RNA G-quadruplexes are globally unfolded in eukaryotic cells and depleted in bacteria

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INTRODUCTION: Many cellular RNAs contain regions that fold into stable structures required for function. Both Watson-Crick and noncanonical interactions can play important roles in forming these structures. An intriguing noncanonical structure is the RNA G-quadruplex (RG4), a four-stranded structure containing two or more layers of G-quartets, in which the Watson-Crick face of each of four G residues pairs to the Hoogsteen face of the neighboring G residues. RG4 regions can be very stable in vitro, particularly in the presence of K+, and thus they are generally assumed to be predominantly folded within cells, which have ample K+. Indeed, these structures have been implicated in mRNA processing and translation, with recently proposed canonical structure is the RNA G-quadruplex (RG4), a four-stranded structure containing two or more layers of G-quartets, in which the Watson-Crick face of each of four G residues pairs to the Hoogsteen face of the neighboring G residues. RG4 regions can be very stable in vitro, particularly in the presence of K+, and thus they are generally assumed to be predominantly folded within cells, which have ample K+. Indeed, these structures have been implicated in mRNA processing and translation, with recently proposed roles in cancer and other human diseases. However, the number of cellular RNAs that can fold into RG4 structures has been unclear, as has been the extent to which these RG4 regions are folded in cells.

RATIONAL: Enzymes and chemicals that act on RNA with structure-dependent preferences provide valuable tools for detecting and monitoring RNA folding. For example, dimethyl sulfate (DMS) treatment of RNA, either in vitro or in cells, coupled with high-throughput sequencing of abortive primer-extension products can monitor the folding states of many RNAs in one experiment. Analogous high-throughput methods use cell-permeable variants of SHAPE (selective 2′-hydroxyl acylation analyzed by primer extension) reagents. These methods reveal important differences between RNA structures formed in vivo and those formed in vitro. However, they are designed to detect Watson-Crick pairing and do not identify RG4 structures or provide information on their folding states. After recognizing that RG4 regions can block reverse transcriptase, we reasoned that this property, together with the known ability of RG4s to protect the N7 of participating G nucleotides from DMS modification, could be used to develop a suite of high-throughput methods to both identify endogenous RNAs that can fold into RG4s in vitro and determine whether these regions also fold in cells.

RESULTS: We first developed a high-throughput method that identifies RG4 regions on the basis of their propensity to stall reverse transcriptase in a K+-dependent manner. Applying this method to RNA from mammalian cell lines and yeast, we identified >10,000 endogenous RNA sequences that contain regions that form RG4s in vitro, thereby expanding by a factor of >100 the catalog of endogenous RNA regions that can fold into RG4 structure.

CONCLUSION: In mammals, thousands of endogenous RNA sequences have regions that can fold into RG4s in vitro, but these regions are globally unfolded in eukaryotic cells, presumably by robust and effective machinery that remains to be fully characterized. In contrast, RG4 regions are permitted to fold in E. coli cells, and the other bacteria have undergone evolutionary depletion of endogenous RG4-forming sequences.

The different mechanisms that eukaryotic and eubacterial cells use to avoid RG4 structures. RG4 regions are abundant but overwhelmingly unfolded in mammalian and yeast cells, implying robust and efficient molecular machinery, presumably involving helicases and RNA-binding proteins, which specifically unfolds RG4 regions and maintains them in an unfolded state (left). In contrast, regions that can fold into RG4 structures are depleted in bacteria, implying fewer progeny of cells that acquire these regions (right).
RNA G-quadruplexes are globally unfolded in eukaryotic cells and depleted in bacteria

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In vitro, some RNAs can form stable four-stranded structures known as G-quadruplexes. Although RNA G-quadruplexes have been implicated in posttranscriptional gene regulation and diseases, direct evidence for their formation in cells has been lacking. Here, we identified thousands of mammalian RNA regions that can fold into G-quadruplexes in vitro, but in contrast to previous assumptions, these regions were overwhelmingly unfolded in cells. Model RNA G-quadruplexes that were unfolded in eukaryotic cells were folded when ectopically expressed in mammalian cells. Model RNA G-quadruplexes that were unfolded in eukaryotic cells were folded when ectopically expressed in mammalian cells.

