mTOR-sensitive translation: Cleared fog reveals more trees

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ABSTRACT

Translation is fundamental for many biologic processes as it enables cells to rapidly respond to stimuli without requiring de novo mRNA synthesis. The mammalian/mechanistic target of rapamycin (mTOR) is a key regulator of translation. Although mTOR affects global protein synthesis, translation of a subset of mRNAs appears to be exceptionally sensitive to changes in mTOR activity. Recent efforts to catalog these mTOR-sensitive mRNAs resulted in conflicting results. Whereas ribosome-profilng almost exclusively identified 5′-terminal oligopyrimidine (TOP) mRNAs as mTOR-sensitive, polysome-profiling suggested that mTOR also regulates translation of non-TOP mRNAs. This inconsistency was explained by analytical and technical biases limiting the efficiency of ribosome-profiling in detecting mRNAs showing differential translation. Moreover, genome-wide characterization of 5′ UTRs of non-TOP mTOR-sensitive mRNAs revealed 2 subsets of transcripts which differ in their requirement for translation initiation factors and biologic functions. We summarize these recent advances and their impact on the understanding of mTOR-sensitive translation.

Selective regulation of mRNA translation via the mTOR pathway

Gene expression is modulated at multiple levels including transcription, mRNA-splicing, -export, -stability, -translation and protein-stability.1 Each regulatory layer contributes to the repertoire and levels of expressed proteins. Modulation of mRNA-translation and/or protein-stability allow cells to rapidly adjust their proteomes in response to external and internal cues without altering mRNA levels.2,3 As a result, protein levels do not always reflect steady-state mRNA abundance.4-7 Moreover, it is thought that only a fraction of all cellular mRNA is translated at a given moment.8-10 Indeed, although still highly debated,11 mRNA translation has been suggested to modulate protein levels to a similar extent as transcription and has therefore emerged as a principal post-transcriptional mechanism affecting the proteome.5,12 Consistently, translational control plays central roles in pivotal biologic processes including control of the immune system, cell proliferation and development; and diseases including cancer.13,14 Common to these contexts is that mRNA translation is selectively modulated to alter synthesis of specific subsets of proteins which are required to mount an optimal response to a variety of stimuli; and when dysregulated can lead to a wide array of pathologies.15 Thus, deciphering mechanisms by which translation efficiencies of individual mRNAs are reprogrammed in response to stimuli and/or in normal vs. dysfunctional cells is crucial for a more complete understanding of many biologic phenomena.

mRNA translation can be divided into 4 phases – initiation, elongation, termination and ribosome recycling.15 To date, the best described examples of modulation of translational efficiencies occur at the rate-limiting initiation step, i.e. the efficiency of ribosome recruitment to mRNA.16 In mammals initiation is facilitated by multiple eukaryotic translation initiation factors (eIFs) including the eIF4F complex. eIF4F recruits mRNA to the ribosome and consists of the mRNA cap binding subunit eIF4E, the scaffolding protein eIF4G and the DEAD box RNA helicase eIF4A.16 The mechanistic/mammalian target of rapamycin (mTOR) complex 1 (mTORC1) stimulates assembly of the eIF4F complex by phosphorylating and inactivating the 4E-binding proteins (4E-BP1, 2 and 3) which otherwise prevent eIF4E:eIF4G interaction and thereby eIF4F complex assembly.17-21 Although eIF4E is required for cap-dependent translation of all nuclear-encoded mRNAs, some transcripts are dramatically more sensitive to changes in eIF4E levels and/or...
availability. Such mRNAs are commonly referred to as "eIF4E-sensitive" and encode key proteins which stimulate proliferation (e.g. cyclins, ODC1, c-MYC) and survival (e.g. BCL-2 family members), many of which have been implicated in cancer.\textsuperscript{22-26} eIF4E-sensitive translation is in part mediated by long and complex 5′ untranslated regions (UTR).\textsuperscript{30} These mRNAs have a higher requirement for eIF4A helicase unwinding activity as compared with other cellular mRNAs.\textsuperscript{30,31} eIF4A activity is bolstered significantly within the eIF4F complex,\textsuperscript{32-34} whereby eIF4E acts as a rate limiting factor for eIF4F assembly.\textsuperscript{30} eIF4E-sensitivity of the mRNAs with long and structured 5′UTRs is therefore thought to stem from eIF4E-dependent recruitment of eIF4A and stimulation of eIF4A's activity.\textsuperscript{33} Accordingly, recent ribosome profiling studies in mammalian cells revealed that eIF4A inhibitors preferentially suppress translation of mRNAs that harbor long 5′UTRs enriched in complex structures, including G-quadruplexes,\textsuperscript{35-37} although this was recently disputed for the class of eIF4A inhibitors belonging to the rocaagate family.\textsuperscript{38} Intriguingly, in contrast to mammals, yeast ded1 (ortholog of mammalian DDX3 helicase) but not eIF4A appears to play a predominant role in stimulating translation of mRNAs with highly structured 5′UTRs.\textsuperscript{39} Given the key role of the mTOR pathway and eIF4E in many biologic contexts and human diseases including cancer, neurologic diseases, diabetes and metabolic syndrome, there has been a considerable interest in applying genome-wide approaches to obtain a complete catalog of mRNAs that are sensitive to changes in mTOR activity and thereby eIF4E availability.

