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Functional 5' UTR mRNA structures in eukaryotic translation regulation and how to find them

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Abstract

RNA molecules can fold into intricate shapes that can provide an additional layer of control of gene expression beyond that of their sequence. In this Review, we discuss the current mechanistic understanding of structures in 5' untranslated regions (UTRs) of eukaryotic mRNAs and the emerging methodologies used to explore them. These structures may regulate cap-dependent translation initiation through helicase-mediated remodelling of RNA structures and higher-order RNA interactions, as well as cap-independent translation initiation through internal ribosome entry sites (IRESs), mRNA modifications and other specialized translation pathways. We discuss known 5' UTR RNA structures and how new structure probing technologies coupled with prospective validation, particularly compensatory mutagenesis, are likely to identify classes of structured RNA elements that shape post-transcriptional control of gene expression and the development of multicellular organisms.

In the ancient RNA world, RNA likely served as the main catalytic, self-replicating and information-carrying component pre-dating cellular life^{1,2}. Such intricate activities can be traced to the ability of RNA to fold into complex secondary structures and tertiary structures. In many cases, these RNA structures appear more dynamic than globular protein domains or the double-helical DNA structure. Primarily in bacteria and archaea but also in algae, fungi and plants, prominent examples of RNA structure-directed functions include ribozymes and metabolite-sensing riboswitches^{3,4}. The roles of RNA structures in splicing and in gene regulation by non-coding RNAs may have been a driving force in the evolution of ancient eukaryotes^{5,6}. Most strikingly, the peptidyl transferase centre of ribosomal RNA (rRNA) is a structured remnant of the RNA world that is central to protein synthesis in all living cells.

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The authors declare no competing interests.

SUPPLEMENTARY INFORMATION

See online article: S1 (box) | S2 (box) | S3 (table)

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We focus here on mRNA structures with functions in translation initiation, which is ultimately linked to the ribosome — the macromolecular complex that harbours this peptidyl transferase activity. The great number of protein and RNA components involved in ribosome initiation, scanning, elongation and recycling of mRNAs highlights that translation — especially translation initiation, which is one of the most crucial steps in translation^{7,8} — is a highly regulated process (BOX 1). Indeed, the ribosome itself, which for a long time was regarded as a constitutive, housekeeping molecular machine, has only recently been appreciated to be functionally heterogeneous with respect to its associated proteins^{9,10}. Translation regulation can also involve structures in the mRNA itself, which are rearranged and unfolded by the ribosome and by RNA remodellers such as RNA helicases. This complex and dynamic interplay of mRNA structures and the translation machinery raises an important question: how much regulatory potential is encoded in mRNA structures or in the structure-mediated sensing and recruitment of interacting factors such as RNA-binding proteins (RBPs) or *trans*-acting RNAs^{11–13}?

One of the great challenges in answering this question has been to accurately find mRNA structures that modulate translation. Although mRNA primarily transmits genetic information from DNA to protein through its coding sequence (CDS), the 5′ and 3′ untranslated regions (UTRs) are non-coding and do not directly contribute to the protein sequence. Free from the constraints of encoding proteins, UTRs can form considerable Watson–Crick and non-canonical base pairing that can potentially impact every step of translation. Indeed, algorithms for modelling RNA secondary structure^{14,15} have suggested that UTRs have the potential to engage in intricate RNA base-pairing patterns, which may change in response to protein binding and may impact the recruitment of ribosomes. Although these algorithms have had difficulty in processing long RNA sequences and predicting long-range interactions or complex structures such as pseudoknots and still fail to account for protein interactions, the past decade has witnessed a burst of advances in integrating these methods with experimental RNA structure probing methodologies. Such integration raises the prospect of determining the complete ensembles of folding states of all transcribed RNAs in a given cell type at a given time and of understanding how these structural ensembles may give rise to intricate gene regulation programmes. Expected improvements of these methods would enable precise validation of functional mRNA structures inside cells.

In this Review, we highlight recent research on the roles of functional 5′ UTR structures in eukaryotic mRNAs as modulators of translation initiation. However, it is important to note that regulatory elements elsewhere in the mRNA, especially in the 3′ UTR, can also modulate translation¹⁶. It is also important to mention that even within 5′ UTRs, unstructured (linear) regulatory elements are likely to have a crucial impact on translation^{17–21}. Linear elements include upstream open reading frames (uORFs)^{22–24} and the sequences around the start codon of the main ORF. Such sequence elements have been dissected and discussed elsewhere^{17,21,25}. As a classic example, a strong Kozak sequence²⁶ improves start codon recognition as a feature of highly translated mRNAs. Analysis of the relative strength of AUG codon recognition of all possible translation initiation sites has recently revealed specific responsible sequence motifs around start sites in mammals²⁷.

In this Review, we focus on how 5' UTR structures in mRNAs may block or recruit ribosomes and other regulatory factors to enable a rapid, dynamic response to diverse cellular conditions to control gene expression. We speculate that such structured elements may be particularly important when newly induced responses through mRNA transcription, processing and nuclear export may be too slow²⁸. Instead, the cell may reversibly change its expression profile by adjusting the stability and translation of pre-existing mRNAs in the cytoplasm. For example, activated macrophages derepress the translation of 90 mRNAs — many of which encode anti-inflammatory regulators — within 1 hour of immune stimulation²⁹. We also highlight how recent technological advances are improving our understanding of RNA structure by chemical probing of RNA conformation *in vitro*, and especially now inside cells, *in vivo*. We anticipate that these methods will be crucial for the elucidation of yet unknown structured RNA elements that guide translation control.

5' UTRs as platforms for RNA structure

During evolution from invertebrates to humans, genome size greatly increased, and UTRs have especially expanded in length^{30,31}, providing a 'playground for mRNA evolution'. As UTRs are usually not coated with translating ribosomes, they are presumably more accessible for interactions with regulatory factors. Whereas the length of the 3' UTR immensely increased during eukaryotic evolution, the 5' UTR has maintained a median length of approximately 53–218 nucleotides^{17,19,32,33}. The longest known median length of mRNA 5' UTRs occurs in humans (218 nucleotides)^{30,31}, exceeding those of other mammals and dwarfing that of budding yeast (53 nucleotides) (FIG. 1). However, 5' UTR lengths vary dramatically among individual genes in higher eukaryotes and can range from a few to thousands of base pairs^{32,34} (FIG. 1). This large range of 5' UTR lengths suggests that there may be greater regulation of specific mRNA subsets.

High GC content and a highly negative folding free energy (ΔG) level of a 5' UTR are often used as parameters for predicting 5' UTR RNA secondary structures. As canonical translation initiation requires that the 43S pre-initiation complex scans the 5' UTR to reach the start codon (BOX 1), such overall high GC content in the 5' UTR has been thought to cause inefficient scanning and a lower rate of initiation. Indeed, in classic examples, the prediction of complex secondary structures in GC-rich 5' UTRs has been correlated with inhibition of translation³⁵, for example, in the mRNA of the metabolic enzyme ornithine decarboxylase³⁶. However, these secondary structure predictions and particularly their functional relevance in cells have not been established. Predictions typically calculate the most stable base pairing of an RNA as the one that has the overall lowest computed ΔG . Although determining the ΔG of an entire RNA structure in an mRNA takes into account the transition of an RNA domain from its fully folded to a completely linear form, scanning is thought to require only local melting of RNA structures rather than linearization of the whole 5' UTR. Indeed, emerging data indicate that strong local RNA structures and protein interactions may have important roles in impeding ribosome scanning³⁷. Given the limitations of predicting global 5' UTR mRNA structures in guiding the discovery of RNA structures, the search for individual functional mRNA structures may be more promising. In the next sections, we take a closer look at interesting candidate 5' UTR RNA structure motifs that regulate mRNA translation and at the methods used to find and confirm them.

5' UTR structures in ribosome scanning

Translation in eukaryotes typically starts at the 5' end of the mRNA, which harbours the 5' cap and a UTR as the entry point for the ribosome (BOX 1; FIG. 2a). Some mRNAs lack a 5' UTR completely, for example, all mRNA species in mammalian mitochondria are leaderless³⁸, but they are generally rare in higher eukaryotes¹⁹. Some human mRNAs with an extremely short 5' UTR (12 nucleotides on average), known as translation initiator of short 5' UTR (TISU), undergo scanning-free initiation³⁹. By contrast, some 5' UTRs are highly structured and can block entry of the ribosome. One of the first and best-studied examples is a small 5' UTR structural element — the iron responsive element (IRE)⁴⁰ — which affects the translation of a subset of mRNAs that are important for iron homeostasis⁴¹. Briefly, a single conserved IRE stem-loop close to the cap of the mRNAs encoding either the iron storage protein ferritin or iron transporter ferroportin is bound by iron-regulatory protein 1 (IRP1) or IRP2 in low-iron conditions. IRP binding represses translation initiation by preventing the 43S pre-initiation complex from associating with the mRNA^{42,43} (FIG. 2b). The IRE-IRP ribonucleoprotein (RNP) complex sterically blocks ribosome access to the cap and 5' UTR. Other stable RNA secondary structures such as cap-proximal hairpins⁴⁴ might block the assembly of the 43S pre-initiation complex onto the 5' UTR^{35,45}. The DEAD-box RNA helicase eukaryotic initiation factor 4A (eIF4A), as part of the eIF4F complex that is assembled at the cap, is thought to be crucial for unwinding such structures and therefore preparing a clear path for ribosome scanning⁴⁶ (BOX 1).

