

REVIEW

Bridging the gap between *in vitro* and *in vivo* RNA folding

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Abstract. Deciphering the folding pathways and predicting the structures of complex three-dimensional biomolecules is central to elucidating biological function. RNA is single-stranded, which gives it the freedom to fold into complex secondary and tertiary structures. These structures endow RNA with the ability to perform complex chemistries and functions ranging from enzymatic activity to gene regulation. Given that RNA is involved in many essential cellular processes, it is critical to understand how it folds and functions *in vivo*. Within the last few years, methods have been developed to probe RNA structures *in vivo* and genome-wide. These studies reveal that RNA often adopts very different structures *in vivo* and *in vitro*, and provide profound insights into RNA biology. Nonetheless, both *in vitro* and *in vivo* approaches have limitations: studies in the complex and uncontrolled cellular environment make it difficult to obtain insight into RNA folding pathways and thermodynamics, and studies *in vitro* often lack direct cellular relevance, leaving a gap in our knowledge of RNA folding *in vivo*. This gap is being bridged by biophysical and mechanistic studies of RNA structure and function under conditions that mimic the cellular environment. To date, most artificial cytoplasm have used various polymers as molecular crowding agents and a series of small molecules as cosolutes. Studies under such *in vivo-like* conditions are yielding fresh insights, such as cooperative folding of functional RNAs and increased activity of ribozymes. These observations are accounted for in part by molecular crowding effects and interactions with other molecules. In this review, we report milestones in RNA folding *in vitro* and *in vivo* and discuss ongoing experimental and computational efforts to bridge the gap between these two conditions in order to understand how RNA folds in the cell.

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1. Introduction

According to the classical view of biology, RNA has three roles, as a messenger (mRNA) that shuttles information between DNA and proteins, as an adaptor (tRNA) that translates the information stored in mRNA into protein sequence, and as a structural molecule (rRNA) that is part of the ribosome (Fig. 1). Research over the last 25 years has revealed that RNA carries out many other essential functions in the cell. RNA regulates gene expression at the transcriptional and translational levels, and this regulation often arises from the structures adopted by various RNA classes, including ribozymes, riboswitches, and RNA–protein complexes (Doudna & Cech, 2002; Serganov & Nudler, 2013). Since RNA is single stranded it can fold back on itself forming a plethora of secondary and tertiary interactions, as well as complex folding motifs, binding pockets, and active site clefts (Fig. 1). Misfolding and mutations of RNA are characteristics of many cancers and diseases; for example, triplet repeat expansion diseases are associated with Huntington’s disease, myotonic dystrophy, and Fragile X syndrome (Osborne & Thornton, 2006). Single nucleotide polymorphisms (SNPs) that alter the structural ensemble of RNA sequences also have been associated with genetic diseases (Halvorsen *et al.* 2010). Accordingly, understanding RNA structures and their dynamic regulation is an integral aspect of understanding RNA function.

The negatively charged phosphate backbone and diverse folds of RNA lead it to interact with cellular components, including metal ions, ligands, and proteins. Binding interactions with these species can change the fold of the RNA (Fig. 2). Monovalent and divalent metal ions are essential for the catalysis of small self-cleaving and large ribozymes both for folding and for active site catalysis (Serganov & Patel, 2007; Swisher *et al.* 2002). Small molecule-binding refolds riboswitches to regulate gene expression in a positive or negative mode (see Fig. 2) (Garst *et al.* 2011; Serganov & Patel, 2007).

Functional RNAs, such as tRNA, ribozymes, and riboswitches, are often found in ribonucleoprotein (RNP) complexes, which can help fold metastable RNA structures or induce a conformational change. The Protein Data Bank (PDB) has over 2000 annotated RNA-binding proteins, which include RNA chaperones, helicases, dsRNA-binding proteins, tRNA synthetases, ribonucleases (RNases) and RNA recognition motifs (RRMs) (Gerstberger *et al.* 2014). Highly studied RNPs include the ribosome, non-plant RNase P, and the spliceosome, which are responsible for the synthesis of proteins, maturation of the 5'-end of tRNAs, and splicing of pre-mRNAs, respectively. Remarkably it is the RNA component that is responsible for catalysis in these three RNPs, while the protein component provides scaffolding (Guerrier-Takada *et al.* 1983; Nissen *et al.* 2000).

Crowding plays critical but poorly understood roles in RNA folding. The cellular environment is very complex with up to 40% of the cytosol taken up by macromolecules (Minton, 2001; Zimmerman & Trach, 1991). In addition, small molecule metabolites, polyamines and other species occupy volume and interact with RNAs. Macromolecular crowding can drive the compaction of RNA and proteins, while small molecules can either stabilize or destabilize RNAs through interactions with the RNA molecule (Minton, 2001).

Nearly all of the biological components that influence RNA structure and function *in vivo* – biological ion compositions, ligands, proteins, and crowding – are missing during typical *in vitro* experiments (expanded upon in Table 1). A major goal of current research is to add back these components in order to more closely mimic *in vivo* conditions. We group studies of RNA folding into three approaches: (1) *in vitro* studies in dilute solutions; (2) *in vivo* studies in living cells; and (3) *in vivo-like* studies that mimic *in vivo* conditions. We also discuss how *in silico* methods facilitate each of these approaches. There are advantages and limitations to working in each of these conditions, and experiments in each can yield unique insights into

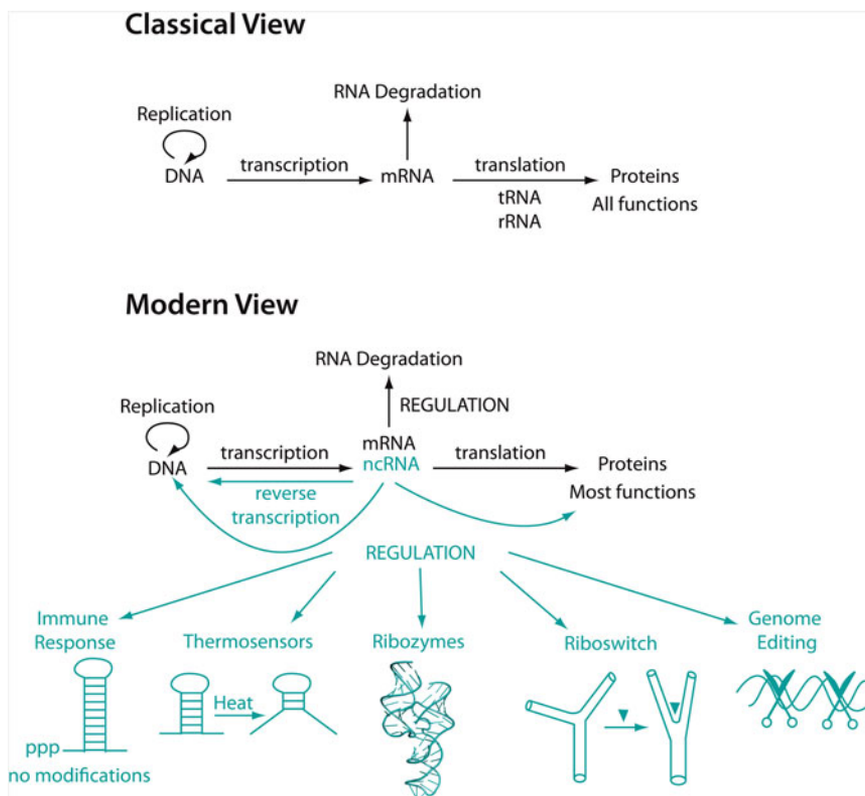


Fig. 1. The Classical View (top) and the Modern View (bottom) of RNA's role in biology. In the classical view of biology, RNA (top) serves as a messenger molecule between DNA and proteins and proteins have all the main functions in cells. Messenger RNA serves to translate information from DNA to proteins. The modern view of biology (bottom) has emerged in the last 25 years as the field learns more about the many functions of RNA. Non-coding RNA (ncRNA) has vast regulatory functions, some of which include immune responses ('ppp' = 5'-triphosphate, which activates PKR) (Nallagatla *et al.* 2007), thermosensors, ribozymes, riboswitches, and genome editing. In the modern view of biology, proteins still have most cellular functions, but RNA plays essential roles in the cell beyond its classical functions.

the biological functions of RNAs. Structures and folding pathways of RNA have been studied mostly in dilute *in vitro* conditions, resulting in fundamental insights into RNA structure and function. However, there is a deep desire to understand how Nature works, and the *in vivo* environment is very different from typical *in vitro* solution conditions (Table 1 and Fig. 3). In particular, the majority of thermodynamic experiments studying the energetics of folding and associated pathways (Freier *et al.*, 1986b; Schroeder & Turner, 2009) have been conducted in non-biological salt concentrations (London, 1991; Lusk *et al.* 1968; Minton, 2001; Romani, 2007; Truong *et al.* 2013). There are also myriad RNA–protein interactions *in vivo*, many of which profoundly affect RNA folding and function.

Over the last few years, *in vivo* experiments probing RNA structure in living cells have revealed significant differences in many RNA structures as compared to *in vitro* (Kwok *et al.* 2013; Rouskin *et al.* 2014; Tyrrell *et al.* 2013). *In vivo* studies, while desirable because of their biological relevance, are at the same time limited in that they typically elucidate only the ensemble structure of each RNA transcript, do not deconvolute RNA–protein interactions versus RNA self-structure, and cannot easily perturb or control solution conditions. In particular, biophysical studies that can be readily conducted under highly controlled *in vitro* conditions are often simply not feasible *in vivo*. In an effort to gain more insight into the structure and function of RNA in the cellular environment, recent studies have focused on the folding pathway, structure, and function of RNAs under *in vivo-like* conditions, which mimic conditions in the cell (Desai *et al.* 2014; Dupuis *et al.* 2014; Kilburn *et al.* 2013; Nakano *et al.* 2014; Strulson *et al.* 2012, 2013).

In silico prediction and modeling of RNA structure is an important tool used in all three of the above approaches to provide additional insight into RNA structure and function (Dawson & Bujnicki, 2016; Seetin & Mathews, 2012a). Prediction of canonical base pairs, for example, provides testable hypotheses for RNA structure and also provides frameworks for interpreting experimental results. Likewise, experimental data aid in improving *in silico* structure prediction.