**Controversy regarding the repertoire of mRNAs showing mTOR-sensitive translation**

Three recent studies using pharmacological inhibitors of mTOR (including the allosteric mTOR inhibitor rapamycin, active-site mTOR inhibitors and the biguanide metformin, which inhibits mTOR indirectly via AMP-activated kinase-dependent and independent mechanisms) were performed to catalog the transcriptome-wide set of mRNAs showing mTOR-sensitive translation.\textsuperscript{40-42} Unexpectedly, these studies reached radically different conclusions. Two studies using ribosome-profiling suggested that mTOR almost exclusively regulates translation of mRNAs harboring a 5′ terminal oligopyrimidine (5′ TOP) motif.\textsuperscript{43,44} The TOP motif consists of a cytosine (C) directly after the mRNA cap followed by a stretch of 4–15 pyrimidines and is mainly found in mRNAs encoding for components of the translational machinery including ribosomal proteins, poly (A) binding protein (PABP) and eukaryotic elongation factor 2 (eEF2).\textsuperscript{45} The mTOR-sensitivity of TOP mRNAs was recognized over a decade ago,\textsuperscript{44} but this appeared to be largely eIF4E-independent.\textsuperscript{44-46} In contrast, the conclusion that TOP mRNAs would essentially be the only targets of mTOR was unexpected, especially as translation of mRNAs encoding growth, proliferation, survival and tumor-promoting proteins, such as cyclins,\textsuperscript{47} ornithine decarboxylase (ODC1),\textsuperscript{48} vascular endothelial growth factor (VEGF)\textsuperscript{49} or c-MYC\textsuperscript{50} had been shown to be sensitive to alterations in eIF4E levels, which is a major mediator of mTOR-dependent translational control. Moreover, the conclusion drawn by the 2 aforementioned studies,\textsuperscript{40,44} that the effects of mTOR on TOP mRNA translation is chiefly mediated via 4E-BPs was in conflict with previously reported findings which showed that TOP mRNA translation is not eIF4E dependent.\textsuperscript{40} Moreover, a recent report revealed that under conditions when mTOR signaling is modulated by physiologic stimuli, TOP mRNA translation is regulated via an 4E-BP-independent mechanism.\textsuperscript{45} Indeed, it appears that mTOR regulates TOP mRNA translation via La-related protein 1 (LARP1).\textsuperscript{51,52} In stark contrast to ribosome-profiling studies, polyosome-profiling suggested that several non TOP mRNAs including those that were previously identified as eIF4E-sensitive (e.g., cyclins, ODC1) exhibit mTOR-sensitive translation. The cohort of mTOR sensitive mRNAs also contained those encoding mitochondria related proteins.\textsuperscript{42} Indeed, modulation of mitochondria related mRNAs by mTOR was further functionally evaluated which revealed that the mTOR/4E-BP/eIF4E axis coordinates energy expenditure by the mRNA translation machinery with mitochondrial ATP production.\textsuperscript{53} In conclusion, strikingly disparate catalogs of mRNAs that are translated in an mTOR-dependent fashion were captured using ribosome- vs. polyosome-profiling, which suggested biases of these approaches in the detection of mTOR-sensitive mRNAs.