The RNA helicase eIF4A unwinds RNA structures

Recent studies suggest that the function of eIF4A in binding and unwinding RNA can have specific effects on target mRNAs, at least in part through structured RNA elements in 5' UTRs^{47–49}. Although scanning is assisted by the ATPase-dependent duplex-unwinding activity of eIF4A⁵⁰, these studies indicate that eIF4A activity is sensitive to both local RNA structures and sequence motifs. Unwinding of local structures may also explain why certain mRNAs that harbour long 5' UTRs and high GC content are still efficiently translated. For example, the 5' UTR of the *LINE1* mRNA has a high GC content of 60% and is 900 nucleotides in length but is still translated in a cap-dependent manner at a rate similar to that of the very well translated β -actin mRNA⁵¹.

Recent unbiased studies based on ribosome profiling have sought to understand the potential specificity of eIF4A in promoting the translation of certain classes of mRNA^{47–49}. Specific small molecule inhibitors of eIF4A are well characterized. For example, silvestrol^{52,53} and rocaglamide^{52–54} increase the affinity of eIF4A for RNA. These drugs block the dissociation of eIF4A from RNA and thereby reduce eIF4A recycling. Ribosome profiling of silvestrol-treated human KOPT-K1 leukaemia cells revealed a decrease in the translation efficiency of mRNAs with long 5' UTRs⁴⁷, including oncogenes, chromatin modifiers and transcription regulators, which may require tight control of their expression. Interestingly, many of these silvestrol-sensitive transcripts appear to encode a specific structural element in their 5' UTRs: 12-nucleotide long (CGG)₄ motifs that can fold into stable, energetically favourable RNA G-quadruplex (RG4) structures *in vitro*⁴⁷ (FIG. 2b), discussed further below. As eIF4A activity is hyperactivated in cancer and silvestrol has been employed in preclinical cancer

studies⁵⁵, RNA structures and the RNA helicase activity of eIF4A could be relevant for cancer research⁴⁷. Consistent with this possibility, ribosome profiling in silvestrol-treated breast cancer cells revealed that translation initiation was reduced at hundreds of mRNAs, and sensitivity to eIF4A inhibition correlated with the complexity of the inferred structures and the increased length of their 5' UTRs⁴⁸. However, only 25% of 5' UTRs of eIF4A-sensitive mRNAs contain (CGG)₄ motifs⁴⁸, so mRNAs with long 5' UTRs and structural features other than RG4s may be required for eIF4A sensitivity⁵⁶. It remains to be determined how many of these 5' UTR motifs in fact fold into RG4 or other structures. Agreeing in part with the drug-based eIF4A inhibition studies, eIF4A knockdown in MCF7 breast cancer cells followed by polysome profiling and RNA-sequencing analysis revealed eIF4A-sensitive mRNAs that harboured 5' UTRs with a highly negative ΔG and GC-rich motifs with potential to form RG4s, these measurements also uncovered U-rich and GA-rich sequence motifs⁵⁷. Moreover, a more recent study using the eIF4A inhibitor rocaglamide A (RocA) found that 5' UTR structures, including RG4s, contribute little to translation repression⁴⁹. Rather, by use of toeprinting and RNase I footprinting, RocA was suggested to clamp eIF4A onto polypurine motifs in the 5' UTRs of target mRNAs. Further increasing this effect of RocA is the possibility that eIF4A 'trapping' on certain mRNAs may sequester the helicase from being recycled to other mRNAs that require resolution of 5' UTR structures, including those with RG4 structures.

In addition to the classic RNA helicase eIF4A, other helicases and initiation factors are also being revealed as key players in translation control. A helicase that may have overlapping activity with eIF4A is the budding yeast helicase Ded1, which appears to be required to scan through long, structured 5' UTRs⁵⁸. Other helicases with redundant function in translation, such as the DExH-box protein DHX29, can partially rescue the unwinding of structured 5' UTRs in the absence of eIF4A activity⁵⁹. In addition, the eIF4A cofactor eIF4B has been found to stimulate translation of long mRNAs containing structured 5' UTRs in budding yeast independently of eIF4A, as demonstrated by ribosome profiling, which also correlates with 5' UTR structure accessibility assessed *in vitro*⁶⁰. Finally, following the release of eIF4A upon recognition of the start site, the budding yeast DEAD-box helicase Dhh1 specifically promotes translation of mRNAs that have long and highly structured coding regions⁶¹. These examples highlight how diverse RNA helicases and initiation factors can target specific mRNAs with structured regions or motifs, which otherwise may serve as roadblocks to scanning and translation initiation.

RNA G-quadruplex structures

RG4s are stable *in vitro*, with melting temperatures that are higher than physiological temperature, especially in the presence of potassium ions (K⁺), which are specifically chelated inside G-quartets. As the cytoplasm contains high concentrations of K⁺, it has been assumed that RG4s also fold *in vivo*. The formation of RG4 structures — if validated inside cells — would represent the most stable RNA structure that could block ribosome scanning. Beyond the helicase eIF4A and in contrast to the extensively studied DNA G-quadruplexes⁶², other physiological roles of RG4s in mRNAs have only fairly recently been explored (reviewed in REFS 63,64) and include roles in mRNA processing and translation regulation (reviewed in REFS 65,66). Most examples of RG4s in 5' UTRs are linked to

translation repression in *cis*^{65,67,68} presumably by preventing the 43S pre-initiation complex from binding to mRNA or by slowing down scanning^{69,70} (FIG. 2b). A scanning block by RG4 structures was first suggested for the human *NRAS* proto-oncogene mRNA using *in vitro* translation assays^{69,71} and for the *Zic1* zinc finger protein mRNA in eukaryotic cells⁷². The stability and position of RG4 structures close to the 5' end of 5' UTRs contribute to translation repression *in vitro*, as tested for the *NRAS*RG4 (REF. 69), as well as in cells⁷⁰. However, the inhibitory effect of 5' UTR RG4s on scanning is still speculative. This effect and the formation of 5' UTR RG4 structures⁶⁷ need to be carefully confirmed and studied inside cells. Probing RNA structures inside cells and on a genome-wide scale has only recently been adapted to address the physiological relevance of RG4s in mRNAs. The current data suggest that, inside cells, RG4s in most mRNAs appear mainly unfolded (Supplementary information S1 (box)).

Scanning inhibition is thought to be further increased by recruitment of RG4-stabilizing proteins^{63,66} such as fragile X mental retardation protein (FMRP)⁷³, which binds to many RG4-harboring mRNAs^{74,75} (FIG. 2b). At least *in vitro*, FMRP appears to bind an RG4 in the CDS of its own mRNA (*FMR1*)⁷⁶ and can regulate its own alternative splicing⁷⁷. However, a clear role for the FMRP–RG4 interaction in translation repression has only been shown for 5' UTR RG4s in mRNAs other than *FMR1* (REFS 76,78). FMRP might inhibit translation initiation or elongation by binding to and stabilizing RG4 structures and recruiting *trans*-acting factors or by direct binding and stalling of the translating ribosome^{79,80}. Other examples exist of RBPs that promote translation by destabilizing RG4 structures in the CDS⁸¹, further highlighting the diverse roles of stable RNA tertiary structures in translation regulation.

Higher-order mRNA structures

RNA secondary structures can form higher-order interactions to assemble tertiary structures or intermolecular RNA complexes. For example, pseudoknots are complex intramolecular RNA structures consisting of at least two intercalated stem–loop structures that form a knot-like three-dimensional shape. A pseudo-knot structure conserved across mammals, along with contiguous helices, has been proposed to reside in the 5' UTR of human interferon gamma (*IFNG*) mRNA^{82,83}, on the basis of *in vitro* ribonuclease mapping, in-line probing and compensatory mutagenesis coupled with structure–function experiments (FIG. 2b). This pseudo-knot signals to another member of the innate immunity pathway, protein kinase R (RNA-activated) (PKR), which is induced by interferon. PKR is typically activated by double-stranded RNAs of >33 bp in length, which do not appear naturally in the cell but are commonly generated by viruses during infection⁸⁴. Normally, initiating ribosomes unfold the pseudoknot in the *IFNG* 5' UTR. The pseudoknot structure refolds as part of a larger, base-paired RNA structure of sufficient length to attract a PKR dimer. The interaction of PKR with the *IFNG* 5' UTR is thought to locally activate the kinase, which phosphorylates eIF2 α and results in repression of *IFNG* translation (FIG. 2b). Thus, as part of a feedback loop, this RNA structure adjusts translation of its mRNA to PKR activity levels to prevent excess interferon synthesis^{82,83}.

In addition to pseudoknots, RNAs have the capability to form numerous higher-order interactions, including complexes with *trans*-acting long non-coding RNAs (lncRNAs) for post-transcriptional control⁸⁵. In the case of the mouse ubiquitin carboxyl-terminal hydrolase L1 (*Uchl1*) mRNA, the antisense lncRNA *Uchl1AS* produced from the same locus undergoes partial base pairing with the *Uchl1* mRNA 5' UTR, and a repeat region of the lncRNA increases ribosome binding and translation by a so far unexplored mechanism⁸⁶ (FIG. 2b).

To our knowledge, additional examples of 5' UTR *IFNG*-like mRNA pseudoknots or *Uchl1*-like RNA–RNA interactions have not yet been discovered. It is possible, however, that such higher-order interactions, which are difficult to identify, are common in eukaryotic translation initiation. Indeed, pseudoknots constitute a well-known structural motif⁸⁷ in bacterial riboswitches and ribozymes and have roles in eukaryotic pre-mRNA processing such as in splicing^{88,89} and adenosine-to-inosine editing⁹⁰. Furthermore, there are prominent examples of *cis*-regulatory pseudoknots in the CDS that interact directly with translating ribosomes to induce programmed frameshifting (reviewed in REFS 91–93). Frameshifting pseudoknots can either lead to the production of different polypeptides, as first described in retroviruses^{94–96}, or act as mRNA-destabilizing signals^{92,97,98} embedded in coding regions that induce no-go decay or nonsense-mediated decay in eukaryotes. Thus, pseudoknots couple translation to *trans*-acting sensors such as PKR or regulate frameshifting and transcript decay in *cis* to fine-tune mRNA expression. Finally, as discussed next, higher-order structures, including tertiary structures such as pseudoknots, recur in 5' UTRs that control translation in a cap-independent rather than a cap-dependent manner.