In this review, we discuss major achievements in describing and understanding RNA folding and structure through *in vitro*, *in vivo*, and *in silico* efforts. The next section introduces the reader to *in vitro* studies of RNA folding, which set the stage for

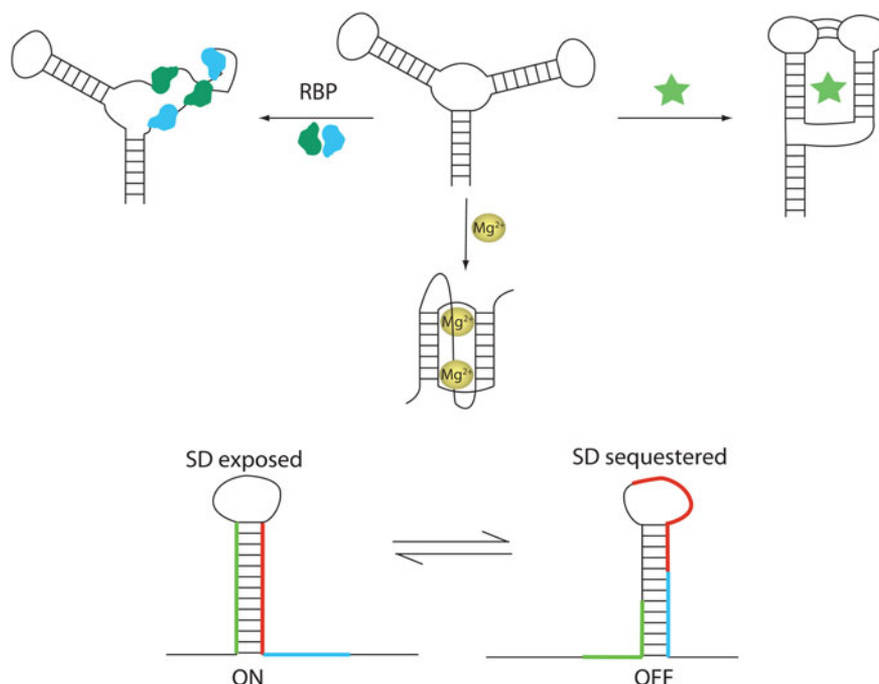


Fig. 2. RNA interactions with RNA-binding proteins (RBP, left), metal ions (central), and ligands (star, right) can result in structure changes. Unlike typical *in vitro* conditions, there are other molecules and complex solution conditions *in vivo* that can interact with RNA and change its structure. These structure changes can result in an RNA with less structure (top left) more structure (top right), or an alternate conformation than the structure that is prevalent *in vitro* (bottom). Also shown (bottom) are the bacterial expression platforms of riboswitches that switch between two mutually exclusive structures that turn a gene ON (left) or OFF (right) by exposing or sequestering the Shine–Dalgarno sequence (blue).

Table 1. Comparison of *in vitro* and *in vivo* solution conditions

Condition	<i>in vitro</i> (Historical in the RNA field)	<i>in vivo</i>
Molecular crowding	0%	20–40% (w/w)
Monovalent salt	0–1 M ^a	140 mM K ⁺ ^b
Divalent salt (free)	0–100 mM ^c	0.5–1.0 mM Mg ²⁺ (eukaryotes) ^d 1.5–3.0 mM Mg ²⁺ (prokaryotes) ^e
Divalent salt (total)	0–100 mM ^c	20 mM
Ionic strength	0–1 M (monovalent only) 0–0.3 M (divalent only)	0.142–0.143 M (eukaryotes) 0.145–0.149 M (prokaryotes)

Conditions *in vitro* are the conditions historically used to study RNA. Typical values in the literature are listed in the table, although actual values differ across various studies. *In vivo-like* conditions, not provided in this table, typically emulate at least one of the conditions missing during *in vitro* experiments.

^a Typically Na⁺ is used *in vitro* although K⁺ is found. Freier *et al.* (1986a), Xia *et al.* (1998).

^b Feig & Uhlenbeck (1999).

^c Typically Mg²⁺ is used. Herschlag & Cech (1990), Tanner & Cech (1996).

^d Alberts *et al.* (1994), London (1991), Romani (2007).

^e Lusk *et al.* (1968), Truong *et al.* (2013).

in vivo and *in vivo-like* studies of RNA folding. We focus on recent efforts to understand how RNA folds in the cell by bridging the gap between knowledge of RNA structure and folding *in vitro* and *in vivo*, which has led to an emerging field that studies RNA under *in vivo-like* conditions. We also discuss ways in which the accuracy of *in silico* modeling could be improved with experimentally derived *in vivo* structure probing data. We conclude by discussing advances needed under cellular-like conditions to better understand how RNA folds in the cell.

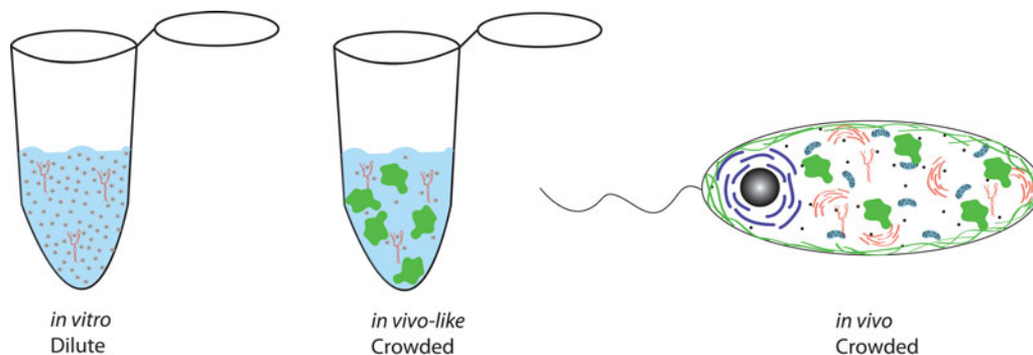


Fig. 3. Artist's rendition of *in vitro* conditions (left), *in vivo* conditions (right) and *in vivo-like* conditions (center). Typical *in vitro* solutions are dilute with high monovalent ion concentrations that are very different from cellular conditions. The cellular environment is complex with monovalent and divalent salts, macromolecules, cosolutes, and organelles. *In vivo-like* conditions (center) bridge *in vitro* and *in vivo* conditions and are more complex than *in vitro* conditions with added synthetic crowding agents and proteins and physiological ion concentrations. However, *in vivo-like* conditions are still much less complex than those prevailing *in vivo*.

2. Setting the stage

2.1 *In vitro* studies of RNA folding

Most of what we currently know about RNA structure and folding comes from studies completed *in vitro*, under experimental conditions that favor a folded state. Such studies are typically conducted in dilute solutions with high concentrations (~ 1 M) of monovalent ions (Freier *et al.* 1986b) and/or (~ 10 mM) divalent ions (Herschlag & Cech, 1990), especially Mg^{2+} , or under conditions that facilitate population of a desired folding intermediate, for example, by renaturing the RNA at an unusual temperature or salt concentration (Baird *et al.* 2005). These solution conditions are advantageous for studying folding because they can be chosen such that the RNA folds in an apparent two-state manner or the RNA populates just a single intermediate, but have the drawback that they differ profoundly from *in vivo* conditions, which have predominantly ~ 140 mM K^+ and 0.5–3 mM Mg^{2+} (expanded upon in Table 1).

An advantage of using high concentrations of monovalent salts is that they compete with trace polyvalent metal ions and hydroxide ions for the phosphate backbone thereby reducing RNA degradation. In addition, high monovalent salt conditions minimize end fraying of RNA hairpins, favoring two-state folding (Freier *et al.* 1986b). As we describe below, the thermodynamics and kinetics of systems, ranging from simple RNAs, such as hairpins and bulges, to complex RNAs and RNPs, such as ribozymes and the ribosome, have been well characterized. Many aspects of the RNA-folding process can be understood by the application of techniques and the systematic manipulation of conditions only possible under *in vitro* or under *in vivo-like* environments.

2.1.1 Major advances: elucidating RNA folding pathways *in vitro*

With the invention of various enzymological methods, such as PCR, cloning, T7 transcription and chemical synthesis, RNA preparation has advanced to the point where RNA of almost any sequence and length can be studied (Hoseini & Sauer, 2015; Li *et al.* 2011; Milligan *et al.* 1987; Mullis, 1990). A wide variety of techniques have been applied to the study of RNA *in vitro* (Table 2). The earliest studies on RNA were conducted on homoribopolymers, such as polyU and polyA, which revealed that stacking – the non-bonded interactions between the surfaces of the bases – contributes to RNA stability (Richards *et al.* 1963; Suurkuusk *et al.* 1977). These studies also provided the first indications that individual RNAs adopt structure. An early breakthrough was from studies of tRNA, which could be isolated from living systems owing to its high cellular abundance, which led to insights into RNA tertiary structure. The cloverleaf base pairing of tRNA had been first predicted from sequence alignments of sequence variants (Levitt, 1969). Solving the crystal structure of tRNA confirmed its cloverleaf secondary structure and revealed novel tertiary interactions (Kim *et al.* 1973; Robertus *et al.* 1974). The crystal structure of tRNA provided the first direct evidence that RNAs can form complex structures, akin to those of proteins, and that stacking, base pairing, and tertiary contacts all contribute to the adoption of complex three-dimensional (3D) structures (Sussman *et al.* 1978). With the advent of chemical synthesis techniques, ~ 100 – 200 mer of DNA and eventually ~ 50 mer RNA of any sequence could be made (Matteucci & Caruthers, 1981; Scaringe *et al.* 1998; Sierzchala *et al.* 2003), with a plethora of atomic modifications. Semi-synthetic approaches were then developed that combine enzymological and chemical synthesis to facilitate the introduction of mutations both at the nucleotide and functional group levels in RNAs of any size (Moore & Sharp, 1992).

Table 2. Common experimental techniques used to study RNA structure and folding

Method	Condition	What is probed	References
Low-resolution methods			
Small-angle X-ray scattering	<i>in vitro</i>	Overall structure	Yang (2014), Pollack (2011)
Optical melting	<i>in vitro</i>	Thermodynamics/thermostability	Schroeder & Turner (2009)
Stopped-flow, hand mixing	<i>in vitro</i>	Kinetics	
Temperature-jump	<i>in vitro</i> and <i>in vivo</i>	Folding kinetics/thermodynamics	Dyer & Brauns (2009), Gao <i>et al.</i> (2016)
Single-molecule FRET	<i>in vitro</i>	Overall structure	Klostermeier & Millar (2001), Roy <i>et al.</i> (2008)
Hydroxyl radical footprinting	<i>in vitro</i> and <i>in vivo</i>	Structure context of nucleotides	
Enzymatic mapping	<i>in vitro</i>	Structure context of nucleotides	Wan <i>et al.</i> (2011), Clatterbuck Soper <i>et al.</i> (2013)
Chemical mapping	<i>in vitro</i> and <i>in vivo</i>	Structure context of nucleotides	Kwok <i>et al.</i> (2015), Weeks (2010)
High-resolution methods			
X-ray crystallography	<i>in vitro</i>	Å-resolution structure	Reyes <i>et al.</i> (2009)
Nuclear magnetic resonance	<i>in vitro</i>	Å-resolution structure	

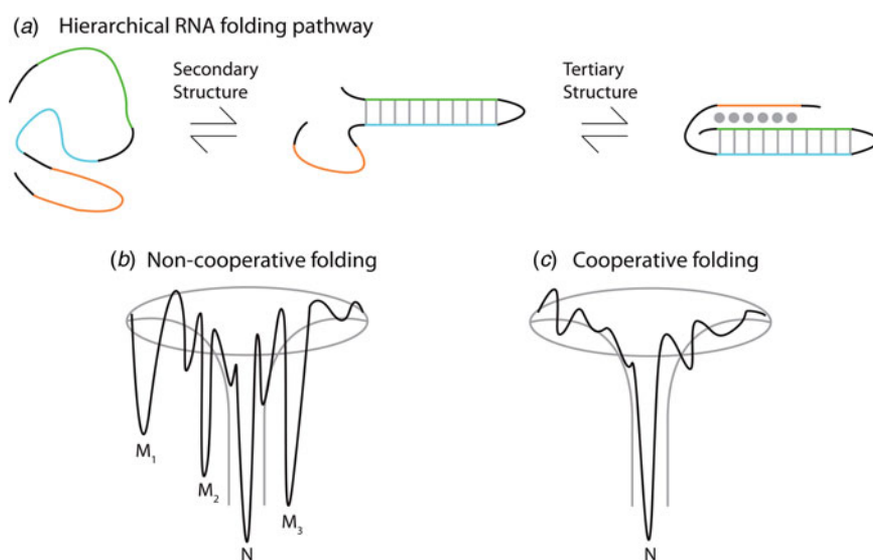


Fig. 4. Depiction of the hierarchical RNA folding pathway and folding funnels for non-cooperative and cooperative folding. (a) RNA folds in a hierarchical manner in which secondary structures form followed by tertiary structure. Hierarchical folding can be (b) rugged and non-cooperative in which the pathway intermediates are populated and the RNA can form misfolds (M_i) before populating the native state (N), or folding can occur in a (c) cooperative manner in which the intermediates do not populate and the RNA folds in a single transition.