**Ribosome-profiling introduces biases in identification of mTOR-sensitive translation**

Because translational efficiency is primarily regulated at the initiation step, efficiently translated mRNAs are associated with more ribosomes than inefficiently translated mRNAs leading to more proteins being synthesized.\textsuperscript{54} This tenet underpins current transcriptome-wide approaches to study changes in translational efficiency, including polysome- and ribosome-profiling. During polysome-profiling efficiently translated mRNAs (commonly those associated with more than 3 ribosomes) are isolated and quantified using DNA microarrays or, more recently, RNA-sequencing (RNAseq). Polyosome-profiling therefore directly assesses changes in translation efficiency from an “mRNA perspective,” by physically separating efficiently and non-efficiently translated mRNA molecules by ultracentrifugation on sucrose gradients (Fig 1A).\textsuperscript{54} During ribosome-profiling, ribosome protected fragments (RPF), i.e., RNA fragments protected by the ribosome from RNase-mediated degradation, are isolated and quantified using RNAseq (Fig 1B).\textsuperscript{55} Thus, in contrast to polysome-profiling, ribosome-profiling has a “ribosome perspective” wherein translational efficiency is determined indirectly by counting the number of RPFs from both efficiently and inefficiently translated mRNAs. Although several factors affect the performance of ribosome-profiling,\textsuperscript{56,57} this technique holds a great promise by providing unprecedented single nucleotide resolution of ribosome positioning on the mRNA.\textsuperscript{58,59} In contrast, polysome-profiling does not reveal ribosomal location on the mRNA, but it allows isolation of intact mRNAs from the polysomes that can be further studied (see 5′UTR profiling below).

We recently showed that a key difference between polysome- and ribosome-profiling occurs when mRNAs that exhibit different magnitudes in their changes in translational efficiency are studied in parallel. For example, a treatment with a pharmacological inhibitor may alter ribosome association of a class of mRNAs from a mean of 4 to 2 ribosomes, while
others may shift from 7 to 1 ribosomes. The differences in shifts reflect intrinsic properties of these mRNAs and are thus not expected to directly mirror their relative importance for the biology or pathology studied. In ribosome profiling translation efficiency is inferred indirectly based on the number of RPFs, and therefore the size of the shifts in ribosome association will result in directly proportional effects on fold-change estimates (i.e., 2 vs 7-fold from the example above). This effect is much less pronounced in polysome-profiling studies, as herein translation efficiency is estimated directly from mRNAs associated with heavy polysomes (i.e., >3 ribosomes) (Fig. 1A-B). Moreover, mRNAs that show large shifts in translational efficiency and are also very abundant will cause a seemingly global effect on translation. As a result, changes in translation of mRNAs that show less dramatic shifts in translational efficiency are masked. This is because RNAseq applied during polysome- or ribosome-profiling generates relative quantification, whereby changes in translational efficiency are compared with the ‘mean change’ in global translation. Smaller fold-changes of low abundant mRNAs will therefore be strongly down weighted. Altogether, this indicates that ribosome-profiling is biased toward detecting changes in translation of highly abundant mRNAs that exhibit large shifts in polysome association but is considerably less sensitive in identifying those mRNAs with smaller shifts and lower abundance (Fig 1B). A prototypical example of a very abundant mRNA-class that shows a large shift in translational efficiency upon changes in mTOR activity are TOP mRNAs. Many other mRNAs, such as those encoding for cyclins or mitochondria-related proteins, show relatively smaller shifts and are expressed at a much lower level as compared with TOP mRNAs. Thus, application of ribosome profiling to conditions when these 2 mRNA populations change their translational efficiency at the same time (high expression/big shift vs. moderate expression/moderate shift) leads to preferential identification of TOP mRNAs (Fig. 1B). These technical differences, together with the issues related to sequencing depth discussed below, appear to explain why ribosome profiling studies essentially only identified TOP mRNAs as mTOR sensitive, while polysome profiling revealed a more diverse set of genes including TOP and non-TOP mRNAs (e.g., those encoding cyclins and mitochondria-related proteins).

