mRNA structure dynamics identifies the embryonic RNA regulome

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RNA folding plays a crucial role in RNA function. However, our knowledge of the global structure of the transcriptome is limited to steady-state conditions, hindering our understanding of how RNA structure dynamics influences gene function. Here, we have characterized mRNA structure dynamics during zebrafish development. We observe that on a global level, translation guides structure rather than structure guiding translation. We detect a decrease in structure in translated regions, and we identify the ribosome as a major remodeler of RNA structure in vivo. In contrast, we find that 3'-UTRs form highly folded structures in vivo, which can affect gene expression by modulating miRNA activity. Furthermore, we find that dynamic 3'-UTR structures encode RNA decay elements, including regulatory elements in nanog and cyclin A1, key maternal factors orchestrating the maternal-to-zygotic transition. These results reveal a central role of RNA structure dynamics in gene regulatory programs.

Introduction

RNAs carry out a broad range of functions, and RNA structure has emerged as a fundamental regulatory mechanism, modulating various post-transcriptional events such as splicing, subcellular localization, translation and decay. RNA probing reagents combined with high-throughput sequencing allow the interrogation of RNA structure in a transcriptome-wide manner in vitro and in vivo. These approaches have identified common features in RNA structures and their regulation at steady-state, suggesting that RNAs tend to be unfolded in vivo. However, the cellular factors that remodel RNA folding in vivo remain unknown.

While the ribosome possesses a constitutive mRNA helicase activity, stable RNA structures found within coding regions can reduce the rate of translation in vitro. Similarly, the intrinsic structure adopted by coding regions instruct the rate of translation in bacteria, where sequences forming stable structures show reduced translation in vivo. In addition, stable structures located in 5'UTR or surrounding the AUG initiation codon repress translation and modulate protein output in bacteria and eukaryotes. However, in plant and yeast, highly structured mRNA in vitro correlated with higher ribosome density and protein output, yet, other studies found that RNA structure in yeast were not correlated with translation efficiency.

Previous studies have focused on the global analysis of RNA structures in cells at steady-state. However, in an organism, critical developmental and metabolic decisions are often made when cells transition between cellular states. Consequently, to understand the relationship between RNA structure, regulation, and function in vivo, it is paramount to determine how RNA structures change in dynamic cellular environments. During the maternal-to-zygotic transition (MZT) in animals, the embryo is initially transcriptionally silent, and gene expression is primarily orchestrated by post-transcriptional regulation of maternally-deposited mRNA translation and decay. These regulatory pathways are central to embryogenesis, controlling mRNA translation, the cell cycle, and the activation of the zygotic genome. Among the set of maternally-deposited mRNAs, the transcription factors nanog, oct4, and sox19b are highly translated. Their combined function is required to activate a large fraction of the zygotic program. One of the first zygotically transcribed RNAs is the microRNA miR-430, which causes translation repression, deadenylation and clearance of hundreds of maternal mRNAs during the MZT. Thus, the MZT provides an ideal system to understand how the entire cellular population of RNA structures is remodeled across changing cellular states, and how these changes may impact gene regulation. Here, we analyze mRNA structural dynamics during zebrafish embryogenesis to investigate the

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relationship between RNA structure and gene regulation in vivo.

Results

Dynamic structure reflects translation changes

We analyzed the structure of the zebrafish transcriptome during the MZT using DMS-seq (Fig 1a), which we validated as an accurate readout of in vivo RNA structure (Extended Data Fig. 1). Our analysis revealed global changes in mRNA structure during the MZT (Extended Data Fig. 1i-k). Changes in translation, analyzed by ribosome footprinting, were correlated with global mRNA accessibility changes in coding sequences (CDS) and 5'-UTRs, but not in 3'-UTRs (Fig. 1b and Extended Data Fig. 1l). We analyzed differentially structured regions over development as 100-nt sliding windows (Fig. 1c) and observed that mRNAs with decreasing rates of translation between 2 and 6 hours post-fertilization (hpf) increase in structure (orange), and those whose