IRES structures and function

Perhaps the best understood examples of RNA structure and function in translation control are represented by internal ribosome entry sites (IRESs) in viral genomes (Supplementary information S2 (box)). Viruses evolved IRESs to efficiently hijack the host translation machinery for replication and to overcome the cellular block of cap-dependent translation initiation upon viral infection. Many viral mRNAs rely solely on internal initiation of uncapped viral RNA by specific 5' UTR RNA sequence elements or secondary structures that directly recruit the ribosome for their translation (FIG. 2b). Interestingly, alternative modes of internal but cap-dependent translation initiation are also well established in viruses and are suggested to occur in some eukaryotic mRNAs (Supplementary information S2 (box)).

In contrast to viral RNAs, all cellular mRNAs are capped and thus can undergo cap-dependent translation initiation. However, internal ribosome recruitment by IRES-containing cellular mRNAs is activated or favoured following environmental changes as a means to bypass 5' cap usage to sustain protein expression when cap-dependent translation is diminished⁹⁹ (FIG. 2b). Indeed, most proposed cellular IRESs reside in mRNAs that rely on internal initiation for sustained translation in conditions of stress, mitosis or apoptosis, which reduce global cap-dependent translation (reviewed in REFS 100–105). The first cellular IRES, for example, was discovered in the mRNA encoding immunoglobulin heavy

chain-binding protein (BiP; also known as GRP78), a stress-induced chaperone¹⁰⁶. The IRES sustains BiP translation¹⁰⁷ especially during viral infection¹⁰⁸.

Cellular IRES activity during stress

Ten to fifteen per cent of mammalian mRNAs are predicted to contain IRESs¹⁰⁴, and over 100 proposed IRES-containing mRNAs have been reported¹⁰⁹. These mRNAs mostly encode transcription factors, growth factors and transporters¹⁰⁵. Nevertheless, after decades of work, few examples have been well characterized. Only in the past 10 years or so have new experimental tools and controls been developed that stringently assess the activity of proposed cellular IRESs¹¹⁰. The importance of RNA structure for IRES activity has only been indicated for some cellular IRESs, and RNA structures have been chemically and enzymatically probed for only a handful of examples (Supplementary information S3 (table)). Overall, cellular IRESs appear to be less structured than viral IRESs^{111,112}, with few structural similarities to each other¹¹³, and their mechanisms of action are largely unknown. The activity of cellular IRESs can depend on rather short motifs¹¹⁴ and on the assistance of translation initiation factors or RBPs that serve as IRES *trans*-acting factors (ITAFs; reviewed in REFS 112,115,116) (FIG. 2b). Although several ITAFs are common to viral and cellular IRESs^{112,117}, only a few ITAFs are well characterized. IRES–ITAF interactions may contribute to stabilizing unstable IRES structures or to inducing a conformational change of the IRES RNA that enables the recruitment and correct positioning of the ribosome. We propose that cellular IRESs can largely be categorized into three groups based on which factors or mRNA elements interact with the IRES structure: assisting ITAFs that remodel IRES structures; uORFs that sequester ribosomes and affect IRES structures; or RG4 structures as part of IRES structures (FIG. 3).

ITAFs remodel cellular IRES structures

Several cellular IRESs are thought to be activated through a structural change in their RNA motifs following a change in cellular conditions. This often requires RBPs to act as RNA chaperones. Examples include the IRESs in the *MYC*¹¹⁸, apoptotic peptidase activating factor 1 (*APAF1*)¹¹⁹ and BCL-2 associated athanogene 1 (*BAG1*)^{120,121} mRNAs. The proto-oncogene *MYC* IRES is activated by genotoxic stress, viral infection or apoptosis. A point mutation found in the *MYC* IRES structure¹²² in cell lines derived from individuals with multiple myeloma was predicted to lead to the formation of an additional stem–loop that increased both IRES activity¹²³ and binding of the ITAFs Y-box-binding protein and polypyrimidine tract-binding protein 1 (PTB)¹¹⁸. In the highly structured *APAF1* IRES RNA, binding of NRAS upstream gene protein (UNR) to a purine-rich region in a stem–loop opens two stem–loop structures and allows neural PTB (NPTB) binding¹¹⁹, which generates a single-stranded site for 40S recruitment (FIG. 3a). Similarly, poly(rC)-binding protein 1 (PCBP1) opens the structure of the *BAG1* IRES, but ribosome recruitment requires the subsequent binding of PTB, for example upon heat stress^{120,121}. Thus, different IRESs require sequential or combinatorial binding by ITAFs to induce the structural changes that serve to recruit ribosome subunits in a cap-independent manner. At present, only a handful of ITAFs have been characterized. In the future the identification of additional RBPs that can either stabilize IRES structures or favour the internal recruitment of ribosomes will be an important area of investigation.

Cap-inhibitory elements favour IRES-dependent translation

A different mechanism to favour IRES-mediated translation over cap-dependent translation is stalling of cap-initiated ribosomes in a short uORF²⁴. Examples include the uORFs upstream of amino acid transporter, cationic 1 (*CATI*; also known as *SLC7A1*)^{124,125} and fibro-blast growth factor 9 (*FGF9*)¹²⁶ IRESs and within the vascular endothelial growth factor A (*VEGFA*)¹²⁷ IRES, which are all in mRNAs that encode regulators of differentiation and cell growth. In the *CATI* mRNA, structural remodelling of the 5' UTR *in cis* by uORF translation mediates unfolding of inhibitory 5' UTR structures and a switch to a translationally active state of the IRES structure upon amino acid starvation^{124,125,128} (FIG. 3b). In contrast to IRES activation by uORF translation, uORFs can also repress IRES activity, as seen in the *VEGFA*¹²⁷ and *FGF9* (REF. 126) mRNAs. In a *VEGFA* mRNA isoform, a uORF embedded in the IRES is translated in a cap-independent manner and this uORF suppresses IRES-mediated expression of the main ORF¹²⁷, possibly by unfolding the IRES structure. In the *FGF9* mRNA, translation of a uORF upstream of the IRES suppresses FGF9 protein synthesis in normal conditions, whereas hypoxia-induced inhibition of cap-dependent translation and a switch to IRES-dependent translation increases FGF9 protein levels¹²⁶.

In addition to ITAF-induced structural remodelling and uORF translation, RG4s have also been implicated in the regulation of IRES-mediated translation (FIG. 3c). Sequences in 5' UTRs capable of forming RG4s are a functionally important part of the *VEGFA* IRES^{129,130} and a structural feature of the *FGF2* IRES¹³¹, contributing to IRES-mediated translation of both mRNAs. Although the exact functional role of the *FGF2* RG4 has not yet been studied, the *VEGFA* RG4 within the IRES, which *in vitro* folds independently of the IRES, is required to directly recruit the 40S ribosomal subunit, as shown by *in vitro* footprinting and structure mapping¹³², although this remains to be confirmed *in vivo*.

In summary, structured cellular IRESs in 5' UTRs employ different strategies to overcome translation silencing in response to environmental change (FIG. 3c).

The physiological importance of cellular IRESs

Although several cellular IRESs have been investigated mainly under cellular stress conditions, IRES elements are emerging as important regulators of normal gene expression programmes that underlie embryonic development. A recent study identified and characterized conserved structured IRESs in the 5' UTRs of a subset of genes in the homeobox (HOX) gene family¹³³, which encode key regulators of embryonic development and tissue patterning (FIG. 3d). The HOX 5' UTRs repress their cap-dependent initiation in normal physiological conditions through the cooperation of RNA elements — an IRES and a newly described 5' proximal translation inhibitory element (TIE)¹³³, which acts as a potent inhibitor of cap-dependent translation (FIG. 3c,d). The TIE acts in a highly modular fashion, as its placement upstream of the well-initiated 5' UTR of the mRNA encoding β -globin suppresses cap-dependent translation of a reporter mRNA. Thus, translation initiation of the HOX mRNAs is enabled from IRESs in physiological conditions, as TIE motifs enable these mRNAs to bypass the cap-dependent pathway. This TIE–IRES coupling perhaps enables more intricate control of expression in time and space by ribosomes, as suggested by the

finding that some HOX IRESs selectively require a specific ribosomal protein, RPL38, for their activity¹³³ (FIG. 3d). For example, RPL38 expression is markedly enriched in specific regions of the embryo, such as developing somites (precursors of vertebral elements), and is required at these locations to control HOX gene expression at the post-transcriptional level through IRES elements within HOX 5' UTRs. Thereby, certain ribosomal proteins, such as RPL38 (REF. 133) and RPS25 (REF. 134), as well as the pseudouridylation modification of rRNA¹³⁵, can specifically regulate IRES-dependent translation and highlight the specific contribution of the ribosome itself to gene expression¹³⁶.

HOX IRESs are examples of structured RNA elements that physiologically regulate gene expression during embryonic development. Another example is the IRES in the human *FGF1* mRNA. A domain of the *FGF1* IRES is conserved in sequence and structure among six mammals¹³⁷. The importance of the HOX and *FGF1* IRES structures for their function was confirmed by structural and compensatory mutagenesis. Additionally, the *Hoxa9* IRES structure was characterized *in vitro* by selective 2'-hydroxyl acylation analysed by primer extension (SHAPE), mutate-and-map¹³³ and multiplexed •OH cleavage analysis with paired-end sequencing (MOHCA-seq)¹³⁸ analyses.