Thermodynamic and kinetic studies under *in vitro* conditions provide insight into the complex folding pathways of many functional RNAs. Ribozymes and riboswitches are ideal for the study of RNA folding because their function serves as a read-out for the occupancy of the native state (Banerjee *et al.* 1993; Crothers *et al.* 1974; Mitchell & Russell, 2014; Mitchell *et al.* 2013; Rook *et al.* 1998). Major themes are that large RNAs fold on a rugged pathway through populated intermediates, largely in a hierarchical manner, where secondary structures form before tertiary contacts, as demanded by the topologies of these complex RNAs (Fig. 4) (Brion & Westhof, 1997; Mitchell & Russell, 2014; Solomatin *et al.* 2010; Tinoco & Bustamante, 1999; Wan *et al.* 2010). It is informative to consider these principles on several specific RNAs. Using temperature-dependent nuclear magnetic resonance (NMR) and relaxation kinetics, the mechanism of tRNA unfolding was elucidated (Crothers *et al.* 1974; Hilbers *et al.* 1976; Stein & Crothers, 1976). Five distinct transitions were mapped to the four arms and the tertiary contacts (Crothers *et al.* 1974). Secondary structures form on a fast timescale (μ s to ms) followed by folding of the tertiary structure on a slower timescale (ms to s). In the presence of monovalent metal ions, multiple thermal unfolding transitions are observed for these processes (Stein & Crothers, 1976). These transitions merge into one as Mg^{2+} concentrations are increased, revealing that Mg^{2+} induces an apparent two-state folding. Larger functional RNAs, ribozymes, and riboswitches also fold in a hierarchical manner *in vitro* (Fig. 4a).

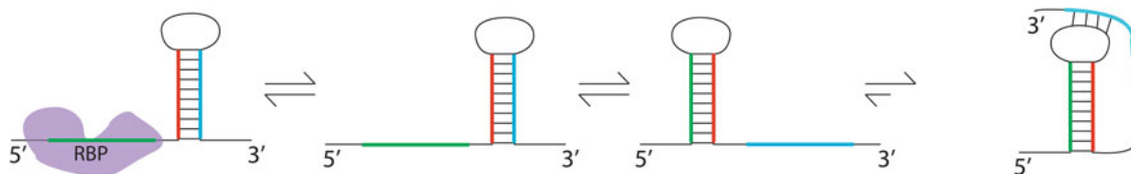


Fig. 5. Different RNA structures can be populated under *in vitro*, *in vivo*, and *in vivo-like* conditions. RNA structures induced by the cellular environment, including proteins and crowding, are shown in the two outermost structures. The conditions *in vitro* favor the population of a structure that may not always be the functional RNA structure (center two structures). Depending on the *in vivo-like* conditions chosen, specific RNA structures will be populated.

The *Azoarcus* group I ribozyme was used to determine the influence of tertiary interactions on RNA folding (Fig. 5). This ribozyme has been shown to fold quickly, with ~80% of the ribozyme folded into the native state in under 50 ms in 15 mM Mg^{2+} (Rangan *et al.* 2003). To determine the roles of tertiary interactions in ribozyme folding, the tertiary contact between the P9 GAAA tetraloop and its J5/5a receptor were perturbed (Chauhan & Woodson, 2008). While the WT ribozyme folded in a cooperative manner to the native state, the tetraloop mutant occupied many previously hidden intermediates on the folding pathway, even at 50 mM Mg^{2+} . This study indicated that tertiary contacts promote cooperative RNA folding.

More recently, methods have been developed to study RNA folding on the nucleotide level and at the millisecond timescale (Merino *et al.* 2005; Scalvi *et al.* 1997; Zhuang *et al.* 2000). Experiments using hydroxyl radical mapping yielded insight into the pathway of tertiary structure formation and folding kinetics in the *Tetrahymena* Group I Intron (Sclavi *et al.* 1998). Combined with time resolved small-angle X-ray scattering (SAXS) (Roh *et al.* 2010), hydroxyl radical footprinting on the *Tetrahymena* ribozyme folding pathway uncovered an initial collapse of structure on the millisecond timescale during the dead time of the instrument. During the subsequent time course, tertiary contacts and several intermediates were elucidated (Sclavi *et al.* 1998).

The folding pathways of large functional RNAs have proven to be quite complex with intermediates that can be trapped for minutes to hours (Banerjee & Turner, 1995; Chadalavada *et al.* 2002; Zarrinkar *et al.* 1996). For example, 90% of the *Tetrahymena* ribozyme is found in a misfolded state that transitions to the native state with hour timescale kinetics (Banerjee & Turner, 1995), and the hepatitis delta virus (HDV) ribozyme folds through numerous intermediates, some long-lived (Chadalavada *et al.* 2002). Long-lived misfolded intermediates are often very similar in structure to the native RNA and typically arise from a secondary structure mispairing or an incorrect 3D topology (Mitchell *et al.* 2013; Treiber *et al.* 1998; Wan *et al.* 2010). For instance, a long-lived intermediate occurs in the *Tetrahymena* ribozyme where P3 is docked correctly but the topology of the ribozyme is incorrect (Mitchell & Russell, 2014; Mitchell *et al.* 2013). To fold into the native state, this misfold needs to undergo a global unwinding of structure. Importantly, the extent to which these pathways and intermediates are populated *in vivo* is unknown. Indeed, some of these folding intermediates are affected by the method by which the RNA is purified. For example, the wild-type HDV ribozyme has the optimal rate of catalysis when the ribozyme is folded co-transcriptionally, as opposed to being renatured prior to assay (Chadalavada *et al.* 2007). In addition, choice of flanking sequences can profoundly affect the activity of small and large ribozymes (Cao & Woodson, 1998; Chadalavada *et al.* 2000).

2.1.2 Major advances: applying biophysical techniques to study RNA folding *in vitro*

Using optical melting, a set of thermodynamic parameters have been established to estimate folding free energies from sequence and structure alone (Andronescu *et al.* 2014; Lu *et al.* 2006; Turner & Mathews, 2010; Xia *et al.* 1998). The nearest-neighbor model predicts the free energy and stability of an RNA from each base pair's nearest neighbor, along with initiation, symmetry, and terminal-AU base pair terms. Nearest-neighbor terms for certain loops, those regions without canonical base pairs, have also been determined (Mathews *et al.* 2004). As noted below, these experimental parameters have been incorporated in RNA structure prediction programs that find the lowest free energy structures for an input RNA sequence (Mathews, 2006; Reeder *et al.* 2006; Seetin & Mathews, 2012a). Parameters to account for complicated tertiary interactions and loops are still being revised (Liu *et al.* 2010b, 2011; Lu *et al.* 2006). The nearest-neighbor parameters currently available were measured under highly folding *in vitro* conditions of 1 M NaCl.

Low-resolution methods provide information about the structure of RNA on both the global and nucleotide length scales. Although these techniques do not give atomic resolution, they have significantly faster throughput than crystallography or NMR structures while still providing insight into the fold and function of RNA. SAXS and Förster Resonance Energy Transfer (FRET) provide low-resolution information on the overall fold of an RNA. RNA is particularly amenable to SAXS because the phosphate backbone is electron-rich and scatters X-rays well. Different solution conditions can be prepared

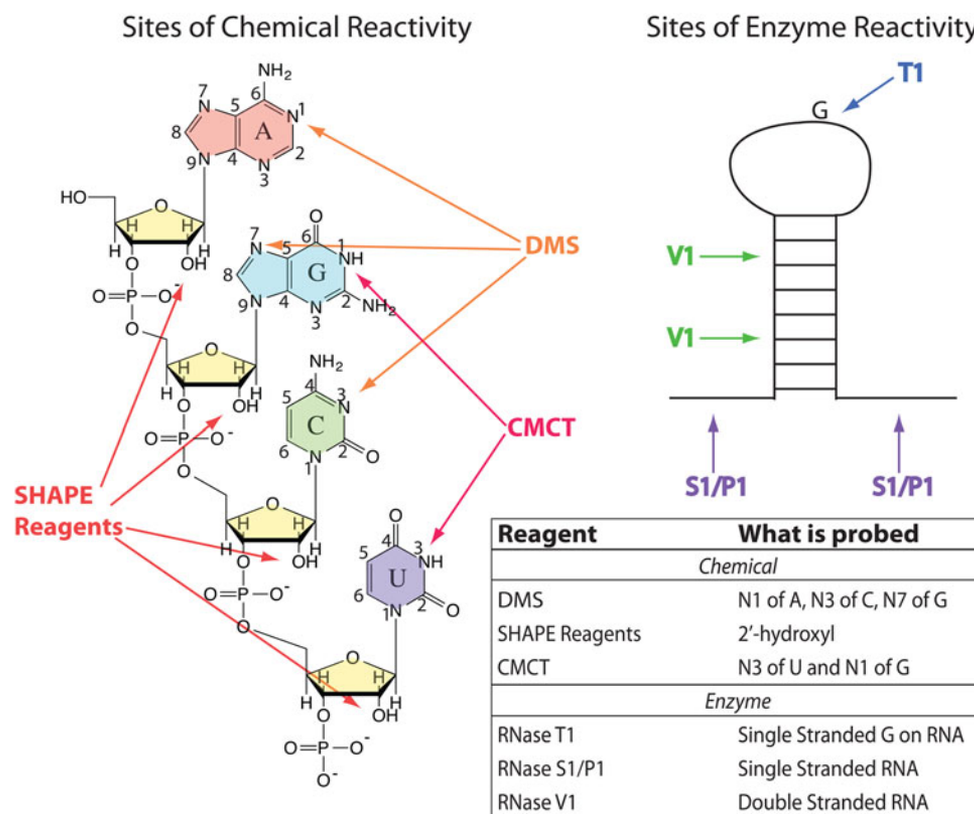


Fig. 6. RNA modifications by DMS, SHAPE reagents (selective 2'-hydroxyl acylation analyzed by primer extension), and CMCT (1-cyclohexyl-(2-morpholinoethyl)carbodiimide metho-p-toluene). SHAPE reagents modify the 2'-hydroxyl on the sugar of all four ribonucleobases. SHAPE reagents include 1M7 (1-methyl-7-nitroisatoic anhydride), NMIA (N-methylisatoic anhydride), and NAI (2-methylnicotinic acid imidazolide). DMS modifies the N1 of A and the N3 of C as well as the N7 of G. CMCT modifies N3 of U and N1 of G. The chemical modifications (except N7 of G) can be detected immediately by RT followed by gel electrophoresis or high-throughput sequencing, and the enzymatic cleavages, which cleave single- and double-stranded RNA, can be read out through gel electrophoresis or high-throughput sequencing.