Mammalian RNAs contain regions that fold into stable structures required for function (<sup>1,2</sup>). These structures can be studied with chemical probes that modify accessible or flexible nucleotides (<sup>3-5</sup>). For example, dimethyl sulfate (DMS) methylates A and C residues that are not protected by Watson-Crick pairing or other interactions, and because these modifications stall reverse transcriptase, primer-extension reactions can detect modification and thereby report on the folding state of these nucleotides. DMS also penetrates living cells and modifies RNAs within these cells, and with high-throughput sequencing of global primer-extension products, the intracellular folding of numerous RNAs can be simultaneously monitored in a procedure called DMS-seq (<sup>6,7</sup>). Analogous high-throughput methods have also been developed with cell-permeable SHAPE (selective 2′-hydroxyl acylation analyzed by primer extension) reagents (<sup>8,9</sup>). These methods reveal important differences between RNA structures formed in vivo and those formed in vitro (<sup>7,9</sup>). However, these high-throughput methods are designed to detect Watson-Crick pairing, which leaves the folding states of noncanonical structures difficult to assess.

One such noncanonical structure is the RNA G-quadruplex (RG4), in which four strands of RNA interact, either intramolecularly or intermolecularly, through the formation of two or more layers of G-quartets, in which each of four G residues pairs to two neighboring G residues (Fig. 1A) (<sup>10,11</sup>). Owing to the extensive hydrogen-bonding and base-stacking interactions, RG4 structures can be very stable, with in vitro melting temperatures well exceeding physiological temperatures. This stability typically depends on the presence of K<sup>+</sup>, which is the optimal size to bind at the center of two stacked G-quartets and thereby counter the otherwise repulsive partial negative charges that converge at the quadruplex core (Fig. 1A).

Because of the high stability of RG4 structures in vitro and the high concentration of K<sup>+</sup> in cells (typically >100 mM, well above that required for quadruplex formation), regions that fold into RG4 structures in vitro are generally assumed to fold into these structures in cells. Indeed, RG4s are implicated in control of mRNA processing and translation, with recently proposed roles in human diseases, such as cancer (<sup>12</sup>) and neurodegeneration (<sup>13</sup>). Supporting the idea that RG4s are folded in cells, immunostaining with G4-specific antibodies yields a detectable, albeit weak fluorescence signal in vivo and in vitro in cell nuclei (<sup>14</sup>). However, these methods are indirect, and the data do not rule out the possibility that RG4s are present in cells but not in solution.

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Fig. 1. Strong RT stops at G-rich regions in the mESC transcriptome. (A) The RNA G-quadruplex. The schematic (top) depicts a three-tiered RG4, with a parallel RNA-backbone orientation (solid line) and three G-quartets stabilized by two K<sup>+</sup> ions (spheres). The chemical structure of a G-quartet (bottom) highlights hydrogen bonding (dashed lines) to the N7 positions (red) and K<sup>+</sup>-facilitated convergence of the exocyclic oxygens of the four G residues (R, ribose). (B) Schematic of RT-stop profiling. See text for explanation. (C) RT-stop profiles for Eef2 mRNA (box, coding sequence) and Malat1. Bars representing each RT stop are colored according to the identity of the template nucleotide at the 5′-terminal cDNA nucleotide assigned position +1. Template nucleotides are plotted at heights indicating the information content of their enrichment (bits).
weak, ribonuclease-sensitive signal in the cytoplasm (14). However, these immunostaining results leave open the possibility of folding during the processes of fixing, permeabilizing, or staining cells, and even if this signal represented quadruplex formation in cells, it could not provide information regarding either the sequence identities or the overall fraction of RG4 regions that fold in vivo.

**Many RNAs with quadruplex-forming capacity**

To systematically search for structure-forming potential in mammalian cellular RNAs, we exploited the ability of stable structures to stall reverse transcription. Polyadenylate (poly(A))-selected mRNAs from mouse embryonic stem cells (mESCs) were randomly fragmented, and 60- to 80-nucleotide (nt) fragments were ligated to a common 3' adapter used for global primer extension. Complementary DNAs (cDNAs) resulting from reverse transcription (RT) that stalled after only 20 to 45 nt of extension were purified and sequenced to identify the RT stops (Fig. 1B), by means of a procedure resembling that developed for DMS-seq (7). As illustrated for the Eef2 (Eukaryotic elongation factor 2) mRNA and the Malat1 (Metastasis-associated lung adenocarcinoma transcript 1) noncoding RNA, most of the strong RT stops (65%) were at G nucleotides (Fig. 1, A and D, and fig. S1; P < 10^{-15}, χ^2 test). Analysis of the flanking sequences of these strong RT stops at G nucleotides showed that the 30 nucleotides upstream of the RT stops were also enriched in G (and depleted in C), particularly at positions −1 and −2 (92 and 68% G, respectively; Fig. 1E). In contrast, no enrichment was detected downstream, except for weak G enrichment at position +1 (38% G; Fig. 1E).