The targeted knockout of the *Hoxa9* IRES in mice reveals the crucial role for this element in translation of the mRNA in specific regions of the embryo, such as the developing somites and neural tube¹³³. The *Fgf2* and *Myc* IRESs in reporter constructs are also active in the developing mouse embryo, and their abnormal IRES activation has been linked to cellular transformation^{123,139,140}. *VEGFC* IRES activity is specifically increased in metastatic lymph nodes¹⁴¹, for example, whereas *Fgf2* IRES activity may be important for the spatiotemporal regulation of FGF2 expression in neuronal maturation during brain development in mice¹⁴² and in hypoxia¹⁴³. The finding of cellular IRESs that are active in physiological conditions paves the way for systematic approaches¹⁴⁴ aimed at discovering functional, tissue-specific IRESs genome-wide (Supplementary information S2 (box)). A central question, especially in the case of cellular IRESs, is whether their activity relies on local RNA structures or sequences and/or on assisting proteins. Analysis of IRES structures by structure probing (Supplementary information S3 (table)) should be complemented with compensatory mutagenesis, ideally for every base pair in proposed structural models, to assess their contribution to function. It is also tempting to speculate that structural changes in each IRES RNP are likely to occur in a tissue-specific or stimuli-dependent manner and therefore indicate potentially important 5' UTR elements whose RNA structure should be determined *in vivo*.

Direct eIF3 recruitment by RNA structures

More evidence of translation initiation bypassing the 5' cap of cellular mRNAs comes from two recent studies focusing on the multisubunit initiation factor eIF3. Outside its central role in the 43S pre-initiation complex — physically connecting eIF4G at the 5' cap to the 40S ribosomal subunit — eIF3 appears to repress or activate a specific subset of mRNAs by directly binding to stem-loop structures in their 5' UTRs¹⁴⁵ (FIG. 2b). Transcriptome-wide analysis in human cells by photo-activatable ribonucleoside-enhanced crosslinking and immunoprecipitation (PAR-CLIP) showed that eIF3 directly interacts with 3% of all

mRNAs, particularly in the 5' UTR. These interactions were studied further in two mRNAs encoding the cell proliferation regulators c-Jun (also known as AP1) and B cell translocation gene 1 protein, which are subject to eIF3-mediated translation induction and repression, respectively. Characterization of the *Jun* 5' UTR led to the identification of a conserved hairpin that may serve to directly recruit eIF3 for translation activation. Although disruptive mutations in the hairpin abolish eIF3-dependent translation activation, it remains unknown whether the RNA structure is necessary for function, as compensatory mutations are required to demonstrate the relevance of the structure. Interestingly, a putative IRES in the *Jun* 5' UTR that directs internal translation in glioblastoma cells¹⁴⁶ has previously been mapped to the same region as the suggested eIF3-bound stem-loop. It remains unclear whether the putative stem-loop is part of this IRES or is required for IRES activity. Analysis of additional eIF3 target mRNAs is necessary to determine how widespread this alternative initiation pathway is and to what extent 5' UTR RNA structures are required for it. In this respect, a specific eIF3 subunit, eIF3d, was surprisingly found to directly bind to the 5' cap¹⁴⁷ (FIG. 2b), a role previously thought to be exclusive to eIF4E. This non-canonical cap interaction may serve to regulate transcript-specific translation of certain eIF3-bound mRNAs such as *Jun*¹⁴⁷. Overall, eIF3-specific translation represents an additional role for eIF3 in selective translation.

eIF3 binds RNA modifications to mediate cap-independent initiation

Recent work discovered that RNA modifications also employ the eIF3 complex for cap-independent ribosome recruitment (FIG. 2b). Reversible RNA modifications have long been overlooked with respect to their impact on mRNA structure and translation (reviewed in REFS 148,149). The most prevalent internal chemical modification of RNA, N⁶-methyladenosine (m⁶A)¹⁵⁰ (FIG. 4a), was recently shown to stimulate internal ribosome recruitment¹⁵¹ (FIGS 2b,4b). *In vitro* footprinting and translation assays showed that initiation complexes bind to m⁶A-containing 5' UTRs, which stimulates cap-independent translation through the direct interaction of m⁶A with eIF3 (REF. 151) (FIG. 4b). This type of initiation requires scanning and a free 5' end but is independent of the 5' cap itself or cap-binding proteins. This mechanism may indicate that m⁶A in the 5' UTR can serve as an alternative to the cap and is selectively bound by eIF3 to stimulate initiation at sites termed 'm⁶A-induced ribosome engagement sites' (REF. 151). This effect of 5' UTR m⁶A on translation becomes especially important in stress conditions, such as heat shock¹⁵², which selectively increases and redistributes m⁶A in 5' UTRs, especially in those of newly transcribed mRNAs (FIG. 4b). During heat stress, the m⁶A 'reader' YTH domain-containing family protein 2 (YTHDF2) localizes to the nucleus and promotes the maintenance of 5' UTR m⁶A levels by inhibiting binding of an m⁶A 'eraser', fat mass and obesity-associated protein (FTO)¹⁵². This process mediates selective cap-independent translation of stress response transcripts, such as those of the heat shock-induced heat shock protein 70 (HSP70; also known as HSPA) gene family, which are m⁶A-modified at a single site¹⁵². A single m⁶A in a 5' UTR is also sufficient to recruit eIF3, which by itself can attract the 43S pre-initiation complex¹⁵¹. As m⁶A content increases in stress conditions, which also activate IRES-dependent translation of specific mRNAs, it would be interesting to determine to what extent m⁶A modifications in cellular IRESs are responsible for mediating cap-independent translation.

RNA modifications promote translation

RNA m⁶A modifications may also increase translation efficiency by unfolding RNA structures to recruit proteins^{153,154} or to aid ribosome scanning¹⁵⁴. Using nuclease probing, m⁶A was recently found to disrupt local RNA structures termed m⁶A switches (REF. 153) in mRNAs and lncRNAs^{153–155}, thereby rendering previously paired RNA motifs less structured and thus accessible to ‘indirect readers’ (FIG. 4c). The RNA duplex-destabilizing effect of m⁶A is supported by nuclear magnetic resonance and thermodynamic measurements *in vitro*¹⁵⁶ and by the correlation of *in vivo* click SHAPE (icSHAPE)-determined local RNA structural changes at or near predicted m⁶A sites in the transcriptome¹⁵⁴. It will be crucial to define to what extent the locally unfolded 5′ UTR sequences around m⁶A modifications favour translation initiation by the scanning ribosome¹⁵⁴. Furthermore, m⁶A readers integrate many cues from the 5′ UTR¹⁵² and 3′ UTR^{157–159} that have been associated with increased translation efficiency or mRNA decay (FIG. 4b). In addition to m⁶A, the less abundant N¹-methyladenosine (m¹A), which occurs mainly in structured regions of mRNA 5′ UTRs¹⁶⁰, and hydroxymethylcytosine (hmrC) in coding regions¹⁶¹ have been associated with increased translation initiation and elongation, respectively. However, increased co-transcriptional methylation of A to m⁶A in mRNA coding regions during slow transcription¹⁶² as well as disrupted tRNA selection by m⁶A in the CDS¹⁶³ was recently shown to result in decreased translation efficiency and elongation dynamics, respectively, of m⁶A-modified mRNAs (FIG. 4b).

Recently, circular RNAs (circRNAs) that are formed by the joining of 5′ and 3′ ends through back-splicing, were found to be translated into functional proteins^{164,165} (FIG. 2b). As they do not have a cap, ribosomes need to be internally recruited onto circRNAs, which may be mediated by m⁶A modifications before the start codon^{164,166}. Together, these recent data suggest that reversible RNA methylation, directly or through its impact on local RNA structural topology, is a general feature of mRNA function and acts to increase translation initiation.

5′ UTR RNA structure probing *in vivo*

RNA structures are presumably more dynamic in living cells than in typical *in vitro* experiments, owing to the engagement of *trans*-acting RNAs, small molecules and proteins. However, most RNA structural data of 5′ UTRs have in fact been obtained *in vitro* (Supplementary information S1 (box)). Although the cell membrane is an obvious barrier to getting large RNA-modifying molecules into cells, cell-permeable chemicals such as dimethyl sulfate (DMS) and SHAPE reagents have proved successful for RNA structure probing inside living cells (FIG. 5a,b), and they are now being applied to map the folding of the transcriptome *in vivo* (reviewed in REFS 167–171) (FIG. 5c). Transcriptome-wide comparison of RNA structures *in vivo* and *in vitro* by DMS treatment followed by deep sequencing (DMS-seq) has revealed that in cells, mRNAs are generally less structured than they are *in vitro*¹⁷² (Supplementary information S1 (box)). Analysis of living cells by icSHAPE has also shown that RNAs are less folded *in vivo*, although the extent of unfolding varies between RNA classes¹⁵⁴, and RNA structural features *in vivo* may be used to distinguish between coding and regulatory RNAs. For example, the region just upstream of

the start codon appears particularly less structured in cells¹⁷³. Nevertheless, the structural interpretation of these data remains uncertain, as cellular mRNAs are covered with RBPs, RNA helicases and translating ribosomes that constantly remodel the mRNP and influence the RNA structure^{172–174}. Indeed, the extent of unfolding of the transcriptome is thought to be ATP dependent¹⁷², which indicates a major role for RNA helicases in shaping RNA structures.

New psoralen-based technologies that map sites of *cis* and *trans* RNA–RNA interactions (Supplementary information S1 (box)) may also provide new insights into the folding status of the cellular transcriptome and of specific RNA duplexes. As these methods have only recently been established, data remain sparse for RNAs of relatively low abundance, including most mRNAs. As biases in protocols are reduced, sequencing depths increase, data normalization is standardized, and specific mRNAs are targeted for in-depth analysis^{175–177}, it may become possible to investigate the roles of RNA–RNA interactions and RNA structures in translation initiation.