and examined quickly by SAXS to elucidate RNA structural changes. The structures of several functional RNAs and RNA-protein complexes have been explored using SAXS, including ribozymes, riboswitches bound and unbound to ligand, and the spliceosome (Pollack, 2011). FRET studies, in which acceptor and donor fluorophores are attached to the RNA at key locations, have helped elucidate folding intermediates (Walter, 2001). Using single-molecule FRET, or smFRET, the *Tetrahymena* ribozyme was found to fold into multiple conformations, nearly all of which were active, indicating that the ribozyme populates multiple native states (Solomatin *et al.* 2010). Upon exposure to denaturant, the ribozyme re-populated the native conformations, indicating the results are independent of original conformation. Both SAXS and smFRET have been applied to RNA folding under *in vivo-like* conditions, as discussed below (Paudel & Rueda, 2014; Strulson *et al.* 2013).

Structure probing methods serve essential roles in elucidating the structures of functional RNAs at the nucleotide level. Several chemical probes have been employed to attack and modify the RNA bases, sugar, and backbone, in order to reveal the base pairing status of the nucleotides. Commonly used chemical probes include dimethyl sulfate (DMS), carbodiimide tosylate (CMCT), and SHAPE reagents, which allow selective 2'-hydroxyl acylation – each of which is analyzed by primer extension via reverse transcription. Commonly used enzymatic probes are RNases T1, V1, and S1. Targets of these probes and methods of readout are provided in Fig. 6. Structure probing of RNAs *in vitro* has revealed very complex structures, as well as binding sites of ligands, metal ions, and proteins. As discussed below, structure probing with chemical probes can be used *in vivo* as well.

Very recently, RNA structure *in vitro* has been probed genome-wide at the nucleotide level, utilizing the power of next-generation sequencing. Several methods have been developed to map entire transcriptomes. Parallel Analysis of RNA Structure (PARS) cleaves double-stranded regions with RNase V1 and single-stranded regions with RNase S1, and FragSeq cleaves single-stranded regions with nuclease P1 (Kertesz *et al.* 2010; Underwood *et al.* 2010). In PARS, RNA is extracted from cells and aliquots are separately exposed to each nuclease, the digested RNA is converted to cDNA through reverse transcription, and then deep sequenced to map the reverse transcriptase stops to the genome. A PARS score is determined from



the log ratio of V1/S1 sequencing reads, where a high PARS score indicates more RNA structure (Kertesz *et al.* 2010). In FragSeq, RNA is extracted from cells, and one aliquot is treated with P1 nuclease and a second aliquot is untreated (Underwood *et al.* 2010). RNA-seq is then performed on each aliquot, and a cutting score is determined for each mapped nucleotide that indicates the propensity to be cut by P1 nuclease. The cutting score is then used to annotate RNA secondary structures and/or to restrain RNA secondary structure prediction. Genome-wide studies in several organisms, both *in vitro* and *in vivo*, have found that there is significantly more structure in the coding regions than the untranslated regions of RNAs (Ding *et al.* 2014; Kertesz *et al.* 2010; Li *et al.* 2012; Wan *et al.* 2014; Zheng *et al.* 2010). There is also less structure in the start and stop codons than in the rest of a transcript, which presumably facilitates read-through by the ribosome.

Using a method similar to PARS but differing in that the RNA structure is probed at several temperatures, PARTE (Parallel Analysis of RNA Structures with Temperature Elevation) was used to obtain the folding free energies for yeast transcripts genome-wide *in vitro* (Wan *et al.* 2012). RNA from yeast was folded between 30 and 75 °C and exposed to RNase V1 followed by deep sequencing. By examining the melting temperatures (T_m) of RNAs, non-coding and coding RNAs could be distinguished and RNAs with distinct cellular functions could be identified. Functional non-coding RNAs (ncRNAs) were found to have a higher T_m on average than mRNAs.

Three methods that utilize DMS chemistry to determine transcriptome-wide RNA structure were recently published: Structure-seq, DMS-sequencing (DMS-seq), and modification sequencing (Mod-seq) (Ding *et al.* 2014; Rouskin *et al.* 2014; Talkish *et al.* 2014). To date, only DMS-seq has been applied *in vitro* and all of the methods have been applied *in vivo*. These methods are described in more detail below.

2.1.3 Benefits and limitations of *in vitro* studies

Many of the foundational experiments on RNA folding and structure have come from *in vitro* experiments, and numerous underlying mechanisms of RNA folding and function have been discovered *in vitro*. Studies *in vitro* have revealed the folding pathways and structures of RNAs. More recently, methods have been developed to probe the structure of RNAs genome-wide. Major advances include elucidating fast formation of secondary structure and slow formation of the tertiary contacts, understanding of RNA folding energetics, establishment of nearest-neighbor parameters, and determination of structures of functional RNA motifs. The complex structures that RNA adopts enable diverse functions. Experimental techniques, ranging from structure probing to kinetic methods, have been applied to RNA across diverse pH, salt, and temperature conditions.

The major limitation of *in vitro* experiments is that the solution conditions are very different from the cellular environment and unavoidably lack many of the components present in cells, which can influence RNA folding and function. These limitations necessitate the development of experiments and techniques under *in vivo* and *in vivo-like* conditions to determine how RNAs fold and respond to cellular environmental conditions.

2.2 *In vivo* studies of RNA folding

In the previous section, we provided an overview of RNA folding *in vitro*. In this section we discuss recent advances made *in vivo* to understand RNA folding. We note that RNA structure has also been explored to a lesser extent in cellular extracts. Experiments in extracts contain more proteins bound to RNA than *in vitro* experiments but less than *in vivo* studies, as supported by recent comparisons of low DMS reactivity assignments amongst *in vitro*, extract, and *in vivo* studies (Ding *et al.* 2015). Studies in extracts for RNAs with high positive predictive value (PPV) between reactivities *in vitro* and *in silico*, such as the ribosome, have been shown to be biologically relevant (Ding *et al.* 2015; Moazed *et al.* 1986a). Likewise, for RNAs with low PPV between reactivities *in vitro* and *in silico*, studies in extracts might not provide the full complement of interactions. While experiments in cell extracts share many similarities with *in vivo* conditions, thermodynamic assays cannot be easily performed in extracts due to the denaturation and signal of other biomolecules.

An ultimate goal of RNA-folding studies is to understand how RNA behaves in the cell. The majority of the methods developed to study RNA *in vivo* are structure probing, where several chemicals known to penetrate the cell membrane are applied to modify RNA. Structure probing has been used to study the structures of RNAs *in vivo* on both the single gene and genome-wide levels, and has resulted in a breadth of information regarding structures that RNA forms inside living cells. These studies have revealed novel *in vivo* RNA folds, RNA–protein interactions, and novel regulatory roles.

2.2.1 Major advances: transcript-specific RNA structure mapping *in vivo*

Structure probing of RNA *in vivo* uses small chemicals such as SHAPE reagents, DMS, and CMCT, which penetrate cells and modify solvent-accessible regions of the RNA (Bloomfield *et al.* 2000; Ehresmann *et al.* 1987). Structure probing methods using chemicals have revealed that for some transcripts there are significant differences between RNA structures formed



in vivo and *in vitro*. We first describe *in vivo* structure probing experiments on single transcripts, followed by experiments across a genome.

The first *in vivo* nucleic acid structure probing study was from the Gilbert laboratory, where binding of multiple proteins to their cognate sites was observed using DMS modification (Nick & Gilbert, 1985). Structure probing is outlined in Fig. 6. Briefly, DMS methylates adenine and cytosine on the Watson–Crick face and guanine on the Hoogsteen face. The modification on A and C is read out directly by stops in reverse transcription (RT) one position before the methylated base, while the methylated G is treated with aniline to create an abasic site followed by RT read out, which again stops one position before the modified base (Bloomfield *et al.* 2000; Ehresmann *et al.* 1987). The RT can be read out in a gene-specific fashion by polyacrylamide gel electrophoresis (PAGE) or capillary electrophoresis (CE), and in a library fashion with next-generation sequencing (see the next section) (Kwok *et al.* 2013).

The first report of RNA structure comparisons *in vivo* and *in vitro* came from the Cech laboratory (Zaug & Cech, 1995). Structure probing with DMS was used to map the structures of two known protein-bound RNAs, telomerase RNA and U2 snRNA, as well as the *Tetrahymena* ribozyme. Protections from reactivity *in vivo* compared with *in vitro* indicate either protein protection or gain of base pairing, while enhancements of reactivity indicate refolding to expose RNA bases. Telomerase RNA and U2 snRNA showed different reactivity patterns *in vivo* versus *in vitro*, consistent with the influence of protein binding on DMS reactivity. As expected, the group I ribozyme had very similar nucleotide reactivity *in vivo* and *in vitro*, demonstrating that the ribozyme is not protein-bound and self-splices without protein assistance *in vivo*.

Our group investigated structures of high and low abundance RNAs, also on a gene-specific basis, and compared DMS and SHAPE reactivities *in vivo* and *in vitro*. For low abundance RNAs we developed a gene-specific ligation-mediated PCR (LM-PCR) approach (Kwok *et al.* 2013). These studies, which were in the model plant species *Arabidopsis thaliana*, revealed *in vivo* footprinting on high abundance 25S rRNA and 5-8S rRNA, as well as on the low abundance U12 snRNA. We showed that different bases in 5-8S rRNA are methylated *in vivo* and *in vitro*, which provided evidence for 5-8S rRNA refolding *in vivo*. These studies also provided critical control reactions that strongly supported DMS modification of RNA occurring *in vivo* and DMS being completely quenched prior to workup of the *in vivo* reaction. These controls apply equally to the genome-wide studies in the next section.