The upstream G enrichment, together with recent studies of individual transcripts (25), suggested that formation of intramolecular RG4 structures caused these strong RT stops. To test this possibility, we examined whether these RT stops were sensitive to the identity of the monovalent counterion and found that substituting K+ in the RT reaction with either Na+ or Li+ greatly diminished the RT stops at G residues (Fig. 2, A and B, and fig. S2A). Another diagnostic feature of RG4s is their sensitivity to modification of the N7 position of G (15). Methylating this position by using DMS under denaturing conditions (95°C, 0 mM K+), or treating the RNA with DMS under denaturing conditions before RT (DMS 95°C). At each position within the mRNA, the fold enrichment was calculated as the number of RT-stop reads observed at that position, divided by the average number of RT-stop reads observed for all the mRNA positions with the same nucleotide identity. (B) Global analyses of strong RT stops, comparing the enrichment of stops observed in the original conditions (untreated; K+) to those observed when either substituting K+ with Na+ (top) or pretreating RNA with DMS at 95°C (bottom). RT-stop values are colored according to the template nucleotide at position 0. The distributions are truncated at the left because stops with enrichment by less than a factor of 20 in the untreated K+ sample were not classified as strong stops. (C) Overlap between K+-dependent and DMS-sensitive strong RT stops (yellow and blue, respectively). (D) Abundance of RG4 regions within mRNA translated and untranslabeled regions. The expected abundances were estimated on the basis of the relative number of G nucleotides within these three regions of detected mRNAs.

Fig. 2. Folded RG4 structures cause strong RT stops. (A) RT-stop profiles of Eef2, showing enrichment observed in the original conditions (K+) and that observed when either substituting the monovalent cation used during RT (Li+ or Na+) or treating the RNA with DMS under denaturing conditions before RT (DMS 95°C). At each position within the mRNA, the fold enrichment was calculated as the number of RT-stop reads observed at that position, divided by the average number of RT-stop reads observed for all the mRNA positions with the same nucleotide identity. (B) Global analyses of strong RT stops, comparing the enrichment of stops observed in the original conditions (untreated; K+) to those observed when either substituting K+ with Na+ (top) or pretreating RNA with DMS at 95°C (bottom). RT-stop values are colored according to the template nucleotide at position 0. The distributions are truncated at the left because stops with enrichment by less than a factor of 20 in the untreated K+ sample were not classified as strong stops. (C) Overlap between K+-dependent and DMS-sensitive strong RT stops (yellow and blue, respectively). (D) Abundance of RG4 regions within mRNA translated and untranslabeled regions. The expected abundances were estimated on the basis of the relative number of G nucleotides within these three regions of detected mRNAs.

suggesting that at least some of these RT stops also involved RG4 structures that caused RT to stall before reaching the 3’-terminal G nucleotides. Collectively, these results indicated that most G-rich regions that caused strong RT stops did so by forming RG4 structures in vitro.

The four strands of RG4 structures typically assume a parallel orientation (17) (Fig. 1A). The circular dichroism spectra of the 60-nt regions upstream of K+-dependent strong RT stops in Eef2 and Malat1, as well as that of a canonical RG4 sequence G3A2G3A2G3A2G3 (hereafter referred to as the G3A2 quadruplex), exhibited a K+-dependent increase at 263 nm, diagnostic of parallel RG4 structures (17) (fig. S3).

**Features of mammalian RG4 regions**

Of the many endogenous RNA sequences with predicted RG4-forming potential (18), only ~100 have been experimentally tested (11, 19). Therefore, the 6140 RG4 regions in the mESC transcriptome, 4034 of which were nonoverlapping, considerably expanded the repertoire of endogenous RNA sequences with experimentally supported RG4-forming capacity. Nonetheless, cellular transcripts presumably contain additional regions with intrinsic RG4-forming potential not detected in our experiment. For example, our strategy would miss (i) structures with stabilities insufficient to block RT; (ii) structures spanning more than ~60 nt, which would be too large to reside within the RNA fragments assayed for RT stops; or (iii) regions within transcripts that were not expressed in mESCs at levels sufficient to be detected in our sequencing.