A substantial improvement of SHAPE¹⁷⁵ and DMS-based¹⁷⁸ RNA structure probing has been achieved by reading out nucleotides affected by chemical probes as mismatches introduced at the adduct site during reverse transcription (FIG. 5b) and is being adapted for application in cells. For example, DMS mutational profiling with sequencing (DMS–MaPseq), an *in vivo* adaptation of DMS–seq, was recently used for global or targeted RNA structure probing of low-abundance RNAs in *Drosophila melanogaster* ovaries, as well as in yeast and mammalian cells¹⁷⁷. This approach includes compensatory mutations of stem structures modelled in the 5′ UTR of FMR1 autosomal homologue 2 (*FXR2*) mRNA, which showed rescue of activity, although the effect was weak¹⁷⁷.

What are the next steps in RNA structure probing?

The current generation of high-throughput methods for probing RNA accessibility and base pairing inside cells represent substantial advances over prior techniques. However, these methods still appear to be too limited to test the repertoire of proposed mRNA regulatory structures or to establish new ones (Supplementary information S1 (box)). There are two major limitations to these *in vivo* methods, both inherited from their ‘parent’ *in vitro* methodologies. First, structure modelling from these data remains mostly tied to chemistries and modelling methods whose accuracies are poor even *in vitro*^{179,180}. Any such inaccuracies are further exacerbated by the numerous ‘unknowns’ of RNA interactions in cells, from RBP interactions to mRNA domains potentially forming a heterogeneous ensemble of structures. Second, although chemical probing methods coupled with computation can provide base-pair-level structural models, they have generally not been tested in prospective experiments — not even *in vitro*. This absence of standard validation methods is a major obstacle for our understanding of how mRNA structures impact mRNA translation.

Fortunately, solutions for both problems have recently been found for *in vitro* RNA structure modelling and could be brought to bear in the next generation of *in vivo* chemical probing methods. New *in vitro* methods infer effects of mutations at each RNA nucleotide from changes in chemical accessibility elsewhere in the RNA (mutate-and-map)^{181,182} (FIG. 5d).

These methods have enabled highly accurate structure determination in structure prediction competitions^{180,183}. Numerous strategies are now available to generate RNA libraries with mutations inside cells^{176,184,185}, which raises the prospect of carrying out the same mutate-and-map analysis *in vivo*, high-throughput versions of methods to correlate hydroxyl radical damage at nearby nucleotides¹³⁸ to study tertiary structure are now available and were recently applied to study chromatin structure *in vivo*¹⁸⁶. These methods promise a considerable boost in the accuracy of RNA structure modelling from *in vivo* data, even of molecules with complex secondary and tertiary structures¹⁸³.

The gold standard for validating RNA structures remains rescue by compensatory mutagenesis, although this approach has typically required quantitative functional readouts that need to be individually developed for each RNA molecule (see, for example, REF. 177). Recently, compensatory mutagenesis has been analysed by using quite general chemical probing readouts (FIG. 5d), even allowing correction of erroneous structures from conventional SHAPE or DMS-based structure mapping¹⁸². Correlation of *in vivo* RNA structures of operon units in bacterial polycistronic mRNAs with their translation efficiency has been supported by *in vivo* DMS-seq of compensatory mutants in one mRNA¹⁸⁷. These early results foreshadow that the detection of secondary and tertiary RNA structures as well as high-throughput compensatory mutagenesis, if adapted *in vivo* and to eukaryotes, may soon enrich our understanding of structured RNA elements that control eukaryotic translation.

Conclusions

An accumulating body of knowledge highlights the diverse repertoire of mechanisms through which translation initiation can be controlled by 5' UTR structures in eukaryotic mRNAs. Nevertheless, for decades, research has focused on linear RNA sequence motifs bound by RBPs that control post-transcriptional processes^{12,188,189}. For many of these motifs, it is not clear whether RNA structures improve or hinder access of proteins by exposing or burying them, respectively, in local structure³⁷. Thus, an unstructured element embedded in an RNA structure may actually be the driving force of a structured motif. In the case of IRESs, this notion could provide an explanation for the discovery of short linear poly(U) motifs that harbour IRES activity¹⁴⁴, for the finding that the ITAF PTB recognizes and binds to a polypyrimidine-rich motif, (CCU)_n, in structured cellular IRESs¹¹⁴, or for the detection of an unstructured region that is important for the activity of the turnip crinkle virus IRES¹⁹⁰, despite the fact that viral IRESs are thought to largely be structured. Thus, local mRNA structures as well as unstructured motifs therein can regulate translation.

How can we better find and confirm functional mRNA structures, both *in vitro* and *in vivo*? Many compact and rapidly evolving prokaryotic genomes have allowed the discovery of novel RNA motifs by means of genome alignments, conservation and sequence covariation analyses. In eukaryotes, evolutionary covariation analysis has been difficult and may be fundamentally impossible if 5' UTR sequences are highly divergent¹⁹¹, even as some mRNA structures are suggested to be abundant and conserved enough¹⁹² to define distinct families^{193,194}. As new global probing tools can now assess nearly all RNA structures in cells, they may enable the discovery of novel RNA structures in complex genomes and

analysis of their contribution to gene regulation¹⁶⁷. These tools may also permit revisiting the analysis of decades-old examples of highly translation-ally regulated mRNAs to assess potential underlying 5' UTR structures such as the long, GC-rich 5' UTR of cyclin D1 mRNA¹⁹⁵. Its translation is thought to depend on eIF4E-assisted structure unwinding by the helicase eIF4A, but the responsible RNA structures are not known.

Just as important as finding regulatory RNA structures based on structure probing or prediction is their experimental validation. We encourage the field to more routinely apply compensatory mutagenesis to evaluate the physiological relevance and functional contribution of candidate mRNA structures. As experimental methods continue to improve, a major goal for global structure probing is their application across diverse cell types and tissues in the context of development, cell differentiation, stress or disease. This approach is especially important as intricate modes of gene regulation such as translation may be highly cell-type specific. Emerging technologies that are highly complementary to *in vivo* structure probing include advances in cryo-electron microscopy¹⁹⁶, which may soon reveal the intricate interactions of the ribosome and structured mRNA elements¹⁹⁷, including cellular IRES elements. Moreover, cryo-electron tomography *in situ*¹⁹⁸ may soon provide three-dimensional snapshots of translating ribosomes and their template mRNAs in native cells. These methods may also profit from machine learning algorithms and conventional biophysical modelling to more accurately predict and model structures in mRNA 5' UTRs¹⁹⁹. We speculate that such advances will enable an mRNA-centric view of translation, in which one can visualize how the ribosome and RBPs change the accessibility of RNA structured domains to control translation in time and space. It has long been appreciated that the ribosome is a dynamic, multi-state machine that has originated from the ancient RNA world. Upcoming advances raise the promise of understanding whether mRNAs and their structures provide, on top of the ribosome, a new layer of gene regulation that allows the control of translation to meet the needs of modern eukaryotic biology.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Glossary

Secondary structures

Patterns of Watson–Crick base pairs (G–C, A–U and G–U) that define the double helices of an RNA.

Tertiary structures

Interactions that orient RNA double helices into specific three-dimensional arrangements, often involving non-Watson–Crick base pairs.

Ribozymes

RNA molecules that catalyse specific biochemical reactions.

Riboswitches

Non-coding mRNA structures that sense the environmental status of a cell by directly binding to small molecule ligands, such as a metabolite or an ion. This interaction changes the RNA conformation and controls gene expression through alternative splicing, transcription or translation.

Peptidyl transferase centre

The site in the large ribosomal subunit that catalyses peptide bond formation and peptide release; it is composed entirely of RNA.

Pseudoknots

RNA tertiary structures formed by base pairing of a single-stranded loop of a hairpin with a complementary sequence outside that hairpin.

Upstream open reading frames (uORFs)

Small ORFs located in the 5' UTR of some mRNAs. Translation of uORFs can regulate the translation of the downstream ORF.

Kozak sequence

A favourable sequence (GCCRCCAUGG in mammals, where R is a purine) surrounding the translation start codon (AUG); also called the Kozak consensus or Kozak context.

GC content

Percentage of guanine and cytosine nucleotides in an RNA molecule. G–C base pairs are more stable than A–U base pairs and can form more stable RNA structures.

Free energy (ΔG)

The stability of an RNA secondary structure, estimated as the sum of the free energies assigned to all loops and base pair stacks of a folded RNA, based on a computational folding algorithm.

5' cap

A modified guanine nucleotide, m⁷GpppN (where m⁷G is 7-methylguanosine, p is a phosphate group and N is any base), located at the 5' end of eukaryotic mRNAs.

Ribonucleoprotein (RNP)

A complex of proteins in association with an RNA (RNP) or mRNA (mRNP).

Ribosome profiling

Sequencing of the RNA fragments protected by ribosomes, providing a quantitative signature of the translated mRNAs at a given time.

RNA G-quadruplex (RG4)

Extremely stable RNA structure formed in G-rich regions by the stacking of at least two G-tetrads, each of them forming a square-shaped structure by non-Watson-Crick interactions between two or more layers of paired G-quartets.

Toeprinting

Nucleotide-resolution assay that uses primer extension inhibition induced by the complex of a protein, or the ribosome, bound to an mRNA, to report its position on the mRNA.

In-line probing

Nucleotide-resolution structure probing method that uses the natural instability of RNA: the 2' hydroxyl of each nucleotide can attack its phosphodiester backbone at a rate dependent on local structure.

No-go decay

An mRNA and translation quality-control mechanism that recognizes and degrades mRNAs following prolonged ribosome stalling during translation.

Nonsense-mediated decay

An RNA surveillance mechanism that recognizes and degrades mRNAs containing premature termination codons to prevent their translation.

Selective 2'-hydroxyl acylation analysed by primer extension (SHAPE)

SHAPE and *in vivo* click SHAPE (icSHAPE) use cell-permeable reagents to acylate the 2'-OH of accessible, single-stranded RNA at all four nucleotide bases.