Figure 1. Relationship between mRNA structure and translation during the MZT. a, Schematic view of the profound transcriptomic remodeling that occurs during the maternal-to-zygotic transition. After a transcriptionally silent period (gray), the maternal program (pink) transitions to a zygotic program (blue). b, Correlation between the per-transcript changes in translation efficiency (6-2 hpf) and CDS accessibility (6/2 hpf). c, Methodology used to identify local RNA structure changes between conditions. Differentially structured (DS) sliding windows are defined as those that are significantly changing (P<0.05) based on the Kolmogorov-Smirnov test, and show an increase (>1.1) or decrease (<0.9) in their Gini index ratio. d, In the left panel, structure changes of each 100-nt window (Gini ratio 6/2 hpf) along each transcript (total of 1,273) are shown. Transcripts are ranked by their changes in translation efficiency during the MZT (6-2 hpf). Each transcript region (5'-UTR, CDS, 3'-UTR) has been normalized by its length. In the right panel, cumulative distribution of global CDS accessibility in vivo, showing that highly translated mRNAs exhibit increased CDS accessibility in vivo at 2 hpf, a stage at which the rate of translation spans ~10,000-fold between highly and poorly translated mRNAs. Transcripts have been binned into quintiles according to their translation efficiency. P-value was computed using a one-sided Mann-Whitney U test.
translation increases become less structured (turquoise) (Fig. 1d). These results indicate that translation and mRNA structure are anti-correlated in vivo, in contrast to previous reports. Moreover, we compared the accessibilities of the different mRNA regions (5'-UTR, CDS and 3'-UTR) and found that highly translated mRNAs display a greater accessibility in their CDS ($P=2.3e-118$) and, to a lesser extent, in their 5'-UTRs ($P=1.9e-11$) (Fig. 1e and Extended Data Fig. 2a).

Pre-existing RNA structure differences might determine mRNA translation, where less structured mRNAs would be more accessible and, consequently, translated more effectively (Fig. 2a, upper panel). Alternatively, high translation rates might lead to lower structure in vivo due to constant mRNA unfolding by the ribosome (Fig. 2a, lower panel). To distinguish between these scenarios, we compared the accessibility of transcripts analyzed in vitro, binned by their translation efficiency in vivo (see Methods) (Fig. 2b and Extended Data Fig. 2a). We found that lowly and highly translated mRNAs show similar CDS accessibility in vitro, distinct from in vivo (Fig. 2b, c and Extended Data Fig. 2a-c), suggesting that the differences in mRNA accessibility between highly and lowly translated mRNAs are not intrinsic to the nucleotide sequence. While very stable structures in the 5'-UTR clearly disrupt translation, we observed no correlation between translation and structure of 5'-UTRs or the AUG initiation codons in vitro (Fig. 2d, e and Extended Data Fig. 2d-g). Thus, on a global level, RNA structure in 5'-UTR and CDS regions is not a major determinant of translation across the transcriptome in a vertebrate embryo.

Figure 2. Globally, the mRNA intrinsic structure is not a main driver of translation. a, Schematic representation of the two models that can explain the observed anti-correlation between translation efficiency and RNA structure in CDS regions in vivo. In the first model, pre-existing CDS RNA structures would drive translation (discarded by our analysis, upper panel). In a second model, translating ribosomes would be responsible for the unfolding of CDS RNA structures, (consistent with our analysis, lower panel). b, Cumulative distributions of global CDS RNA accessibilities in vitro at 2 hpf, displaying no difference among mRNAs with different translation efficiency. Transcripts have been binned into quintiles according to their translation efficiency. P-value was computed using one-sided Mann-Whitney U test. c, Scatterplot between in vivo and in vitro CDS accessibilities at 2 hpf. Note higher accessibility in vivo for highly translated mRNAs (red dots) and lower accessibility in vivo for poorly translated mRNAs (blue dots). d, Schematic view of the two models where RNA structure at the AUG initiation codon is either the cause (top) or the result (bottom) of mRNA translation. e, Correlation between translation efficiency and the structure at AUG regions, for both in vitro (left) and in vivo (right) conditions.
Ribosomes shape mRNA structures