To benchmark our method with previously supported examples, nearly all of which are in human transcripts (19), we performed RT-stop profiling on mRNA from human cell lines. We identified 12,099 and 12,035 RG4 regions (6506 and 6281 nonoverlapping regions) in the human embryonic kidney 293T (HEK293T) and HeLa transcriptome, respectively; of these, 7852 nonoverlapping regions were identified in at least one of the two cell lines, and 4835 were identified in both cell lines (table S2). Of the known RG4 regions within detected mRNAs, about half were detected as K+-dependent strong RT stops (fig. S4A and table S2). Recently, a high-throughput method has been developed to identify genomic sequences that can fold into DNA G-quadruplexes in vitro (20). Because DNA and RNA of the same sequence often have distinct three-dimensional structures and some
Fig. 3. RG4 regions are unfolded in mESCs. (A) Schematic of transcriptome-wide probing of RG4 folding. (B) Probing RG4 folding in vitro. The RT-stop enrichment observed after DMS treatment of RNA folded in K+ (150 mM K+) was compared to that observed after DMS treatment of RNA folded without K+ (0 mM K+). Values for regions with differences equal to or greater than a factor of 2 are indicated (blue). (C) RT-stop profiles of Eef2 and Malat1, showing results observed after DMS treatment in vitro, either with or without K+, and those observed after DMS treatment in vivo. RT-stop enrichment values corresponding to RG4 regions are in blue; values for other RT stops at G nucleotides are in gray. In vivo folding scores for the RG4 regions are shown. (D) The distribution of in vivo folding scores of the 1141 mESC RG4 regions that were examined. The 0 and 1 reference values, which represent the signal observed when treating with DMS in vitro after folding with or without K+ (B), are marked (dashed lines). (E) Distribution of in vivo folding scores observed for RG4 regions after either treating the cells with 2 μM PDS for 24 hours (PDS, red) or mock treatment (control, black). *P < 10^{-6}, paired t test. (F) Gene-specific primer extension of the ectopically expressed G3A2 quadruplex probed in vitro (after folding in either 0 or 150 mM K+) or in vivo. Shown is a phosphorimage of a denaturing gel that resolved the extension products of a P32-radioabeled primer. In the stop control, the β-mercaptoethanol quench was added to cells before DMS. The stronger RT stops are colored according to the nucleotide at the stall (A, red; C, orange; G, blue). The RG4 region is indicated (vertical line). For additional controls and sequencing ladders, see fig. S6E.

regions of DNA are either not expressed as RNA or are expressed as spliced transcripts that do not match the DNA, we expected that many DNA- or RNA-specific G4 regions would exist. Indeed, only 0.16% of the recently identified DNA G4 regions corresponded to RG4 regions found in HeLa and HEK293T cells, and of the nonoverlapping HeLa and HEK293T RG4 regions that uniquely mapped to the human genome, only 19% mapped to identified DNA G4 regions (fig. S4B).

Compared to control regions with matched nucleotide composition, the identified RG4 regions were more likely to have the four G-triplets needed to match the canonical RG4 motif (fig. S3C), G_{N_2}N_{3N_2}G_{N_3} in which each N_2 represents a linker of any sequence ranging from 1 to ~7 nt in length (18). However, 37% of these regions had fewer than four G-triplets within 60 nt upstream of the RT stop (fig. S4C) and thus would be missed by most G4-searching algorithms (18). The 6140 RG4 regions from mESCs were found in 2792 transcripts (table S1), which included both mRNAs and noncoding RNAs, such as Malat1, which was sufficiently abundant to be analyzed despite its lack of a poly(A) tail. As previously predicted (18), RG4 regions were enriched within untranslated regions (UTRs) relative to mRNA coding sequences (CDSs) (Fig. 2D; P < 10^{-5}, χ^2 test), as might be expected if some of these regions have regulatory functions. However, G nucleotides within the 60-nt regions upstream of RT stops were not more conserved than G nucleotides within flanking regions (fig. S5), suggesting that the RG4 structure—forming capacity of most RG4 regions was not evolutionarily conserved.

In sum, RT-stop profiling identified thousands of RG4 regions in the mammalian transcriptomes, thereby expanding the catalog of experimentally supported endogenous RG4 regions by a factor of >100. As predicted computationally (18), regions that form RG4 structures in vitro are not an esoteric feature of dozens of mRNAs but rather are ubiquitous within mammalian transcriptomes, bringing to the fore the question of their in vivo folding status.

Globally unfolded RG4 regions in mESCs

To identify RG4 regions that are folded in cells, we combined RT-stop profiling with elements of DMS-seq (7) to develop a method that measures, transcriptome-wide, the in vivo folding states of endogenous sequences with RG4-forming potential (Fig. 3A). In this method, cells are first treated with DMS, which rapidly enters and randomly methylates accessible N7 positions of G residues (4). RNA isolated from these cells is then subjected to RT-stop profiling. Although DMS modifies the N7 position of G more efficiently than it modifies the N1 and N3 positions of A and C residues, respectively (27), modification at N7 does not prevent Watson-Crick pairing and thus does not cause an RT stop. Nonetheless, RT-stop profiling can distinguish between RG4 regions that are folded in cells from those that are not because those that are folded in vivo are protected from modification at positions participating in the RG4 structure, enabling them to later refold during RT to generate RT stops, whereas those that are unfolded in cells can be irreversibly modified at residues that would otherwise participate in quadruplex formation in vitro, resulting in RT read-through and correspondingly attenuated RT stops (Fig. 3A).