Figure 1. Schematics illustrating properties underlying a bias toward identification of mRNAs that show large shift in translational efficiency as differentially translated when applying ribosome-profiling. (A) Shown in gray is a model UV absorbance profile from a polysome-preparation where ribosome subunits (40S or 60S), monosomes (80S) or polysomes (i.e., mRNAs associated with >1 ribosome) are separated on sucrose gradients by ultracentrifugation. In polysome-profiling, translational efficiency is measured by quantifying the amount of mRNA that is efficiently translated (i.e., associated with >3 ribosomes, which is indicated by a pink dotted line). Transcripts differ in their basal translational efficiency (green). Left panel represents TOP mRNAs that are associated with the heaviest polysomes and are thus more efficiently translated than non-TOP mRNA (e.g., cyclin) which are associated with intermediate polysomes. Upon mTOR inhibition (blue), TOP mRNAs shift more dramatically than non-TOP mRNAs (i.e., shifts are indicated by arrows). (B) In ribosome-profiling, the amount of mRNA fragments protected by ribosomes (RPFs) is quantified using RNAseq and compared between conditions. This generates radically different fold-changes for TOP and non-TOP mRNAs, as the former are much more abundant and exhibit larger shifts. Such fold-changes are directly proportional to the magnitude of the mRNA shifts leading to a bias favoring identification of TOP mRNAs as differentially translated. This bias is more pronounced when fold-change based vs. statistical analysis is performed.
An additional source of bias in ribosome-profiling studies of mTOR-sensitive translation\textsuperscript{30,44} likely stems from the sequencing depth. Optimal RNA sequencing depth allows reliable quantification across all expression levels and conditions.\textsuperscript{62} When performing ribosome-profiling this is an important consideration, as highly expressed genes that are also very efficiently translated (e.g., TOP mRNAs) will dominate the sequencing library and hence constitute a large proportion of the RPF RNAseq reads. Sub-optimal RNAseq depth greatly influences the signal to variance relationship (low number of RNAseq reads is associated with increased noise in the data set), preventing identification of differential translation for less expressed, and less efficiently translated mRNAs.\textsuperscript{50} Indeed, ribosome-profiling studies of mTOR-dependent translation related very low sequencing depth and accordingly almost exclusively detected effects on translation of the most abundant cellular mRNAs including TOP mRNAs.\textsuperscript{40,41,60}

Altogether, the disparity between the repertoires of mTOR-regulated mRNAs observed between ribosome- and polysome-profiling studies appears to stem from technical biases and insufficient sequencing depth which limited detection of mTOR-dependent changes in the translatome in studies which used ribosome-profiling.

### Profiling 5′UTRs unraveled 2 distinct subsets of mTOR-sensitive mRNAs encoding proteins with different cellular functions

mRNAs exhibit different translational properties, which is in part conferred by their 5′UTR features.\textsuperscript{2,22,24,60,63,64} A number of regulatory elements in 5′UTRs are implicated in translational control of gene expression including upstream open reading frames (uORFs), stem loops (e.g., the iron-responsive element; IRE) and internal ribosome-entry sites (IRESes) which allow cap-independent translation (reviewed in Hinnebusch et al.\textsuperscript{30}). In addition, it is thought that the position of these regulatory elements and/or structural features relative to the mRNA cap or the initiation codon, as well as the length of the 5′UTR play a major role in determining translation efficiency (reviewed in Hinnebusch et al.\textsuperscript{30}). Analysis of 5′UTRs, however, relies on data repositories such as RefSeq and UCSC, which provide a wealth of data and tools to understand the genomic contexts of many species. These data repositories, thereby, offer a framework to interrogate and interpret observed changes in translational efficiency in the context of e.g., 5′UTRs. It is widely thought, however, that these databases contain a repertoire of 5′UTRs that may not necessarily reflect those that are expressed in the cell of interest, thereby potentially leading to faulty conclusions about the role of 5′UTRs in translational control. Several initiatives to accurately pinpoint transcription start sites (TSSs) including the FANTOM Consortium (The FANTOM consortium and the RIKEN PMI and CLST [DGT] 2014) or TSS-seq\textsuperscript{65,66} are currently ongoing but information is still missing for most cell lines and tissues, especially if one considers that 5′UTRs in the same cell line may be context dependent due to e.g., stress induced alternative transcription site selection or alternative splicing.\textsuperscript{67} Thus, for precise understanding of the relationship between 5′UTR features and translational control, both TSSs and translational efficiency should be determined in the same cell line. Importantly, ribosome-profiling does not allow for such assessment, as the nuclease digestion degrades both 5′ and 3′ UTRs of mRNA. In contrast, polysome-profiling can be readily coupled with techniques such as Nano Cap Analysis of Gene Expression (nanoCAGE),\textsuperscript{68} an RNAseq library construction method that allows nucleotide resolution mapping of TSSs and, as a result, 5′UTRs. Application of this approach to mTOR-sensitive translation unveiled that nearly 30% of 5′UTRs were substantially shorter than those indicated in the RefSeq database.\textsuperscript{60} Moreover, nanoCAGE revealed 2 distinct non-TOP, mTOR-sensitive mRNA subsets: one whose members harbor long 5′UTRs encoding cell-cycle and survival-promoting proteins and a second whose members have extremely short 5′UTR (< 30 nucleotides) encoding proteins with mitochondrial function. These 2 subsets could not have been separated using current databases, as these suggested radically different 5′UTRs than the ones identified by nanoCAGE for the subset of mitochondrial-related mRNAs with very short 5′UTRs.\textsuperscript{50}