The helicase activity of the ribosome unfolds RNAs in vitro during translocation\textsuperscript{16-18}. However, in yeast there is no correlation between RNA structure and translation efficiency (TE)\textsuperscript{13}, suggesting that ribosome activity does not account for the differences between mRNA structures in vivo and in vitro. To determine the role of ribosomes in structural remodeling in vivo, we examined RNA structural changes in embryos incubated with inhibitors of translation initiation (pateamine A; PatA)\textsuperscript{36,37} or elongation (cycloheximide; CHX)\textsuperscript{38}. Ribosome profiling of 64-cell embryos (2 hpf) treated with PatA revealed a prominent decrease (~34-fold) in translation efficiency compared to untreated embryos (Fig. 3a, b). mRNAs most sensitive to PatA treatment had longer 5'-UTRs ($P=2.1e-42$), consistent with a loss of the PatA target eIF4A during scanning by the pre-initiation complex\textsuperscript{36,37} (Fig. 3a, c). In contrast, mitochondrial transcripts showed no change in translation (Fig. 3d), consistent with the observation that their translation is independent of eIF4A\textsuperscript{39}. In PatA-treated embryos, we found an overall decrease in accessibility of the CDS regions of highly translated mRNAs ($P=5.3e-35$) (Fig. 3e and Extended Data Fig. 3a). Similarly, regions with higher ribosome footprint densities form more stable RNA conformations after PatA treatment, similar to those observed in vitro (Fig. 3f and Extended Data Fig. 3b, c). In contrast to PatA-treated embryos, CHX treatment did not decrease average CDS accessibility (Extended Data Fig. 3d).

Figure 3. Ribosomes promote alternative mRNA structures in the early embryo. a, Schematic view of the PatA translation inhibitory mechanism. PatA reduces the levels of functional eIF4F initiation complex, which is essential for cap-dependent translation, by trapping eIF4A onto mRNAs and ectopically enhancing its RNA helicase activity. b, Distribution of translation efficiencies (TE) of PatA-treated (blue) and untreated (gray) embryos. c, Correlation between 5'-UTR length and sensitivity to PatA treatment. The least and most sensitive to PatA mRNA subgroups correspond to the top and bottom quintiles of ΔTE (PatA-untreated), respectively. $P$-value was computed using a one-sided Mann-Whitney $U$ test. Box spans first to last quartiles and whiskers represent 1.5× the interquartile range. d, Scatter plot of the per-transcript ribosome footprints in PatA-treated versus untreated embryos. Mitochondrial genes (red), which are unaffected by PatA treatment, remain unchanged. e, Schematic representation depicting the impact of PatA, along with its corresponding cumulative distribution of CDS accessibility ratios (untreated/PatA). Note the decrease in accessibility (red arrow) in highly translated mRNAs in PatA compared to untreated. $P$-value was computed using a one-sided Mann-Whitney $U$ test. f, Arc plots of DMS-seq guided RNA secondary structure predictions of a high ribosome occupancy region found in the basigin gene, for in vivo (yellow) in vitro (green), untreated (yellow), PatA-treated (purple) or CHX-treated (blue) samples. Each arc represents a base pair interaction. g, Arc plots of DMS-seq guided RNA secondary structure predictions of the full-length cnot8 mRNA using SeqFold. The CDS region has been shadowed in gray. h, i, Principal Component Analysis biplots of the SeqFold mRNA structure predictions of the 4 tested conditions (untreated, in vitro, PatA-treated and CHX-treated), of CDS (h) and 3'-UTR (i) regions.
Data Fig. 3d), suggesting that even when the ribosomes are stalled, the residency of the ribosome on the mRNA during translation plays a major role in remodeling mRNA structure. Therefore, translation initiation and ribosomal entry are needed to locally favor the formation of less stable alternative RNA structures.

To evaluate the impact of the ribosome remodeling activity on mRNA structures, we compared the RNA conformations adopted by full-length transcripts in each probing condition using SeqFold. We found that the same transcript could favor different conformations depending on its translation status (Fig. 3g and Extended Data Fig. 3e). Principal component analysis (PCA) on the structural data from 1,143 transcripts (see Methods) revealed that in absence of ribosomes (in vitro or in the presence of PatA), individual mRNAs exhibit similar CDS and 3’-UTR structures, distinct from those formed in the presence of ribosomes (untreated and CHX) (Fig. 3h, i). Finally, translation of upstream open reading frames (uORF) located in 5’-UTRs leads to a restructuring of 5’-UTR structures in vivo (Extended Data Fig. 4). Thus, the ribosome increases mRNA accessibility, and promotes alternative RNA conformations throughout the mRNA.