Reasoning that the RT-stop signals of different RG4 regions might have different sensitivities to DMS treatment, we first determined, for each RG4 region, the difference in RT-stop signal observed when mRNAs were modified in vitro either with or without K+. On average, the mESC RG4 regions that were refolded and DMS-treated in the presence of K+ had RT stops that were stronger by a factor of 2.5 than those observed when refolding and treating in the absence of K+ (median, factor of 2.1), and 1342 regions had a difference of a factor of >100. As predicted computationally (18), regions that form RG4 structures in vitro are not an esoteric feature of dozens of mRNAs but rather are ubiquitous within mammalian transcriptomes, bringing to the fore the question of their in vivo folding status.
nucleotides), which showed that, as expected (7), DMS probing in vivo captured known Watson-Crick pairing within the 5.8S rRNA, as well as the intermolecular pairing between the 5.8S and 28S rRNAs (fig. S6A). Moreover, the RT-stop signals for RG4s were highly correlated between biological replicates (fig. S6B, Pearson’s r = 0.88). Inspection of the RG4 regions in both E. coli mRNA and Malat1 indicated that these RG4 regions were accessible to DMS modification in vivo, as revealed by greatly reduced RT-stop signals (Fig. 3C). The signals observed for the in vivo-modified sample resembled those observed when omitting K+ from the in vitro folding and modification reaction, which indicated that these RG4 regions were unfolded in mESCs (Fig. 3C).

To infer the folding state, DMS-probing assays must be performed within their dynamic range; beyond this range, a transiently unfolded region might instead appear to be mostly unfolded, as most of the molecules eventually become modified. The RT-stop signal at RG4 regions diminished in the unfolded reference (0 mM K+) but did not reach baseline (Fig. 3, B and C), which indicated that our in vitro treatment left a fraction of these molecules unmodified and thus showed that our in vitro modification was within its dynamic range. Moreover, DMS modification of A’s and C’s in vivo resembled that observed for our in vitro references (fig. S6C), which indicated that our in vivo probing was also within the dynamic range of the assay.

We next expanded the analysis to 1141 regions that retained a strong RT-stop signal (a factor of 10 above background) when treated with DMS in the presence of K+ in vitro and had at least a 50% reduction in that signal when K+ was excluded. For each of these regions, an in vivo folding score was calculated in which the RT-stop signal observed in vivo was expressed relative to the range of signal observed in vitro, assigning scores of 1 and 0 to the signals observed in vitro with and without K+, respectively. In vivo folding scores for the 1141 RG4 regions centered near 0 (median = 0.06) (Fig. 3D, table S4B). The folding scores of endogenous RG4 regions in mammalian cells were expressed relative to the range of the ectopically expressed G3A2 quadruplex probed in vitro (after folding in either 0 or 150 mM K+) but did not reach baseline (Fig. 3F). Table S4A, 133 of which were K+-dependent stops at G nucleotides. Among them, 47 showed a difference of a factor of 2 or more in RT-stop signal when comparing samples probed after folding with and without K+ (Fig. 4A and table S4B). The folding scores of endogenous RG4 regions centered near 0 (median = −0.15) (Fig. 4, B and C), again indicating a globally unfolded state. As observed in HEK293T cells, the ectopically expressed G3A2 quadruplex was also highly accessible to DMS, as indicated by an RT stop matching that observed for RNA modified in vitro without K+ (Fig. 4D). These results indicate that the globally unfolded state of RG4 regions is a broadly conserved feature of eukaryotic cells.

SHAPE probing of RG4 regions

In addition to the chemical probes that modify the bases, such as DMS, probes that modify ribose 2’-hydroxyl groups with efficiency depending on the local chemical environment, known as SHAPE reagents, can provide useful tools for studying RNA structures (3, 5). Among these reagents, 2-methylthioimidate acid imidazolide (NAI) has been used to probe Watson-Crick RNA structures in cells (9, 22). To test whether NAI can also distinguish the folding states of RG4 regions, we used it to treat the G3A2 quadruplex folded in vitro with or without K+ and quantified its reactivity at each nucleotide using gene-specific primer extension, substituting Na+ for K+ in the RT reaction, so that modifications within RG4 regions could be detected (Fig. 5A). Whereas the formation of Watson-Crick structure typically decreases SHAPE reactivity, formation of the G3A2 quadruplex in the presence of K+ increased NAI reactivity (Fig. 5A). Furthermore, the enhanced reactivity occurred at the last G residue of each of the first three G tracts of the G3A2 quadruplex (Fig. 5A), consistent with a recent report describing in vitro NAI probing of two other quadruplexes (23). Perhaps the transition between a G tract and a short loop in a parallel RG4 structure bends the RNA backbone to expose the 2’-hydroxyl of the last residue of the G tract (fig. S7A). In vivo NAI treatment of the G3A2 quadruplex ectopically expressed in S. cerevisiae generated a modification pattern resembling that observed for this region folded in vitro without K+ (Fig. 5A), supporting the conclusion that this quadruplex is unfolded in yeast cells.