2.2.2 Major advances: genome-wide RNA structure mapping *in vivo*

Recently, several groups including ours have developed high-throughput methods to probe RNA structure in living cells transcriptome-wide. These studies revealed significant differences in RNA structure *in vivo* compared to *in vitro* and *in silico* predicted (Ding *et al.* 2014; Kwok *et al.* 2015; Rouskin *et al.* 2014; Talkish *et al.* 2014). Three separate methods using DMS to probe RNA structure *in vivo* were published in 2014: Structure-seq (Ding *et al.* 2014), DMS-seq (Rouskin *et al.* 2014), and Mod-seq (Talkish *et al.* 2014), each of which utilizes the next-generation sequencing to probe RNA structure transcriptome-wide.

Each of these studies revealed novel information on RNA structure and possible regulatory functions of those structures. In Structure-seq, the PPV describes the fraction of base pairs in the *in vivo* DMS-restrained predicted structure that are also predicted in the unrestrained *in silico* predicted structure (Ding *et al.* 2014). Of the greater than 10 000 mRNAs evaluated in this fashion, most had a PPV value far from unity, with a maximum PPV of the distribution slightly <0.4. This observation indicates that the *in vivo* structures of many RNAs cannot be predicted well purely *in silico*, using only sequence information and thermodynamic parameters originally derived *in vitro*. We also observed that the mRNAs with the lowest PPV distribution (bottom 5%) were enriched in annotations of biological function of stress and stimulus response, while the mRNAs with the highest PPV distribution (top 5%) were enriched in housekeeping functions (Ding *et al.* 2014).

One possibility is that housekeeping RNAs have well-defined folds, while stress-related RNAs have ill-defined folds or adopt many folds. DMS-seq in yeast found that certain mRNAs are less structured *in vivo* than naked, protein-free RNA *in vitro*, and under *in vivo* ATP depletion the mRNAs on a whole become more structured, with the implication that ATP-dependent processes contribute to RNA unfolding. It is likely that a range of factors *in vivo* contribute to RNA structure (Rouskin *et al.* 2014). Mod-seq was used to reveal the binding location of the L26 protein by deletion in yeast; upon L26 deletion, 58 nucleotides became more reactive to DMS *in vivo* and most of these nucleotides were located in the 5-8S–25S rRNA interface where L26 is known to bind (Talkish *et al.* 2014).

Individual copies of a given RNA sequence can adopt different conformations owing to the single-stranded nature of RNA. Indeed, this may be the origin of the low PPV value in the stress-related genes (Ding *et al.* 2014) in that structure probing methods reveal the average of all populated structures at some instant in time. There is experimental evidence that some



transcripts appreciably populate multiple structures *in vitro*. Using the PARS method, ~4% of mRNAs had both high RNase V1 and RNase S1 activity, which cleave paired and unpaired RNA, respectively, under *in vitro* conditions (Wan *et al.* 2014). The high extents of cleavage by both nucleases suggest that populations of those mRNAs adopt multiple conformations simultaneously *in vitro*, and potentially *in vivo*.

Genome-wide studies revealed a triplet periodicity in mRNA nucleotide reactivity in yeast, mouse, and humans *in vitro* (Incarnato *et al.* 2014; Wan *et al.* 2014), as well as in *Arabidopsis in vivo* (Ding *et al.* 2014; Kertesz *et al.* 2010). The triplet repeat in reactivity is observed in the coding sequence but not in the untranslated regions. At present the mechanism behind the periodicity is not understood. Observation of the repeat *in vitro* suggests that occupancy of ribosomes is not necessary. Additional studies under *in vitro*, *in vivo*, and *in vivo-like* conditions will be necessary to attain a molecular-level understanding of the triplet periodicity in mRNA.

High-throughput sequencing has been coupled with CLIP (crosslinking and immunoprecipitation) to probe RNA-binding protein sites transcriptome wide in HITS-CLIP (high-throughput sequencing of RNA isolated by crosslinking immunoprecipitation) and PAR-CLIP (photoactivatable-ribonucleoside-enhanced crosslinking and immunoprecipitation) (Hafner *et al.* 2010; Licatalosi *et al.* 2008; Weyn-Vanhenhenryck *et al.* 2014). Studies using both of these methods on specific proteins have revealed novel sites of protein binding to RNA as well as possible protein regulatory functions (Hafner *et al.* 2010; Licatalosi *et al.* 2008). Briefly, in HITS-CLIP, RNA is crosslinked to proteins, the protein of interest is isolated through IP, the RNA is reverse transcribed and amplified through PCR, then high-throughput sequencing is performed and reads are mapped to the genome (Licatalosi *et al.* 2008). In PAR-CLIP, cells are grown with a photoactivatable nucleoside (4-thiouridine or 5-bromouridine) in the media to facilitate crosslinking with proteins upon exposure to 365 nm radiation (Hafner *et al.* 2010).

Genome-wide structure data have recently been used to identify certain sites of RNA–protein interactions. The method icSHAPE was used to probe RNA structure in mouse embryonic stem cells *in vivo* and *in vitro* (Spitale *et al.* 2015). The difference in nucleotide reactivity *in vitro* and *in vivo* matched binding sites of the protein Rbfox2, previously identified with iCLIP experiments. This methodology was tested again and successfully identified RNA-binding sites of another RNA-binding protein, HuR. Using this type of analysis, certain RNA–protein interactions and associated RNA structural rearrangements can be distinguished using bioinformatics with experimental genome-wide mapping data.

2.2.3 Major advances: quantification of cellular factors *in vivo*

In vivo quantification of all the cellular factors known to affect RNA folding would both allow more accurate interpretation of *in vivo* RNA structure datasets and allow design of *in vivo-like* experiments that would more faithfully mimic *in vivo* conditions. Although such a comprehensive view of the inner workings of living cells has yet to be achieved, tremendous strides have been made in technique development for *in vivo* monitoring of cellular parameters relevant to RNA structure, including divalent ion concentrations, pH, reactive oxygen species (ROS), certain cosolutes, and RNA molecules themselves. Almost all of these techniques *in vivo* rely on a fluorescent readout, and thus advances in probe technology have gone hand-in-hand with advances in microscopy, although only the former topic is discussed here.

Fluorescent reporters are of three types: synthetic dyes, genetically encoded reporters, and reporters that incorporate both synthetic dyes and genetically encoded elements. Genetically encoded reporters typically rely on the cellular factor interacting with and altering the readout from a naturally fluorescent protein from jellyfish, green fluorescent protein (GFP), or its engineered variants (Tsien, 2010), the gene for which can be transformed into the system of interest. The ideal sensor will be minimally invasive and will have high specificity, brightness, and signal-to-noise ratios, a dynamic range that can accurately report the range of concentrations observed *in vivo*, and response kinetics that are as fast as the natural changes in the probed constituent. The best sensors are also ratiometric, which allows signal normalization to take into account such factors as photobleaching and heterogeneous dye distribution. It is important to note that the cellular environment differs among various cellular compartments and organelles. For example, the microenvironments of mitochondria (De Michele *et al.* 2014) and chloroplasts (Stael *et al.* 2011) (both of which have their own genomes and thus local RNA transcription) are quite different from the microenvironment of the nucleus, and both differ from the cytosolic environment. Ideally, a sensor would also have the capacity to be specifically targeted to an organelle or subcellular location where RNA-folding events of interest occur; for example, sensors that are genetically encoded can be fused to sequences that confer organelle-specific targeting (Choi *et al.* 2012).

Cations of particular relevance to RNA structure are heavy metals, which tend to destabilize and degrade RNA, Mg^{2+} , which tends to promote RNA folding, and H^+ (pH), which affects RNA catalysis. In addition, K^+ and Na^+ promote formation of the special RNA structure, the G-quadruplex. *In vivo* concentrations of Mg^{2+} (London, 1991; Lusk *et al.* 1968; Romani, 2007; Truong *et al.* 2013) and K^+ as well as pH changes are all within the concentration ranges that can affect RNA structure.



Among these cations, sensors based on GFP and its variants are available for Mg^{2+} (Lindenburg *et al.* 2013), Pb^{2+} (Nadarajan *et al.* 2014), Hg^{2+} (Hu *et al.* 2013), and H^+ (Tantama *et al.* 2011). A number of synthetic pH sensors are also available (Yang *et al.* 2014). Both genetically encoded and synthetic sensors of ROS are also available (Pouvreau, 2014; Swanson *et al.* 2011), which could be applied to study how ROS are associated with genetic diseases (Fimognari, 2015) or environmental conditions (Jaspers & Kangasjärvi, 2010) that affect RNA structure *in vivo*.

As discussed in Section 3.2.2, synthetic and biological cosolutes typically destabilize RNA structure. In one early report, sucrose, which is the circulating ‘energy currency’ in plants, was reported to destabilize RNAs *in vitro* (Gao *et al.* 2016; Lambert & Draper, 2007). While the *in vitro* effects occurred at significantly higher concentrations than prevail in the cytosol proper, in microdomains close to the sites of sugar transporters, sucrose, and other sugars could perhaps be present at significantly higher concentrations and consequently affect RNA structure locally; moreover, weakly folded RNAs, such as certain mRNAs, may be more susceptible to such cosolutes. In a possibly analogous situation, while resting Ca^{2+} levels in the cell cytosol are 100–200 nM, Ca^{2+} concentrations as high as 100 mM have been reported at the mouths of Ca^{2+} channels (Tang *et al.* 2015a). Lipid anchoring of recently developed sucrose and glucose sensors (Fehr *et al.* 2003; Lager *et al.* 2006) to probe the near membrane microenvironment of sugar transporters could allow evaluation of this hypothesis.

The physical microenvironment and the localization of RNA, both of which can impact RNA structure, vary across cellular regions and organelles. Accordingly, methods that allow visualization of the spatial location of any specific RNA of interest are also highly desirable (Buxbaum *et al.* 2015). One of the first technologies developed for RNA visualization was molecular beacons (Santangelo *et al.* 2006), which are oligonucleotides tagged with a synthetic fluorophore at one end and a synthetic quencher on the other end. Molecular beacons take on a non-fluorescent stem-loop structure in the absence of a complementary RNA due to the close proximity of the quencher and fluorophore, but exhibit fluorescence upon unfolding and hybridization to the target RNA. Various strategies (Santangelo *et al.* 2006) can be employed to introduce molecular beacons into mammalian cells, but they are not genetically encoded. A more widely used strategy for visualization of specific RNAs employs a genetic approach in which an RNA sequence that binds the bacteriophage MS2 protein is inserted into the UTR of the transcript of interest and the organism is engineered to express GFP-tagged MS2, which then binds to the transcript of interest, marking its location (Buxbaum *et al.* 2015).