Mutate-and-map

Two-dimensional expansion of chemical probing that infers RNA base pairing by mutating each nucleotide and detecting increased chemical accessibility at other nucleotides.

m⁶A switches

mRNA sequences that adopt a different secondary structure depending on N⁶-adenosine methylation.

Dimethyl sulfate (DMS)

A cell-permeable reagent that methylates adenine and cytosine nucleotides that are not protected by base pairing. Modification sites stall or induce a mutation during primer extension by reverse transcriptase, and sequencing the resulting complementary DNA indicates the folding state of a nucleotide in an RNA.

Psoralen

A natural plant product that intercalates into DNA and RNA and reversibly crosslinks interacting RNA duplex molecules at UpA motifs upon irradiation with ultraviolet light.

Sequence covariation

Nucleotide substitutions that differ between two or more homologous genes but retain the potential for RNA base pairing in each sequence.

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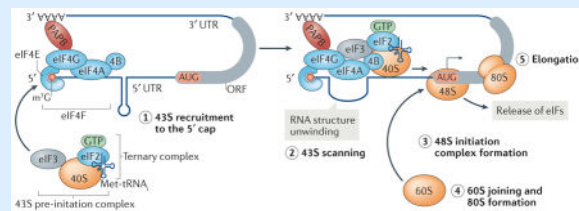
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Box 1**Canonical cap-dependent translation initiation**

The translation of most eukaryotic mRNAs is initiated by ribosome recruitment to the 5' cap, followed by ribosome scanning towards a start codon (see the figure). Canonical cap-mediated initiation mainly occurs by recruitment of the 40S small ribosomal subunit and its associated eukaryotic initiation factors (eIFs) to the 5' end 7-methylguanosine (m⁷G) cap structure (reviewed in REFS 7,8,20). The first step of initiation is the assembly of the trimeric eIF4F cap-binding complex at the 5' cap, wherein cap-binding protein eIF4E interacts with the scaffolding initiation factor eIF4G and the RNA helicase eIF4A. Poly(A) binding protein (PABP) on the 3' poly(A) tail interacts with eIF4G at the cap, thereby circularizing the mRNA. Through its interaction with eIF3, eIF4G recruits the 43S pre-initiation complex (step 1), which consists of eIF3, the 40S ribosomal subunit, the ternary complex of GTP-bound eIF2 and the initiator tRNA (eIF2-GTP-Met-tRNA_i), and eIF1, eIF1A and eIF5 (not shown). Once the 43S complex binds the mRNA near the cap, it travels ('scans') along the 5' untranslated region (UTR) in the 5' to 3' direction (step 2) in an ATP-dependent reaction, with partial hydrolysis of the eIF2-bound GTP to GDP in the ternary complex, until it encounters a start codon (AUG). The RNA helicase eIF4A migrates with the 43S complex and unwinds inhibitory RNA secondary structures in the 5' UTR (a stem-loop structure is shown). Stable binding of the 43S complex at the start codon yields the formation of the 48S initiation complex (step 3) and triggers GTP hydrolysis and release of eIFs. Subsequently, the 60S large ribosomal subunit joins the 40S ribosomal subunit to form the elongation-competent 80S ribosome (step 4), which then proceeds to translation elongation (step 5). Starvation and other stress conditions inhibit the formation of the ternary complex through phosphorylation of eIF2 α and block eIF4F assembly by sequestering eIF4E, which is bound by E-binding protein (not shown), thereby suppressing cap-dependent translation initiation. 4B, eIF4B; ORF, open reading frame.



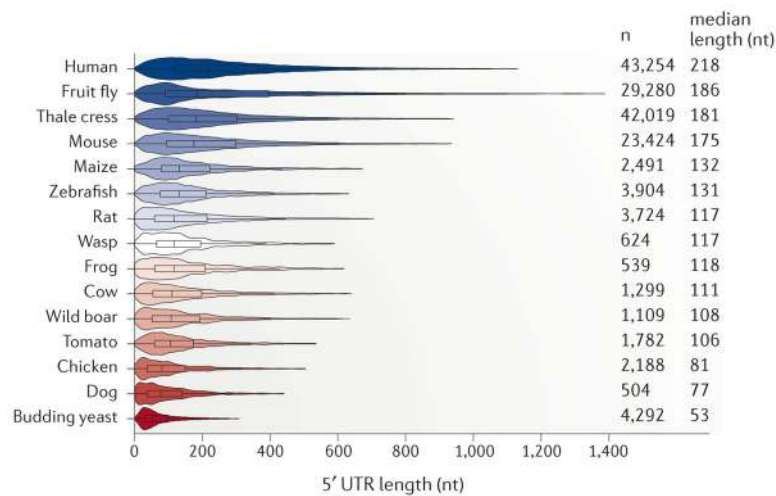


Figure 1. Evolutionary expansion of eukaryotic 5' UTR lengths

The length of 5' untranslated regions (UTRs) has increased in eukaryotes during evolution, with median lengths ranging between 53–218 nucleotides (nt). We compared RefSeq-annotated 5' UTR lengths of reviewed and validated transcripts (n) between species for which at least 100 5' UTRs are annotated. For yeast, we used the 5' UTR lengths as annotated in REFS 34,35. The violin plots depict the distribution of 5' UTR lengths for 15 species sorted according to decreasing median 5' UTR length, including human (*Homo sapiens*), fruit fly (*Drosophila melanogaster*), thale cress (*Arabidopsis thaliana*), mouse (*Mus musculus*), maize (*Zea mays*), zebrafish (*Danio rerio*), rat (*Rattus norvegicus*), wasp (*Nasonia vitripennis*), western clawed frog (*Xenopus tropicalis*), cow (*Bos taurus*), wild boar (*Sus scrofa*), tomato (*Solanum lycopersicum*), chicken (*Gallus gallus*), dog (*Canis lupus familiaris*) and the budding yeast *Saccharomyces cerevisiae*. The data range for each species was trimmed to a maximum of the third quartile plus three times the interquartile range ($Q3 + 3 \times IQR$).

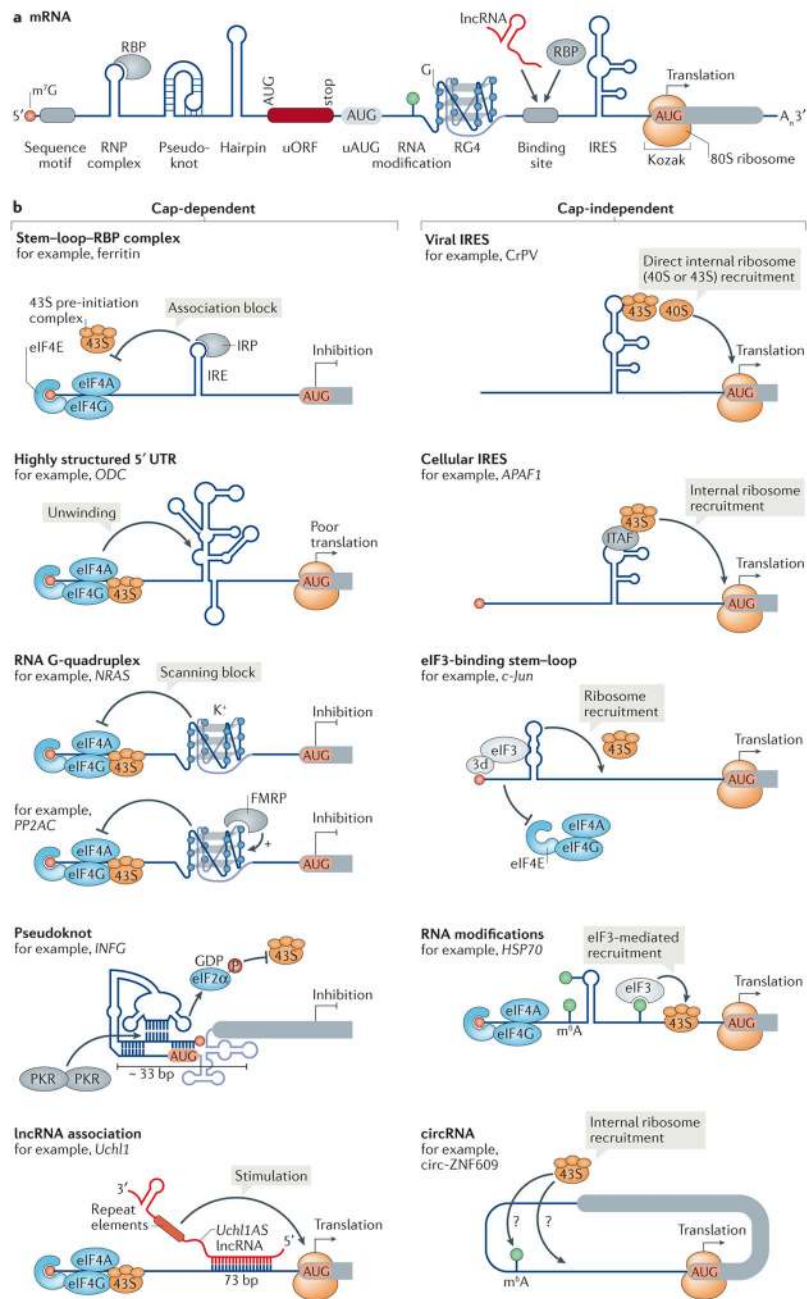


Figure 2. Cis-acting regulatory RNA elements and structures in eukaryotic 5' UTRs influence mRNA translation

a | The 7 methylguanosine (m⁷G) 5' cap structure (circle) at the 5' end of the mRNA and the poly(A) tail (A_n) at the 3' end stabilize the mRNA and stimulate translation. The 5' untranslated region (UTR) contains secondary and tertiary structures and other sequence elements. RNA structures such as pseudoknots, hairpins and RNA G-quadruplexes (RG4s), as well as upstream open reading frames (uORFs) and upstream start codons (uAUGs), mainly inhibit translation. Internal ribosomal entry sites (IRESs) mediate translation initiation independently of the cap. RNA modifications, or RNA binding proteins (RBPs)