Fig. 4. RG4 regions are unfolded in S. cerevisiae. (A) Probing in vitro folding of yeast RG4s. Otherwise, as in Fig. 3B. (B) RT-stop profiles of YPR088C. Otherwise, as in Fig. 3C. (C) The distribution of in vivo folding scores of the 31 RG4 regions that were examined in yeast. Otherwise, as in Fig. 3D. (D) Gene-specific primer extension of the ectopically expressed G3A2 quadruplex probed in vitro (after folding in either 0 or 150 mM K+) or in vivo. The primer was labeled with P32; otherwise, as in Fig. 3F.
endogenous RG4 regions that had sufficient read coverage (≥100 RT-stop reads at G nucleotides in each sample). Among these, 49 had a ≥0.1 increase in Gini coefficient when comparing the modification observed in vitro after folding with K+ compared to that observed after folding without K+ (Fig. 5D). For these 49 regions, we calculated in vivo folding scores calibrated on the Gini-coefficient differences observed in vitro (table S3D). As observed with the DMS probing, the distribution of in vivo folding scores centered near 0 (median = −0.02) (Fig. 5E), indicating that the in vivo NAI modification patterns of most RG4 regions resembled those of the unfolded state.

NAI probing complements DMS probing in three respects. First, NAI preferentially modifies specific residues of folded RG4s, whereas DMS modifies residues of unfolded RG4s. Second, NAI modification generates RT stops without requiring RG4 refolding, whereas DMS probing requires the refolding of RG4 structures in the presence of K+ to generate an RT stop. Third, NAI probing might detect less stable RG4 structures that do not stall RT in vitro and thereby escape identification by RT-stop profiling. However, unlike DMS probing of RG4 regions, NAI probing does not focus the signal onto a single RT-stop nucleotide, and it requires specific RG4 configurations, such as G tracts followed by short loops, which also reduced the number of quantifiable RG4 regions. Nevertheless, the results from these two complementary chemical-probing methods both indicated that, despite the high intracellular K+ concentration, RG4 regions are overwhelmingly unfolded in eukaryotic cells.

**Robust RG4 unfolding in eukaryotic cells**

Our results in eukaryotic cells resembled those of recent high-throughput studies showing that Watson-Crick secondary structures that form in vitro are frequently unfolded in cells (7, 9), except that the intracellular unfolding of RG4 regions was more pervasive. Whereas the previous studies identified many instances in which Watson-Crick secondary structures do form in vitro, as expected from the known Watson-Crick pairing within ribosomal RNAs, tRNAs, pri-microRNAs, mRNAs, and other RNAs, we found no compelling evidence for the folding of an RG4 region in eukaryotic cells, which implies that these cells have a very effective molecular machinery that specifically remodels RG4s and maintains them in their unfolded state.

This remodeling presumably involves adenosine 5′-triphosphate (ATP)-dependent processes. ATP depletion in yeast causes a global increase in Watson-Crick structures, suggesting that ATP-dependent processes, in particular ATP-dependent RNA helicases, play a major role in the cellular remodeling of these structures (7). Among the characterized ATP-dependent RNA helicases, DEAH box-containing helicase 36 (DHX36) accounts for most RG4-unfolding activity in HeLa cell extracts (24). To test whether DHX36 contributes to the globally unfolded state of RG4 regions in vivo, we applied DMS probing to mouse embryonic fibroblasts (MEFs) in which DHX36 was inducibly deleted through Cre-mediated recombination (25). The global distribution of folding scores was largely unchanged after DHX36 depletion, and values for individual RG4 regions were highly correlated before and after DHX36 depletion (fig. S8, A to C), indicating that DHX36 was dispensable for the global unfolding of endogenous RG4 regions. We also tested whether ATP depletion affected RG4 folding and found that the ectopically expressed G3A2 quadruplex remained largely unfolded (fig. S8D). Although redundant functions with other helicases and the inability to completely deplete ATP might explain the negative results of these experiments, our results show that the mechanism responsible for remodeling RG4s and maintaining their unfolded state is robust to either the deletion of a key helicase known to unfold RG4 structures or the substantial depletion of ATP.

