analyze the resulting Structure-seq2 libraries. The rtsc_correlation.py module was used to identify the RT stops prior to every A and C residue in our assembled transcriptome in .csv format for all +DMS libraries in both treatments (+AA, −AA). These two .csv files were read in and analyzed using R (R Core Team 2019) and the ggplot2 package (Wickham 2016). For every pairwise combination of two replicates from the same treatment, we treated the entire transcriptome as one entity and computed the correlation coefficient of the number of RT stops between every A and C: Values are included on each respective graph and in Supplemental Table T.1. Second, we subdivided the transcriptome by transcript and computed the correlation coefficient for each transcript between each pairwise combination of two replicates. The density of these distributions of r values were likewise plotted, showing the vast majority of per transcript correlation coefficients to be high (Supplemental Fig. S3; Supplemental Table T.1). These results implied that the replicates of both treatments were suitably similar to be pooled to minimize technical variation and provide more robust numbers before proceeding to calculate derived DMS reactivity.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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