and long non coding RNAs (lncRNAs) that interact with RNA binding sites or form ribonucleoprotein (RNP) complexes, as well as the Kozak sequence around the start codon, can additionally regulate translation initiation. **b** | Regulatory 5' UTR RNA structures can influence protein synthesis by promoting or inhibiting either cap-dependent (left) or cap-independent (right) translation. Whereas the structures that regulate cap-dependent initiation — RG4s, stem-loop structures and pseudoknots (but not lncRNAs) — tend to repress initiation, cap-independent regulatory RNA structures, including IRESs, eukaryotic initiation factor 3 (eIF3)-binding stem-loop structures, RNA modifications and circular RNAs (circRNAs), can internally assemble the translation machinery onto the mRNA and generally stimulate translation. 3d, eIF3d; *APAF1*, apoptotic peptidase activating factor 1; circ-ZNF609, circular-zinc-finger protein 609; CrPV, cricket paralysis virus; FMRP, fragile X mental retardation protein 1; HSP70, heat shock protein 70; *IFNG*, interferon gamma; IRE, iron responsive element; IRP, iron regulatory protein; ITAF, IRES *trans*-acting factor; K⁺, potassium; m⁶A, N⁶-methyladenosine; *NRAS*, NRAS proto oncogene, GTPase; *ODC*, ornithine decarboxylase; P, phosphorylation; PKR, protein kinase RNA activated; *PP2AC*, protein phosphatase 2 catalytic subunit alpha (also known as *PPP2CA*); *Uchl1*, ubiquitin carboxyl terminal hydrolase L1.

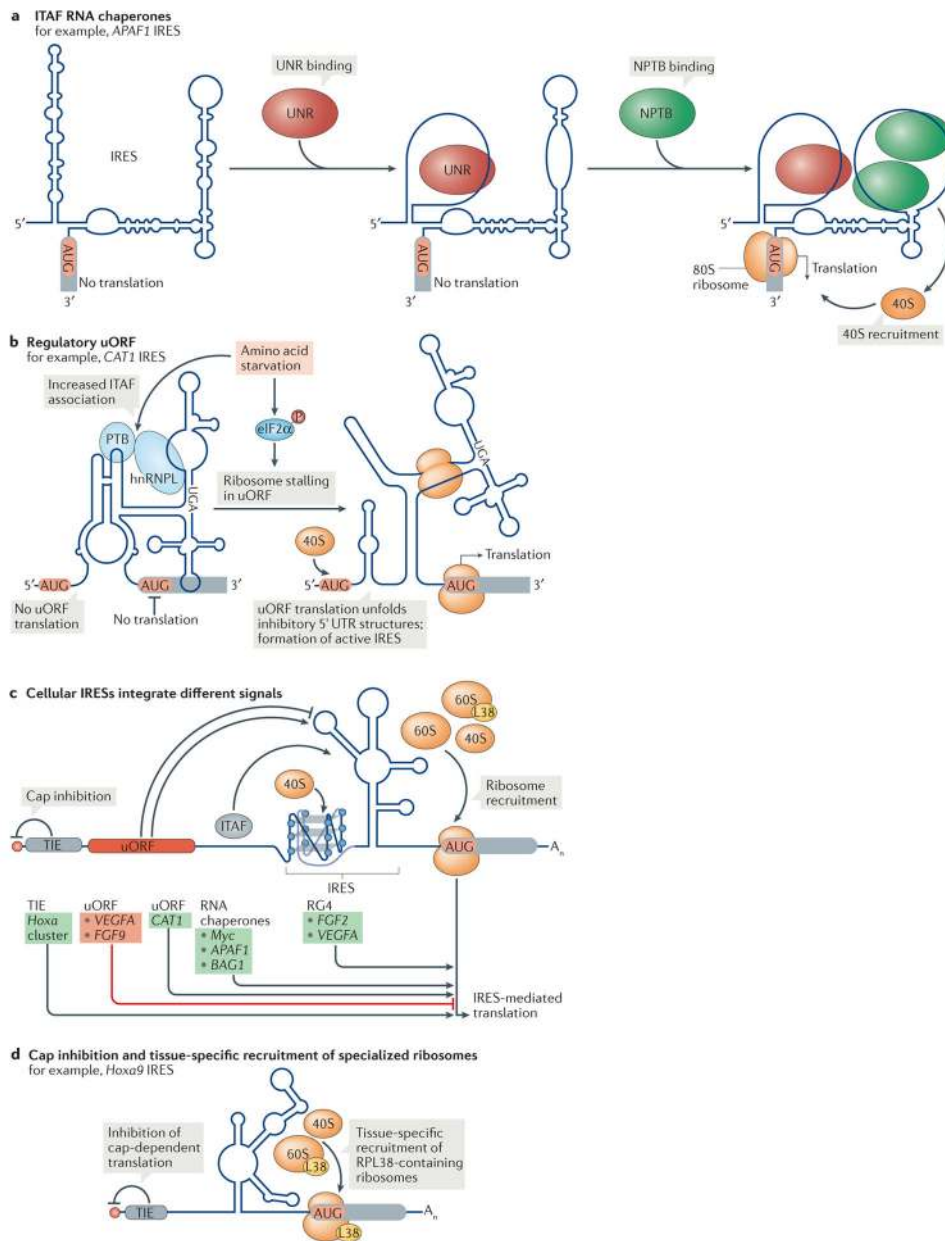


Figure 3. Cellular IRES structures employ different mechanisms for ribosome recruitment
Cellular structured internal ribosome entry sites (IRESs) use diverse modes to recruit ribosomes, and their activity is often induced following a change in cellular condition. **a** | Many IRESs rely on binding by RNA-binding proteins known as IRES *trans*-acting factors (ITAFs) for ribosome recruitment. Such RNA chaperones remodel the IRES structure and thereby prepare a landing platform for the ribosome. In the IRES of apoptotic peptidase activating factor 1 (*APAF1*) for example, binding of NRAS upstream gene protein (UNR) to a purine-rich region in a stem–loop opens two stem–loop structures and allows the binding of neural polypyrimidine tract binding protein (NPTB)¹¹⁹, which creates a ribosome-accessible site for translation. **b** | Cellular IRESs can also use upstream open reading frames (uORFs) to regulate IRES activity. In the 5′ untranslated region (UTR) of the arginine–

lysine transporter amino acid transporter, cationic 1 (*CAT1*) mRNA, translation of a uORF within the IRES is induced upon amino acid stress. This stalls ribosomes in the uORF and causes a structural switch in the IRES to an active conformation, which enables the translation of the main ORF^{124,125}. In addition, the association of the ITAFs PTB and heterogeneous nuclear ribonucleoprotein L (hnRNPL) with the IRES increases and is required for translation during starvation²⁰⁰. **c** | Cellular IRESs can integrate signals in *cis* and *trans* to modulate internal ribosome recruitment. Whereas a translation inhibitory element (TIE) blocks cap-dependent initiation, uORF translation, ITAFs or RNA G-quadruplexes (RG4s) can all increase (green) IRES-mediated translation in a transcript-specific manner; uORF translation can also inhibit (red) IRES activity. **d** | In a subset of homeobox a (*Hoxa*) mRNAs in the mouse embryo, an IRES recruits ribosomes in a tissue-specific manner¹³³. Several of these *Hoxa* IRESs additionally depend on the ribosomal protein RPL38 (L38) for their activity and a TIE at the cap that blocks cap-dependent initiation. *BAG1*, BCL2 associated athanogene 1; eIF2 α , eukaryotic initiation factor 2 α ; *FGF*, fibroblast growth factor; P, phosphorylation; *VEGFA*, vascular endothelial growth factor A.