**Folding of RG4 structures in bacteria**

We next applied our methods to bacterial transcriptomes. Compared to the mammalian transcriptome, the *Escherichia coli* transcriptome was substantially depleted in regions with K+-dependent strong RT stops (Fig. 6A and table S6). Only 35 K+-dependent strong RT stops were identified in *E. coli*, of which only 14 (40%) were at G nucleotides. Among these 14, none had differential DMS accessibility when comparing the results of in vitro modification with and without K+ (table S6). Similar depletion was observed within the transcriptomes of the other two bacteria that we examined, *Pseudomonas putida* and *Synechococcus* sp. WH8102 (Fig. 6A and table S6), even though their genomes are more G-rich than mammalian genomes. Only one region within the transcriptomes of these two species passed our cutoffs for calculating an in vivo folding score (a *P. putida* region with a folding score of 0.5; table S6).

Having acquired evidence for only a single, weak RG4 region in endogenously expressed bacterial RNA, we ectopically expressed the G3A2 quadruplex within the 3′ UTR of an mCherry transcript and probed its folding state in *E. coli*. In contrast to our results in eukaryotic cells, the strong RT stop corresponding to the G3A2 quadruplex was resistant to in vivo DMS modification, indicating that this region was folded in *E. coli* cells (Fig. 6, B and C). Likewise, the G3U quadruplexes, was also folded in *E. coli* cells (Fig. 6C). Since intracellular NAI probing of the G3A2 quadruplex was...
inconclusive, intracellular NAI probing of the G3U quadruplex generated the modification pattern specific to that of the folded G3U quadruplex (Fig. 5B), confirming that RG4 folding rather than protein binding protected the region from DMS modification in vivo. Thus, RG4 regions are permitted to fold in E. coli but are strongly depleted among endogenous E. coli RNAs.

To understand this depletion, we compared the growth of strains that expressed G3A2 or G3U quadruplexes to those of strains that expressed the corresponding quadruplex mutants in which point substitutions abolished RG4-forming capacity and found that the RG4-expressing strains grew more slowly than the corresponding mutant-expressing strains (Fig. 6D). Moreover, these growth defects were exacerbated after introducing stop-codon mutations that caused the mCherry coding sequence to extend through the RG4 regions (Fig. 6D). Although effects from the RG4 regions in UTRs might be attributable to either RNA or DNA quadruplex formation, the enhanced growth defects observed after introducing stop-codon mutations were attributable to only RG4 structures.

To determine the influence of folded RG4 structures on translation, we examined the translation products from each of the strains. Consistent with a previous study (26), RG4 regions downstream of the stop codon did not substantially influence mCherry production. In contrast, the G3A2 quadruplex upstream of the stop codon caused read-through of the stop codon and/or frame-shifting, generating polypeptides that were longer than expected (Fig. 6E). The G3U quadruplex also perturbed translation, causing the production of both longer and shorter polypeptides (Fig. 6E). The products of the expected size dominated when mutant RG4 regions were placed upstream of the stop codon, which indicated that the aberrant translation products were primarily the consequence of stable RG4 structures.

Discussion

The mammalian, yeast, and bacterial cells that we studied all strongly avoid the presence of folded RG4 structures in their transcriptomes but do so through different mechanisms. Based on our in vivo probing, the eukaryotic cells appear to have a robust and effective molecular machinery that specifically unfolds and maintains the thousands of RG4 regions in an unfolded state, whereas bacteria lack this machinery and have instead eliminated sequences with RG4-forming potential over the course of evolution. When considering the impaired growth rates observed for strains ectopically expressing RG4 regions, the bacterial mechanism is easy to understand, but how might the eukaryotic mechanism act? Although the critical factors remain to be identified, this mechanism differs from that which unfolds Watson-Crick structure in two key aspects. First, it is less sensitive to ATP depletion, and second, it is more pervasive, unfolding essentially every RG4 that could be monitored in mESCs, MEFs, and yeast, whereas the activities that unfold Watson-Crick structure allow many RNAs to remain folded.

We suspect that single-stranded RNA-binding proteins lie at the center of the mechanism that unfolds most eukaryotic RG4 regions. A wide variety of abundant RNA-binding proteins bind to G-rich RNA, including the heterogeneous nuclear ribonucleoprotein (hnRNP) F/H family (27, 28), hnRNP D0 (29), hnRNP M (30), hnRNP A/B (31), hnRNP A1 (32, 33), hnRNP A2 (34), CBF-A (35), and SRSP1/2 (35). The solution structures of the three quasi-RNA recognition motifs (qRRMs) of hnRNP F in complex with G-tract RNA show how qRRMs could maintain G tracts in a single-stranded conformation without blocking solvent accessibility to the N7 positions (36), which is consistent with our DMS probing results. Regardless of the identity of the machinery that operates in eukaryotic cells, it must be acting broadly throughout the transcriptome, including on untranslated RNAs and nuclear RNAs, as illustrated by the unfolding of RG4 regions within Malat1 (Fig. 3C), a nuclear noncoding RNA.