A different type of RNA marker has been developed recently based on the GFP fluorophore. GFP is fluorescent because the folded protein immobilizes the 4-hydroxy-benzylidene-imidazolinone (HBI) fluorophore encoded by a cyclized and subsequently oxidized Ser–Tyr–Gly tripeptide. RNA aptamers have been identified that analogously immobilize and thus induce fluorescence of a related synthetic fluorophore, DFHBI [(Z)-4-(3,5-difluoro-4-hydroxybenzylidene)-1,2-dimethyl-1H-imidazol-5(4H)-one]. The sequence of the RNA aptamer is genetically incorporated into the gene of interest and upon RNA expression and administration of the membrane-permeant fluorophore and its immobilization by the RNA aptamer, fluorescence is observed that marks the location of the target RNA (Paige *et al.* 2011). The RNA aptamer, dubbed Spinach, as well as the second generation aptamer Spinach2, both require addition of exogenous Mg^{2+} to fold properly; such addition could obviously also affect native RNA structures. The third generation Spinach reporter, Broccoli, eliminates this requirement (You & Jaffrey, 2015).

Spinach aptamers can be further modified to read out concentrations of cellular metabolites by fusion of the Spinach aptamer with other aptamer sequences (identified by artificial selection) that selectively bind small molecules (Paige *et al.* 2012), or by incorporation of the Spinach aptamer into prokaryotic riboswitches (You *et al.* 2015). Riboswitch-based reporters have the advantage of having undergone natural selection that confers high affinity and specificity for the metabolite of interest, but are not currently ratiometric. Ratiometric sensors based on FRET between CFP and YFP, variants of GFP, have been engineered for several metabolites, including those with relevance to RNA structure. For example, FRET-based sensing of ATP concentration (Imamura *et al.* 2009) could be relevant to RNA structure because of the ATP requirement for the activity of RNA helicases (Rouskin *et al.* 2014). In summary, the future is bright for *in vivo* quantification of a plethora of the metabolites and physical properties that affect RNA structure. Quantification of cellular factors *in vivo* will play an important role in designing artificial cytoplasm to conduct *in vivo-like* studies of RNA folding.

2.2.4 Benefits and limitations of *in vivo* studies

Studies *in vivo* have shown that RNA can adopt different structures *in vivo* and *in vitro*, and have led to fresh insights on how the cellular environment affects RNA folding across a genome. Novel RNA structure motifs and RNA–protein interactions have been demonstrated through genome-wide *in vivo* experiments. In addition, novel RNA regulatory pathways have been identified by such studies.

Since some RNAs have been shown to fold and function differently under cellular conditions, the question arises, “Why not study RNA solely in living cells instead of in dilute solution conditions?” The reality is that methods for directly studying



RNA folding *in vivo* are limited, and most current *in vivo* approaches rely on structure probing methods that do not probe RNA thermodynamics or folding pathways. Experiments done *in vivo* provide information only on the average RNA structure in a cell or organism and lack information on RNA dynamics, the folding process, and the presence of multiple populated structures of the same transcript. These limitations motivate *in vivo-like* studies to understand the influence of cellular conditions on RNA folding. Before moving to the *in vivo-like* section, we consider the important role that *in silico* studies play in both *in vitro* and *in vivo* studies.

2.3 *In silico* studies of RNA folding

Studies *in vitro* and *in vivo* described above yielded insights into RNA folding and structure that were informed by *in silico* structure prediction tools. Structure probing experiments, for example, typically use *in silico* prediction tools to model structure that is guided by the experimental data. In the subsections below, we describe advances in predicting RNA structure from one sequence, from multiple sequences, and with experimental data. Limitations of each approach are provided as well.

2.3.1 Major advances: RNA structure prediction from one sequence *in silico*

The most popular approaches to predict RNA structure use dynamic programming algorithms to efficiently search the set of possible structures (Eddy, 2004) and folding free energy nearest-neighbor rules to estimate folding stability (Turner & Mathews, 2010). The dynamic programming algorithms guarantee that every structure allowed by the set of folding rules is considered, except for those containing pseudoknots (see below). This means, for example, that the lowest free energy conformation will be found for programs that find lowest free energy structures, i.e. the most probable structure at equilibrium.

The accuracy of RNA structure prediction from sequence alone, in terms of fraction of known pairs correctly predicted, is stubbornly limited to ~70% (Hajiaghayi *et al.* 2012; Lu *et al.* 2009), and accuracy is lower for long sequences (>1000 nucleobases) such as small and large ribosomal RNAs and mRNAs (Doshi *et al.* 2004) or for sequences that fold to more than one conformation at equilibrium. *In silico* predictions of base pairs presently rely on a parameterization of stabilities determined *in vitro* rather than *in vivo*, and these parameters are based on relatively few experiments, as compared to all possible folded sequences.

In response to this moderate success rate, a number of *in silico* methods have been developed to predict alternative structures, as reviewed previously (Mathews, 2006). Programs generate sets of alternative hypotheses for the structure (suboptimal structures) (Wuchty *et al.* 1999; Zuker, 1989), feasible structures in equilibrium with each other (stochastic samples) (Ding & Lawrence, 2003), or estimates for base pairing probabilities (partition function calculations) (McCaskill, 1990). Each of these three methods is described in turn. Suboptimal structures are those with similar free energy to the lowest free energy structure. Certain suboptimal structures can sometimes be more representative of the biological structure than the *in silico*-estimated lowest free energy structure, and can be viewed as alternative models or alternative hypotheses for the *in vivo* structure. Stochastic samples are rigorous samples from the equilibrium (Boltzmann) ensemble. They are useful for estimating ensemble statistics for the secondary structure of an RNA. Partition function calculations provide pairing probability estimates; more probable pairs in predicted structures are more likely to occur in the accepted structure (Mathews, 2004).

2.3.2 Major advances: RNA structure prediction from multiple sequences *in silico*

The accuracy of *in silico* folding can be dramatically improved by using additional information to guide the folding. In this section, we discuss using homologous sequences to guide the folding, while in the next section we discuss applying experimental data. Multiple homologous sequences, commonly called an RNA family, can be used to estimate the common secondary structure (Seetin & Mathews, 2012a) because structure is generally conserved to a greater extent than sequence for RNAs. Due to sequence variation, the number of base pairs conserved across a family is smaller than the number of base pairs adopted by each sequence. With enough sequences, conserved pairs stand out as positions of covariation, where compensating base pair changes are observed. Covariation is a change in sequences where one biological species, for example, will have an AU base pair, but another species will have a GC pair at the homologous position. During evolution, two separate changes occurred in sequence (a compensating change) that conserved the base pair.

Three approaches are used to estimate the biologically conserved structure from a set of homologous sequences (Reeder *et al.* 2006; Seetin & Mathews, 2012a). In the first approach, the available sequences are aligned, and then used to restrain the *in silico* prediction. This approach is typically the fastest, but generally works best when the pairwise sequence identity of all the homologs is high (75% or higher). These programs are exemplified by RNAalifold (Bernhart *et al.* 2008) and TurboFold (Harmanci *et al.* 2011). Programs in the second set predict the structures for each sequence first and then compare the predicted structures to find those common to all sequences. This approach works well when the structure is highly



conserved and is exemplified by RNACast (Reeder & Giegerich, 2005). The third approach is to simultaneously align and fold sequences to find the common structure and sequence alignment. This is the best approach to use when the sequences are diverse (pairwise sequence identity for some sequence pairs below 75%) because low pairwise identity makes sequence alignment challenging. Programs in this class include Dynalign/Multilign (Fu *et al.* 2014; Xu & Mathews, 2011), Foldalign (Torarinsson *et al.* 2007), LocARNA (Will *et al.* 2007), PARTS (Harmanci *et al.* 2008), and RAF (Do *et al.* 2008).

The accuracy of *in silico* prediction of conserved structures from a set of homologous sequences can be much higher, than for predictions from single sequences. For example, often an additional 20% or more of the known base pairs can be correctly predicted using multiple homologs as compared to predictions using a single sequence (Xu & Mathews, 2011). For a given set of sequences, however, it is not always obvious which approach or program to use, and, therefore, it is probably best to try more than one program to develop hypotheses about the *in vivo* structure. To date, no program can completely automate comparative sequence analysis. Manual comparison is still required for the most accurate RNA secondary structure determination.

2.3.3 Major advances: RNA structure prediction *in silico* restrained with experimental data

Another type of information used to guide *in silico* prediction of RNA structure is experimental structure mapping. Such mapping data can come from *in vitro* or *in vivo* experiments and are used to restrain structure prediction (Lorenz *et al.* 2016; Sloma & Mathews, 2015). The effects of experimental structure restraints have been well studied using *in vitro* probing data on structured ncRNAs. Over 85% of known pairs can be correctly predicted using *in vitro* SHAPE, DMS, or enzymatic cleavage data (Cordero *et al.* 2012; Deigan *et al.* 2009; Eddy, 2014; Hajdin *et al.* 2013; Ouyang *et al.* 2013; Washietl *et al.* 2012; Wu *et al.* 2015; Zarrinhalam *et al.* 2012) when the extent of accessibility is quantified using capillary/gel electrophoresis or deep sequencing counts. This is a dramatic improvement over the above-mentioned 70% limit in the absence of mapping data. Using *in vivo* mapping data to improve the accuracy of structure prediction has not yet been well studied, although mapping data overlaid on known structures suggests that, for structured ncRNAs such as rRNAs, the existing methods should improve structure prediction accuracy (Ding *et al.* 2014). We recently developed a pipeline called StructureFold to fold RNAs across a genome using restraints from experimental data, which works with Structure-seq data (Tang *et al.* 2015b), and the RNAstructure program and can accommodate other data and folding algorithms.

2.3.4 Challenges with *in silico* modeling of RNA secondary structure

Despite its widespread use, RNA secondary structure prediction has known limitations. First, the nearest-neighbor parameters are based on a limited number of experiments measured *in vitro* in 1 M NaCl rather than *in vivo-like* conditions, and there are probably many sequences that are not predicted well with those parameters (Andronescu *et al.* 2014; Mathews *et al.* 2004). On the one hand, for a limited number of simple RNAs melted in physiological K^+ and Mg^{2+} concentrations, the stability is often similar to that in 1 M NaCl (Diamond *et al.* 2001; Jaeger *et al.* 1990; Jiang *et al.* 2014; Schroeder & Turner, 2000). However, for a 5S ribosomal RNA loop E motif, for example, an appreciable difference in stabilities was found between buffers with and without Mg^{2+} (Serra *et al.* 2002). Second, although enthalpy parameters are available for structure prediction between 10 and 60 °C (Lu *et al.* 2006), predictions are generally made at 37 °C, which is relevant to humans, but not the majority of organisms. Third, finding lowest free energy structures assumes that RNAs fold to equilibrium, i.e. kinetics do not control folding. In favor of this assumption, an *in vivo* study of ribozymes suggested that RNAs fold to equilibrium to a greater extent in yeast cells than *in vitro* (Mahen *et al.* 2005). Also, *in vitro* structure mapping studies of annealed ribosomal RNAs were consistent with *in vivo* structures (Moazed *et al.* 1986b). However, some sequences are kinetically trapped, such as transcriptional riboswitches (Seetin & Mathews, 2012a; Wickiser *et al.* 2005a; Wickiser *et al.* 2005b). Therefore, it is unclear to what extent factors such as non-physiological ionic conditions and cotranscriptional folding play roles in shaping the folding of RNA.