### The distinct subsets of mTOR-sensitive mRNAs require different translation initiation factors and encode for proteins participating in discrete cellular processes

The first subset of mRNAs with long 5′UTRs identified by nanoCAGE (including cyclins, BCL-2, MCL1, BIRC5) supports the long-held idea that downstream of mTOR, changes in eIF4E availability primarily affect mRNAs with long and complex 5′UTRs that critically depend on eIF4A activity for efficient translation.\textsuperscript{30,31} In contrast, mitochondrial-related mRNAs with short 5′UTRs (such as ATP5O, ATP5G1, NDUF6, UQCC2) do not seem to fit into this model.\textsuperscript{60} A subset of mRNAs with short 5′UTRs harboring a Translation Initiator of Short 5′UTR (TISU) element (SAASATGCGGCG, in which S is C or G) which is enriched for mRNAs encoding proteins with mitochondrial functions, however, was previously shown to have eIF4E sensitive but eIF4A insensitive translation.\textsuperscript{69,71} Consistently, we revealed differential requirements of the 2 mTOR-sensitive mRNA subsets: those with long 5′UTR are both eIF4E- and eIF4A-sensitive while those with short 5′UTR are eIF4E-, but not eIF4A-sensitive. Surprisingly, this sensitivity of short mRNAs appeared to be more dependent on the length of the 5′UTR than the presence of the TISU element. Moreover, the dichotomy in eIF4A sensitivity between short and long 5′UTRs of mTOR-sensitive mRNAs appear to in part account for the different responses to mTOR or eIF4A inhibitors. mTOR inhibitors affect translation of mRNAs with both very short (encoding for mitochondria related proteins) and long (encoding for pro-survival proteins) 5′UTRs\textsuperscript{60} which reduces mitochondrial\textsuperscript{63} activity but concomitantly downregulates energy consumption by the translation machinery resulting in metabolic dormancy and a cytostatic effect.\textsuperscript{42} eIF4A inhibitors, on the other hand, display strong cytotoxic effect as they reduce the translation of pro-survival mRNAs with long 5′UTRs, without affecting the translation of short 5′UTR mRNAs encoding for proteins with mitochondrial function, which leads to mitochondria dysfunction and apoptosis.\textsuperscript{60} In addition, eIF4A inhibitors, unlike mTOR inhibitors, reduce
autophagy which removes depolarized mitochondria, thereby further bolstering their pro-apoptotic effects.60

Concluding remarks

In conclusion, analytical and technical biases should be taken into consideration when performing transcriptome-wide analysis of translational regulation. Moreover, the limitations of databases for e.g., 5' UTRs can obscure significant findings due to a lack in accuracy or lack of information for a specific experimental model. Therefore, choosing the appropriate experimental methodology coupled with a tailored analysis can address important gaps in knowledge pertinent to the regulation of translation. Factors that limit these advances include suboptimal study design, lack of appropriate quality control and incorrect statistical methods - which are all inherent difficulties of today's science, and were in fact predicted to undermine systems biology approaches.72 Inappropriate strategies such as limited replication which does not allow application of statistical methods, adopting analytical approaches to identify a set of favorite genes that fit the hypothesis (i.e., cherry picking), or simply a lack of understanding of data analysis methodology can compromise validity of conclusions and study reproducibility.73 We have previously demonstrated that using the ratio between polysome-associated mRNA or RPFPs to cytosolic mRNA (commonly denoted as translation efficiency or TE score) for analysis of differential translation results in spurious correlation thereby favoring false positive and negative findings.74 Furthermore, we showed that using a fold-changes based analysis is similarly inappropriate, especially when assessing differential translation using ribosome-profiling.60 Nonetheless, both approaches are commonly applied, thereby indicating the lack of a consensus regarding optimal data analysis. Similarly, a recent advance in quality evaluation of ribosomal profiling data revealed factors pertinent to the validity of ribosome profiling data that are also likely to skew the interpretation of the results.56 Collectively, a consensus on guidelines for study design, quality control and data analysis appears to be required to improve translatome analysis, and thus to advance the understanding of translation regulation in health and disease.

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