The evolutionary depletion of RG4 regions might be more tenable for bacteria than for eukaryotes for two reasons. First, maintaining machinery dedicated to the remodeling of RG4s would be more costly for species under greater selective pressure to minimize their genomes. Second, species with smaller genomes would face less frequent de novo emergence of new RG4 regions. By contrast, the eukaryotic mechanism provides opportunities for regulation. Indeed, the relative enrichment of RG4-forming regions in untranslated regions hints at the possibility that RG4 regions might be allowed to fold and impart regulatory functions in certain cell types and states or subcellular compartments (37). Alternatively,
these regions might impart function through transient folding that cannot be detected in our steady-state measurements. Another possibility is that the previously reported regulatory roles of RG4 regions, such as translational repression by RG4 regions within 5' UTRs (10), might result from the stable association of the RNA-binding proteins that maintain the RG4 regions in the unfolded state. In this scenario, the bound proteins rather than a folded RG4 would inhibit translation initiation. Clearly, more needs to be learned about this RNA structure in its native cellular contexts, and our results and methods provide the framework for doing so.

Materials and methods

In vivo DMS modification

DMS (Sigma-Aldrich; 50% diluted with ethanol) was added to mESCs or HEK293T cells cultured in 15 cm dishes to a final concentration of 8%, and evenly distributed by slow swirling. After incubating at 37°C for 5 min, the media and excess DMS were decanted, and cells were washed twice with 25% β-mercaptoethanol (Sigma-Aldrich) in PBS to quench any residual DMS. After washing, cells were lysed in 10 mL TRIzol reagent (Invitrogen) supplemented with 5% β-mercaptoethanol, and lysates were stored at −80°C. DMS was added to 10 mL of yeast culture to a final concentration of 8%. After incubating at 30°C with continuous shaking for 5 min, two volumes of 25% β-mercaptoethanol were added to the culture to stop the modification. Cells were harvested by centrifugation at 4,000 rpm for 5 min and washed with 25% β-mercaptoethanol until no residual DMS was observed at the bottom of tubes. After the final centrifugation, cells were resuspended in RNAlater solution (Invitrogen) and stored at −80°C. DMS treatment of the E. coli culture was similar to that of the yeast culture, except it was performed at 37°C. For additional details on culture of and transfection of mammalian cells, culture and induction of yeast cells, culture and induction of bacteria, and construction of RG4 expression constructs, see the supplementary materials.

In vitro folding and DMS modification

Poly(A)-selected RNA in 1 mM Mg2+ and 50 mM Tris-Cl (pH 7.0), either with or without 150 mM K+, was heated to 80°C for 2 min and then rapidly cooled to 0°C for 1 min. This refolded RNA was treated with 80 mM NAI for 5 min at either 37°C (mammalian RNA) or 30°C (yeast RNA). After treatment, RNA was phenol-chloroform extracted and precipitated. NAI-treated RNA was subjected to RT-stop profiling with 150 mM Na+ instead of 150 mM K+ during primer extension. Gini coefficients were calculated for each nonoverlapping RG4-containing region (identified as 60-nt regions upstream of K+-dependent strong RT stops) as

\[
Gini = \frac{\sum_{i=1}^{n} \sum_{j=1}^{n} |r_i - r_j|}{2n^2F}
\]

where \(n\) denotes the number of G residues in the RG4 region, and \(r_i\) denotes the RT-stop read number at position \(i\).

Calculation of in vivo folding scores

For each RG4 region that retained a strong RT-stop signal (10 fold above background) when treated with DMS in the presence of K+ and had at least a 50% reduction in that signal when K+ was excluded, an in vivo folding score (\(s\)) was calculated as

\[
s = \frac{f(\text{in vivo}) - f(\text{in vitro}; 0 \text{ mM K}^+)}{f(\text{in vivo}) - f(\text{in vitro}; 0 \text{ mM K}^+)}
\]

For the regions that had ≥100 RT-stop reads at G nucleotides after NAI probing and a difference of ≥0.1 in Gini coefficients when comparing results of RNA folded in vitro with K+ to those of RNA folded in vitro without K+, an in vivo scoring fold (\(s\)) was calculated as

\[
Gini(\text{in vitro}) - Gini(\text{in vitro; 0 mM K}^+) \quad \text{or} \quad Gini(\text{in vitro; 150 mM K}^+) - Gini(\text{in vitro; 0 mM K}^+)
\]

Although folding scores were calculated using linear functions, the conclusions of this study were not dependent on a linear relationship between the fraction of folded molecules and the extent of DMS or NAI modification.

REFERENCES AND NOTES


RNA G-quadruplexes are globally unfolded in eukaryotic cells and depleted in bacteria
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