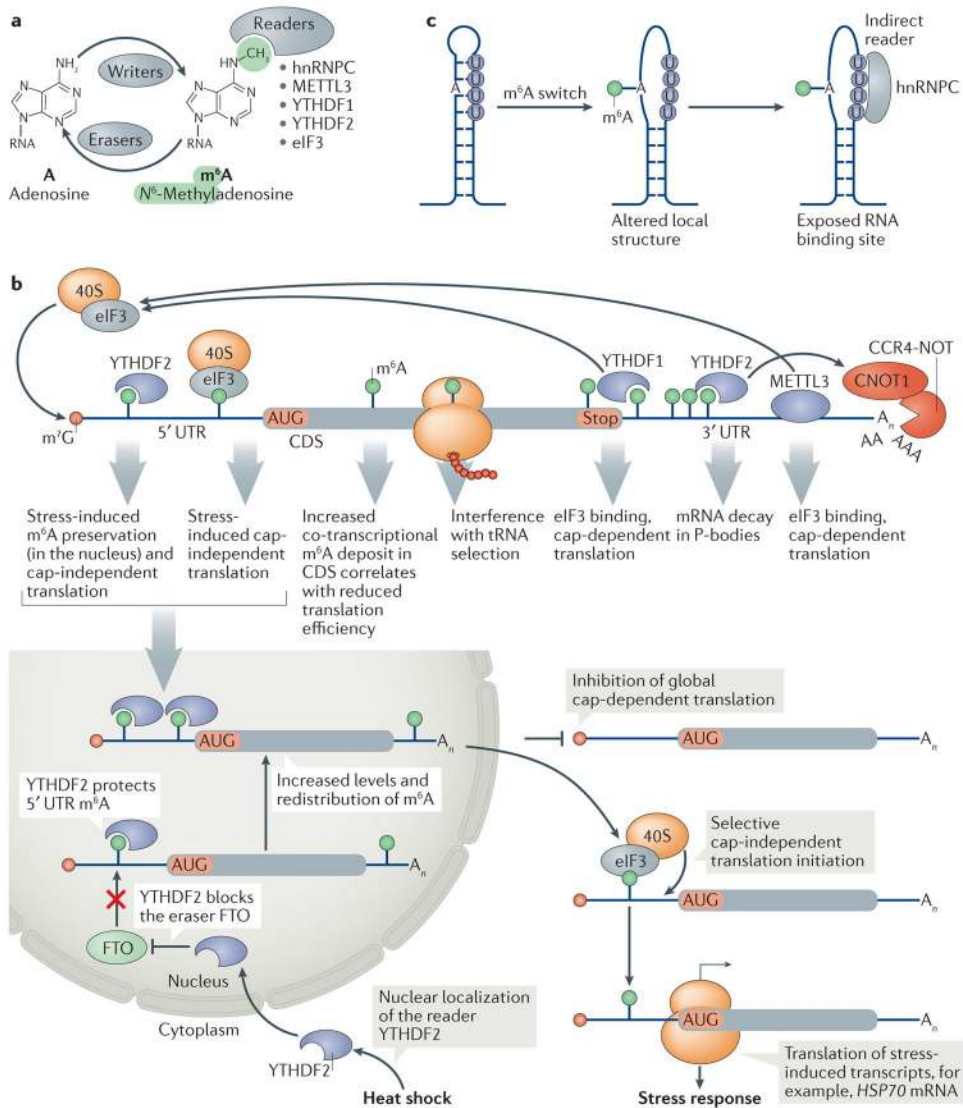


Figure 4. The effects of N^6 -methyladenosine on mRNA translation and decay
a | 'Writer' proteins establish N^6 -methyladenosine (m^6A) at internal RNA sites, 'eraser' proteins remove them, and 'reader' proteins directly bind the N^6 -methyl group of m^6A (reviewed in REF. 148). The listed readers affect the translation and stability of m^6A -modified mRNAs. **b** | m^6A modifications in mRNAs occur with a preference for the last exon and 3' untranslated region (UTR)²⁰¹ and are increased during stress¹⁵². According to its position in the mRNA, m^6A is bound by readers that can induce cap-dependent translation or internal ribosome recruitment. In the 5' UTR, eukaryotic initiation factor 3 ($eIF3$) can directly bind m^6A and facilitate internal translation initiation¹⁵¹. Stress-responsive 5' UTR N^6 -adenosine methylation, for example during heat shock, is preserved by YTH domain-containing family protein 2 (YTHDF2), which blocks binding of the eraser fat mass and obesity-associated protein (FTO), thereby promoting cap-independent translation initiation of stress response mRNAs¹⁵². m^6A in the coding sequence (CDS) is linked to tRNA selection¹⁶³, and at the 3' UTR it is linked to increased translation owing to

YTHDF1 binding to m⁶A and eIF3 recruitment for cap-dependent translation¹⁵⁸. The writer methyltransferase like 3 protein (METTL3) can also directly bind to eIF3 to increase translation of m⁶A-containing mRNAs independently of its m⁶A writer activity¹⁵⁹. By contrast, YTHDF2 promotes degradation of m⁶A-modified mRNAs by recruiting the deadenylase complex CCR4–NOT¹⁵⁷. Together, increased translation efficiency and activated decay of m⁶A-modified mRNAs allow dynamic regulation of protein synthesis. **c** | m⁶A mRNA modifications are associated with unfolded RNA structures. N⁶adenosine methylation in a stem disrupts based paired regions ('m⁶A switch'), which allows binding of the 'indirect reader' heterogeneous nuclear ribonucleoprotein C (hnRNPC) to exposed U rich motifs in the nucleus¹⁵³. CNOT1, CCR4–NOT transcription complex subunit 1; HSP70, heat shock protein 70; P body, processing body. Part **c** is adapted from REF. 153, Macmillan Publishers Limited.

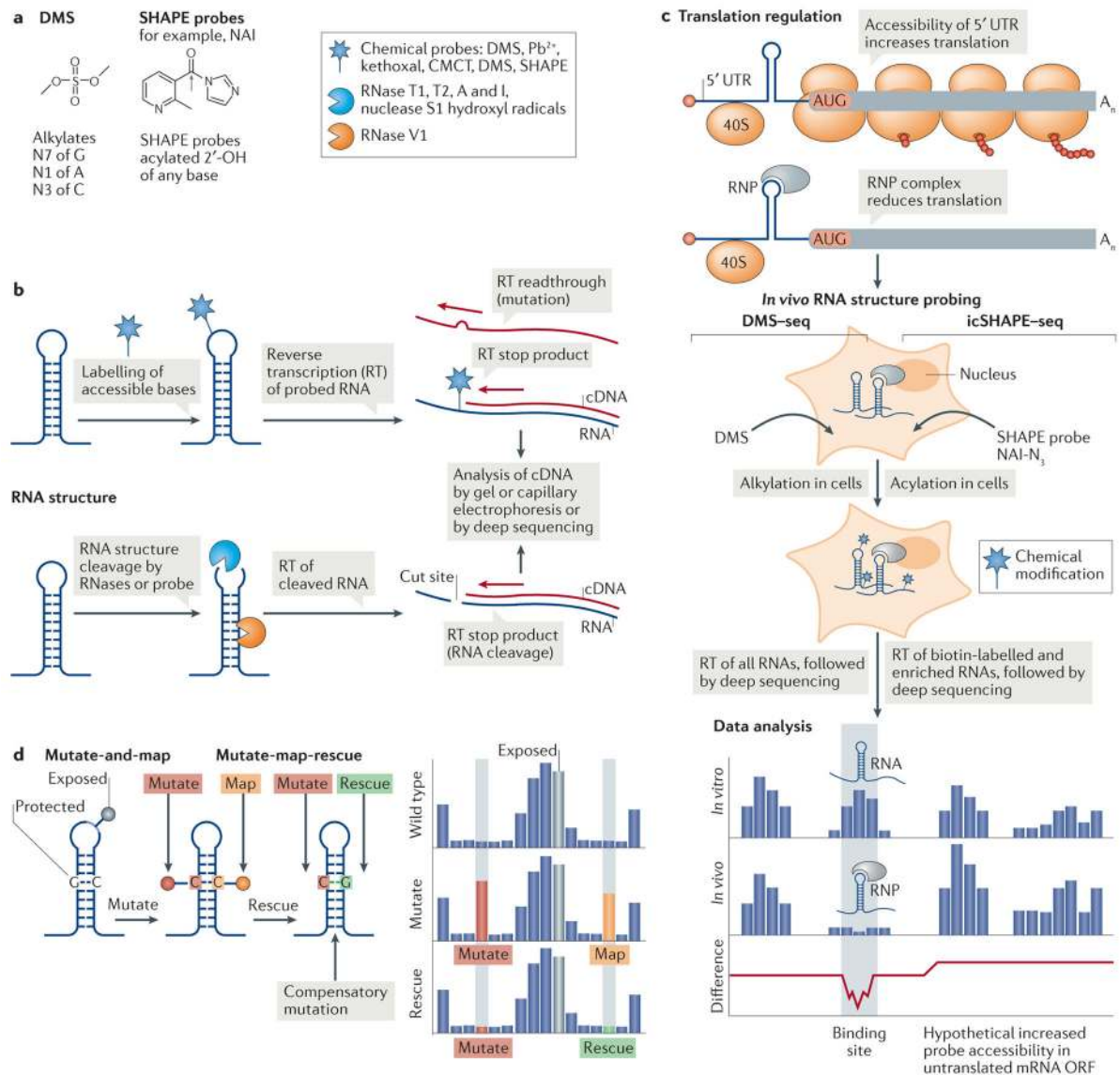


Figure 5. Global RNA structure probing to assess translation regulation

Global RNA structure probing inside cells can assess the transcriptome structure in the presence of proteins. **a** | Chemical schematics of the RNA structure probes dimethyl sulfate (DMS) and the selective 2'-hydroxyl acylation analysed by primer extension (SHAPE) reagent 2 methylnicotinic acid imidazolidine (NAI) and their reactivity. The grey arrow indicates the site of 2'-OH attack of the RNA by the probe. Different probes induce single-strand-specific chemical labelling (star) or cleavage by enzymes or probes (blue Pac Man shape, single strand specific; orange Pac Man, double strand specific). **b** | RNA probing either labels RNA through a covalent reaction of a chemical probe with accessible nucleotides (top) or cleaves the RNA backbone with RNases (bottom). A pool of modified or cleaved RNAs is transcribed into cDNA by reverse transcription (RT), and modified or cleaved sites are identified by their effect on RT. **c** | A 5' untranslated region (UTR) ribonucleoprotein (RNP) complex that inhibits ribosome scanning reduces the accessibility

of the mRNA for the probe in cells. In *in vivo* RNA structure probing, shown here for DMS treatment followed by deep sequencing (DMS-seq) and *in vivo* click SHAPE followed by deep sequencing (icSHAPE-seq), RNA structures are probed in cells by chemical modification. Data analysis of *in vitro*-probed and *in vivo*-probed RNA can indicate the presence of RNA structures as protein binding sites, owing to masked probe accessibility or to remodelling of the structure by protein interaction. Where a 5' UTR RNP complex inhibits the translation of an open reading frame (ORF), the accessibility of the probe to poorly translated coding sequences might increase in the absence of ribosomes. **d** | In multidimensional mutate and map chemical probing, a mutation (red) that eliminates base pairing exposes the mutated nucleotide and its partner nucleotide (orange) to chemical modification (pins). In mutate-map-rescue probing, a mutation is rescued from modification by a compensatory mutation of the partner nucleotide (green). The reactivity profile reflects changes in probe accessibility upon mutation (red), which allows mapping (orange) of the base-paired nucleotide, while rescue (green) confirms base pairing. Nucleotides in loops are exposed and accessible to the probe. Pb²⁺, lead; CMCT, 1 cyclohexyl (2-morpholinoethyl)carbodiimide metho-*p*-toluene sulfonate.