A fourth limitation of the most popular programs for *in silico* folding is that they cannot predict pseudoknots (Liu *et al.* 2010a). A pseudoknot occurs when there are base pairs between nucleotides in two different loops. Formally, a pseudoknot is composed of two or more base pairs, defined by indices i base paired to j and i' base paired to j' , where the order of the nucleotides is $i < i' < j < j'$. Pseudoknotted pairs are a small fraction of total base pairs in known structures but often occur in highly structured and functional RNAs. For programs that predict pseudoknots, the accuracy is shockingly low (<5%) (Bellaousov & Mathews, 2010), although the use of multiple homologous sequences to identify conserved pseudoknots improves the accuracy (Seetin & Mathews, 2012b). Recently, it was also shown that *in vitro* SHAPE mapping data can guide *in silico* structure prediction, including pseudoknots, and achieve over 90% accuracy at predicting known base pairs (Hajdin *et al.* 2013). The program that implements this, ShapeKnots, is limited, however, to sequences of 600 nucleotides or fewer.



Although structure mapping data and sequence comparison are each used to guide *in silico* modeling of RNA secondary structure, little has been done until recently to combine the two approaches for additional synergy. The secondary structures of three long ncRNAs were modeled with the aid of structure mapping data: HOTAIR (with *in vitro* SHAPE, DMS, and terbium) (Somarowthu *et al.* 2015), SRA (with *in vitro* SHAPE, DMS, in-line probing, and RNase V1 digestion) (Novikova *et al.* 2012), and XIST (with *in vivo* DMS mapping) (Fang *et al.* 2015). For each of these studies, sequence comparison, i.e. the verification that the structures are conserved and the identification of compensating base pair changes, was subsequently used to further support the structure model.

Two software programs were enhanced to combine structure mapping data and sequence comparison to improve structure prediction. Sükösd *et al.* (2012) reported PPfold, a program that uses a probabilistic approach to predict structure and can be guided by SHAPE mapping data and/or sequence covariation as estimated from a sequence alignment. Recently, SHAPE data were used to inform sequence alignment and then RNAalifold to predict the conserved structure for the aligned sequences (Lavender *et al.* 2015b). The key observation is that homologous nucleotides, i.e. those that align, have similar SHAPE reactivities and thus the differences in SHAPE reactivity can be included as an additional metric in the scoring of alignments. This approach demonstrated an improved accuracy of base pair prediction by RNAalifold as compared to consensus structure prediction or SHAPE guided structure prediction alone. Both of these approaches were used to model HIV RNA structure using mapping data and sequence comparison (Lavender *et al.* 2015a; Sükösd *et al.* 2015).

3. Bridging the gap between *in vitro* and *in vivo* RNA folding using *in vivo*-like studies

3.1 The gap

The previous sections outlined major contributions of RNA-folding studies *in vitro* and *in vivo* to our understanding of how RNA behaves, while considering the important roles that *in silico* approaches play. *In vitro* studies provide the fundamentals of RNA thermodynamics and kinetics, RNA structural motifs, and genome-wide RNA structure trends. *In vivo* structure probing methods reveal RNA structural trends related to biological functions and regulatory roles of RNA genome-wide. We discussed how several research teams have used genome-wide *in vivo* structural probing to uncover that, in general, RNAs do not adopt the same structures *in vivo* as *in vitro*. Since structure generally dictates function, understanding differences between RNA folding *in vivo* and *in vitro* can illuminate biological function. Toward accomplishing this goal, RNA folding and function studies have been increasingly conducted under conditions that *mimic* the cellular environment.

The dilute solution conditions traditionally used to study RNA *in vitro* are vastly different from the cellular environment. The cellular environment is a complex solution containing biopolymers, metabolites, dilute free salts, and organelles, with 20–40% of the cellular volume occupied by macromolecular crowders (Minton, 2001; Zimmerman & Trach, 1991). As such, there is no single cellular environment to which RNA is exposed. As an mRNA passes from the nucleus to the cytosol, solution conditions change; in eukaryotes, the cell is compartmentalized and as the RNA is transported to different regions its fold can change.

It is of interest to consider the differences between RNA structure in eukaryotic and prokaryotic organisms. Functional RNAs have intricate structures with tertiary contacts that assemble secondary structures close in space. Cations, typically Mg^{2+} , neutralize the negative charge of the phosphate backbone and promote tertiary structures. Free Mg^{2+} concentrations in prokaryotic and eukaryotic cells are different, ~1.5–3.0 and 0.5–1.0 mM, respectively (London, 1991; Lusk *et al.* 1968; Romani, 2007; Truong *et al.* 2013). Structured RNAs such as ribozymes, riboswitches, and thermosensors are found frequently in prokaryotes, where free Mg^{2+} levels are higher. Although a few ribozymes and one riboswitch have been identified in eukaryotes, they appear to be rare, and proteins are typically involved in forming requisite tertiary structures (Kubodera *et al.* 2003; Roth *et al.* 2014; Salehi-Ashtiani *et al.* 2006). Lambowitz and co-workers demonstrated that prokaryotic group II introns fold poorly in eukaryotic cells, although they could select variant RNAs that fold into active conformations at eukaryotic low Mg^{2+} concentrations (Truong *et al.* 2013). Studies in our laboratory indicate that the eukaryotic innate immune sensor PKR is activated by prokaryotic RNAs under eukaryotic low Mg^{2+} conditions, leading to the speculation that riboswitches and ribozymes may be selected against in eukaryotes to aid in discriminating self and non-self at the RNA level (Hull & Bevilacqua, 2015, 2016; Hull *et al.* 2016). To date, there are no studies that compare the structures of eukaryotic and prokaryotic RNAs genome-wide, but such information would be valuable.

Historically, *in vitro* experiments lack many of the components of cellular environments, and, moreover, often have high concentrations of salt to fold RNA for thermodynamic and structural studies (Table 1). Thermodynamic studies cannot, however, readily be performed *in vivo*. The cell prohibits wide variations of temperature, pH, salt, and ligand concentration, all of which are necessary to obtain thermodynamic information. As a result, RNA is being increasingly studied in artificial cytoplasm



that mimic aspects of the cellular environment while allowing biophysical studies. Several recent studies focused on mimicking aspects of the *in vivo* environment *in vitro*; conditions referred to herein as ‘*in vivo-like*’ conditions (Fig. 3). Effects of such conditions as cellular concentrations of monovalent and divalent ions and molecular crowding agents on the folding of RNAs have been a theme in a number of recent studies (Desai *et al.* 2014; Dupuis *et al.* 2014; Nakano *et al.* 2015; Paudel & Rueda, 2014; Strulson *et al.* 2014; Tyrrell *et al.* 2015). Experiments under these *in vivo-like* conditions have the potential to bridge our understanding of observations made *in vitro* and *in vivo*.

3.2 Design of artificial cytoplasm and early experiments

In this section, we discuss various methods of mimicking cytoplasmic conditions, including the use of polymers and cosolutes as crowding agents and the use of protocells and synthetic membranes. We also discuss the outcomes of early experiments under these *in vivo-like* conditions. Finally, directions in which the field needs to move to understand the fold and function of RNA *in vivo* are suggested.

3.2.1 Polymers

Synthetic crowding agents such as polyethylene glycol, dextran, and ficoll, and small cosolute additives such as methanol, proline, and trimethylamine oxide (TMAO) have been used to mimic the crowded environment of the living cell. Functional RNAs that are well studied *in vitro* have been used to test the effects crowding agents have on RNA folding. Various methods, including UV melts, SAXS, kinetic techniques, and smFRET, have been used to study RNA under these *in vivo-like* conditions. Several studies have shown that synthetic crowding agents affect the thermodynamics and function of several RNAs (Dupuis *et al.* 2014; Kilburn *et al.* 2010, 2013; Lambert *et al.* 2010; Strulson *et al.* 2014). Findings of these studies are that RNAs fold cooperatively, structure becomes compact, and ribozymes cleave faster under *in vivo-like* conditions (Kilburn *et al.* 2013; Nakano *et al.* 2009; Strulson *et al.* 2013, 2014).

The kinetics of several small and large ribozymes have been probed under *in vivo-like* conditions and in all reported cases, rates of catalysis have increased in the presence of molecular crowders as compared to dilute solution conditions (Desai *et al.* 2014; Nakano *et al.* 2009; Paudel & Rueda, 2014; Strulson *et al.* 2012, 2013). For example, the hammerhead ribozyme has higher catalytic activity, between 3.5 and 6.5 faster than in dilute solutions, in the presence of 10–30% (wt %) PEG200 or PEG8000, suggesting a more populated active state in crowded conditions (Nakano *et al.* 2009). In addition, *in vivo-like* solution conditions can stabilize ribozymes even in the presence of denaturants. For example, the rate of catalysis of the CPEB3 ribozyme in the presence of 2.5 M of the denaturant urea was recovered by the addition of 30% (w/v) PEG200, PEG8000, or Dextran10, at a rate higher than in buffer alone (Strulson *et al.* 2013). SAXS experiments have provided insight into the structural basis for enhanced catalysis, showing that the natively folded state adopts a more compact structure in the presence of molecular crowders under conditions of biological Mg^{2+} concentrations (Kilburn *et al.* 2010; Strulson *et al.* 2013).

The thermal stability of several functional RNAs has been reported to increase under *in vivo-like* conditions as compared with *in vitro* experiments. For instance, in 20% PEG200 or PEG8000 the hammerhead ribozyme retains catalytic activity up to 60 °C, a temperature that thermally denatures the ribozyme in dilute solutions (Nakano *et al.* 2009). Observation of increased hammerhead catalytic activity, up to 270-fold, at high temperatures in crowded conditions indicates a more thermostable RNA under *in vivo-like* conditions. Interestingly, the individual secondary structure elements of the ribozyme were observed, through optical melting experiments, to be thermally destabilized in molecular crowding agents, suggesting that tertiary structure is stabilized and resulting in more cooperative folding of the ribozyme (Nakano *et al.* 2009). A thermodynamic study from our laboratory using SHAPE structure probing on tRNA^{phe} under *in vivo-like* conditions showed that tRNA folds in a cooperative manner at biological Mg^{2+} concentrations in the presence of molecular crowding (Strulson *et al.* 2014). The observed increase in folding cooperativity with crowding was accompanied by an increase in the temperature of the melting transition for tertiary structure. When the tertiary interactions were removed by mutation of nucleotides in tertiary contacts to uridine, cooperativity was lost and the RNA folded with multiple transitions under all solution conditions, thus indicating that tertiary interactions are vital to cooperative RNA folding under *in vivo-like* conditions. This effect is similar to that observed under *in vitro* conditions mentioned above (Chauhan & Woodson, 2008).

The contribution of molecular crowding agents to RNA catalysis and folding has been found to be largest in a background of physiologically low ionic conditions rather than high ionic conditions (Kilburn *et al.* 2013; Strulson *et al.* 2013). In the absence of crowding, physiological concentrations of Mg^{2+} are not high enough to fold functional RNAs in a two-state manner. This is apparent from the observation of long-lived intermediates and slow folding under these conditions (Banerjee & Turner, 1995; Chadalavada *et al.* 2002; Mitchell *et al.* 2013). However, in the presence of biological crowding conditions and physiological Mg^{2+} , functional RNAs tend to fold in a cooperative manner into compact structures (Desai *et al.* 2014; Dupuis *et al.* 2014; Strulson *et al.* 2014; Tyrrell *et al.* 2015), and ribozymes and riboswitches tend to have higher rates of cleavage and higher

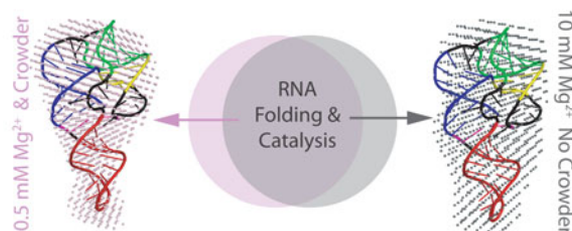


Fig. 7. Under both *in vitro* (right, grey) and *in vivo-like* conditions with molecular crowding (left, pink) RNA fold into their native state that is functional, indicated in this figure by catalysis. High concentrations of Mg^{2+} (10 mM or higher) are needed to achieve the folded state *in vitro* compared with *in vivo-like* crowded conditions where low physiological Mg^{2+} (0.5 mM) folds the RNA. Reprinted with permission from Strulson *et al.* (2013). Copyright 2016 American Chemical Society.

ligand-binding affinity (Paudel & Rueda, 2014). The addition of more Mg^{2+} to these conditions does not result in a further increase in the rate of activity or more cooperative RNA folding, indicating that together physiological crowding and Mg^{2+} conditions fold RNA optimally (Fig. 7).

A recent study explored the structural effects of the molecular crowding agent PEG (ranging in size from the monomer to 35 000 kDa) on the adenine riboswitch (Tyrrell *et al.* 2015). Using SHAPE chemistry, the reactivity of the riboswitch under *in vitro*, *in vivo*, and *in vivo-like* conditions was explored. The authors found that in low molecular weight PEG (<3350 kDa) the riboswitch had low correlation between reactivity *in vivo* and *in vivo-like* conditions, whereas in higher molecular weight PEG (12 000–35 000 kDa) the RNA had a similar reactivity under *in vivo* and *in vivo-like* conditions. While this study was limited to a single molecular crowding agent, it is significant because it showed that certain *in vivo-like* conditions are not accurate cellular mimics.

Recently, the folding of a model RNA was examined *in vivo*. The *Salmonella* fourU RNA thermometer hairpin containing a FRET pair was injected into live mammalian cells and reported to have similar melting temperatures and unfolding free energy *in vivo* and *in vitro* (Gao *et al.* 2016). The addition of 30% (w/v) PEG of varying sizes and Ficoll70 was shown to modify the thermodynamics of the hairpin, and higher molecular weight polymers were found to have similar effects on the RNA as the *in vivo* environment. The *in vivo* data had a very broad distribution of melting temperatures and free energy between both different cells and different cellular compartments, leading to some uncertainty about how the cellular environment is affecting RNA folding.

3.2.2 Cosolutes

While molecular crowding agents generally facilitate the folding of functional RNAs, small cosolutes have varying and complicated effects on RNA thermostability and folding cooperativity. This arises in part because the effect on stability depends strongly on the interactions between the particular cosolute and RNA considered. Cosolutes, also known as osmolytes, regulate osmotic pressure in cells (Record *et al.* 1998; Yancey *et al.* 1982). Effects of cosolutes on RNA folding were not significantly investigated until the last decade. Studies on RNAs with either secondary and/or tertiary structures report that cosolutes such as betaine, proline, and methanol, almost always destabilize secondary structures, while having mixed effects on tertiary structure (Lambert & Draper, 2007, 2012; Lambert *et al.* 2010; Soto *et al.* 2007). Several osmolytes have been shown to interact with the nucleobase, sugar, and phosphate of RNAs, with examples of both favorable and unfavorable interactions (Lambert & Draper, 2007). Stabilizing osmolytes have unfavorable interactions with the unfolded state of RNA, resulting in RNA compaction that buries functional groups and stabilization of the native state, while destabilizing osmolytes have favorable interactions with the unfolded state of RNA, driving unfolding (Holmstrom *et al.* 2015; Lambert & Draper, 2007; Lambert *et al.* 2010).

There are a limited number of studies available for the effect of cosolutes on RNA function. The hammerhead ribozyme was shown to have increased rates of cleavage in 20% cosolutes, such as glycerol and 1,2-dimethoxyethane, in the presence of physiological Mg^{2+} , which was attributed to enhanced electrostatic interactions with Mg^{2+} (Nakano *et al.* 2015). The secondary and tertiary structures of the hammerhead ribozyme were destabilized in the presence of several cosolutes (Nakano *et al.* 2009). In crowded conditions, ribozyme activity also increased, while secondary structure was destabilized and tertiary structure was stabilized (Nakano *et al.* 2009).

The influence of the cosolute TMAO on RNA secondary and tertiary structure, as well as on the phosphate backbone, has been studied (Denning *et al.* 2013; Lambert & Draper, 2007; Lambert *et al.* 2010). TMAO is unusual in having almost no effect on secondary structure stability, while generally stabilizing tertiary structure. A small 58 mer rRNA was found to exhibit



cooperative two-state folding in the presence of TMAO, observed by a single transition in an optical melting experiment (Lambert *et al.* 2010).

3.2.3 Protocells and synthetic membranes

There are several groups focusing on how to model RNA function and structure in early Earth conditions, which also relate to compartmentalization in modern cells. Coacervates and synthetic membranes are often used to mimic early Earth protocells, and RNA function in these protocells is often studied through ribozyme cleavage. Our laboratory studied the activity of a two-piece hammerhead ribozyme in aqueous two-phase systems (ATPS) made of polyethylene glycol and dextran (Strulson *et al.* 2012). The system forms a dextran-rich phase droplet in which the ribozyme preferentially localizes at a concentration up to 3000 times that of the aqueous phase, resulting in a 70-fold increase in the rate of catalysis. This study suggested that RNA catalysis in the early Earth environment could have arisen from compartmentalization increasing the local concentration of RNA, possibly accelerating very slow reactions, so that they could occur on a biologically relevant timescale.

Similar to these droplets, mononucleotides will form microdroplets when mixed with cationic peptides in water (Koga *et al.* 2011). Inside these droplets, nucleotides and peptides can reach concentrations as high as 1.6 M and 400 mM respectively, which is much more concentrated than in the aqueous phase. Cationic and anionic dyes and certain nanoparticles were shown to partition into the droplets, indicating that the droplets are permeable to charged molecules (Koga *et al.* 2011). These droplet phases are another indicator that early life could have arisen in non-membranous compartments. More recently we made coacervates from nucleotides and poly(allylamine) that contain molar concentrations of Mg^{2+} and nucleotides, which could facilitate RNA catalysis in an early life scenario (Frankel *et al.* 2016).

4. Future directions

The majority of what is known about RNA folding and structure comes from studies that were performed *in vitro* on small model systems and highly structured RNAs. In contrast, little is known about how RNA folds and functions *in vivo*. Current *in vivo* methods probe the RNA structure ensemble. While providing a benchmark for new prediction parameters, ensemble methods cannot themselves generate thermodynamic parameters.

The current thermodynamic parameters for RNA structure prediction were established in 1 M NaCl. However, several-transcript-specific and genome-wide studies have shown that certain RNAs do not fold into the same structures *in vivo* and *in vitro* (Kwok *et al.* 2013; Rouskin *et al.* 2014; Tyrrell *et al.* 2013, 2015), so improved software and prediction parameters are needed to model *in vivo* structure. In particular, genome-wide *in vivo* structure probing datasets (Ding *et al.* 2014; Rouskin *et al.* 2014) contain a wealth of information that has not yet been completely realized or understood. One barrier to taking full advantage of the data is that most of the available *in silico* methods assume that RNAs fold to a single structure (Cordero *et al.* 2012; Deigan *et al.* 2009; Hajdin *et al.* 2013; Ouyang *et al.* 2013; Wu *et al.* 2015), while probing data averages across all structures populated by sequences for the duration of the experiment.

Modeling a single structure works well for ncRNA sequences that function with a single structure, such as ribosomal RNAs, but there are many RNAs for which this assumption is not correct, such as RNA switches and open reading frames. A key challenge is developing methods to use the probing data to model ensembles of relevant structures. Three recent papers highlight work to address this challenge. Cordero and Das report an *in silico* method (M^2 -REEFFIT) that models complex mixtures of multiple structures, aided by *in vitro* SHAPE mapping of the wild-type sequence and also a set of mutant sequences, which reveal nucleotide interactions (Cordero & Das, 2015). Multiple structures for the 5' UTR of an mRNA were modeled using *in vitro* SHAPE mapping of a mixture of structures (Kutchko *et al.* 2015). The multiple conformations were modeled *in silico* using stochastic sampling, restrained using the standard SHAPE restraints expressed as free energy terms (Deigan *et al.* 2009). A third *in vitro* approach separated multiple conformations of HIV RNA using native gel electrophoresis, and mapped the structures with SHAPE in the gel (Sherpa *et al.* 2015). This simplified the *in silico* analysis because the SHAPE mapping data were acquired for each conformation independently.

Another key challenge for mapping studies is determining the best way to discover or model interactions of RNAs with proteins or other RNAs. *In vivo*, all RNAs can interact with macromolecules and metabolites. These interactions generally result in protection from probing agents. Deconvoluting *in silico* whether a nucleotide is unreactive because of intramolecular structure or intermolecular interactions is a grand challenge that will likely require new types of experimental information to address. Modeling and predicting 3D RNA structures *in vitro* is an ongoing challenge. A recent RNA puzzle tested blind 3D folding predictions by providing research teams with RNA sequences and chemical probing data for those RNAs (Miao *et al.* 2015). The structures that the teams modeled were compared with crystal structures, and most teams could predict



Watson–Crick base pairs, but struggled in predicting non-canonical WC base pairing and stacking interactions. A long range of goal is to predict relevant RNA 3D structures *in vivo* to understand the biologically relevant confirmation(s).

The study of RNA under *in vivo-like* conditions is relatively young. To better mimic the cellular environment, more complex cytoplasm mimics should be developed. To date, artificial cytoplasms have focused on synthetic polymers and cosolutes, but more accurate ionic conditions, biopolymers and even cell extracts need to be applied. In addition, studies under *in vivo-like* conditions have focused on single transcripts in synthetic crowding and cosolute conditions. Genome-wide comparisons of RNA folding under *in vivo* and *in vivo-like* conditions are needed. Lastly, methods that can probe the thermodynamics and kinetics of RNA folding under complex *in vivo-like* conditions will enhance our understanding of *in vivo* RNA folding. Overcoming the challenges outlined herein will allow the field to accomplish the ultimate goal, to understand how RNA folds in the cell.

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