**Translation-dependent mRNA structure**

We then evaluated translation control mechanisms during the MZT for their impact on mRNA structure regulation (Fig. 4a). Polyadenylation increases translation efficiency during early embryogenesis (Fig. 4a). We found that mRNAs with longer poly(A) tails show higher accessibilities in the CDS at 2 and 4 hpf (rho of 0.42 and 0.36, respectively), but not in the 3’-UTR (Extended Data Fig. 5a). This effect was reduced following PatA treatment, and lost in vitro (rho of 0.29 and 0.01, respectively) (Extended Data Fig. 5b), consistent with a translation effect specific to the CDS. Moreover, mRNAs with extended poly(A) tails during the MZT displayed a decrease in structure, while deadenylated transcripts showed an increased structure (Fig. 4b, c). These results suggest that changes in polyadenylation during development influence mRNA structure, an effect that depends on the translation of the mRNA.

To understand the effect of microRNA-mediated repression on mRNA structural changes, we analyzed miR-430, which regulates hundreds of maternal mRNAs during the MZT (Fig. 1a, 4a). The miR-430 targets are initially translationally repressed at 4 hpf without significant signs of decay. If ribosomes play a crucial role in shaping mRNA structure in vivo, mRNAs targeted by miR-430 should reach an intermediate state at 4 hpf, characterized by an increase in structure due to a depletion of ribosomes. To test this hypothesis, we monitored structural changes in 483 miR-430 targets and a control set of 483 randomly selected mRNAs. We observed a global increase in structure (orange) for miR-430 targets compared to the control set, which is characterized by a decrease in structure (turquoise) between 2 and 4 hpf (Fig. 4d, e). These results support a central role of the ribosome in orchestrating mRNA structure in the cell, suggesting that biological processes regulating translation can have a broad impact on directing mRNA folding.

**3’-UTRs have a distinct folding landscape**

We find that zebrafish transcripts are globally more accessible in vivo than in vitro (Extended Data Fig. 6a). To investigate whether this global unfolding was uniformly distributed along the transcript, we performed a metagene analysis of each structure, for both in vivo and in vitro conditions, using a sliding-window approach (Fig. 1b). Our results reveal that regions with decreased structure in vivo are mainly found in the 5’-UTR and CDS regions of highly translated transcripts (Fig. 5a, b and Extended Data Fig. 6b), in agreement with the remodeling role of the ribosome. Conversely, 3’-UTRs show increased structure in vivo. This increase in RNA structure in vivo doesn’t seem to result primarily from RNA binding protein footprints, since no decrease in accessibility was observed over the binding sites of an AU-rich elements binding protein, KHSRP, and surrounding regions recognized by the Ago2/miR-430 complex (Extended Data Fig. 7). This increase in structure is anti-correlated with translation efficiency, and spreads into the CDS of poorly translated mRNAs, suggesting that the increased structure is specific to ribosome-free regions (Fig. 5b and Extended Data Fig. 6b).

To evaluate the importance of the 3’-UTR RNA folding landscape in the cell, we analyzed the impact of RNA structures on miR-430 activity. Since the accessibility of miRNA binding sites has been shown to affect their activity, we analyzed how miRNA targeting efficiency correlated to the stability (ΔG) of the predicted structure in silico, in vitro and in vivo (Fig. 5c, d). We found that the stability of the RNA structure (ΔG) based on in vivo probing is the best predictor of miR-430 activity (rho = -0.70), outperforming in silico (rho = -0.39) and in vitro (rho = -0.57) structure predictions. Consistent with these
results, endogenous miRNA targets with higher free energy in vivo were more strongly regulated by miR-430 than those with lower free energy (Fig. 5e, f, i, j and Extended Data Fig. 8). To confirm the regulatory role of those in vivo specific RNA conformations, we compared the activity of endogenous targets with different free energy and analyzed the effect that altering their structure had on their regulation. To this end, we quantified the repression of a GFP reporter mRNA in wild type and mutant embryos where the miR-430 locus was deleted (MZmiR430). As a control, we co-injected a DsRed reporter mRNA not targeted by miR-430. Modifying the stability of the in vivo structure in the target site altered the regulatory strength of each site. For example, destabilizing the in vivo structure of the rab33ba target site, which is poorly regulated in vivo (Fig. 5f), increased miR-430-mediated regulation (Fig. 5g, h). Conversely, stabilizing the in vivo structure of
fam171a1, which is strongly regulated by miR-430 (Fig. 5j), abolished miR-430 activity (Fig. 5k, l). Altogether, these results demonstrate that there are increases in 3'-UTR structures in vivo, which can regulate gene expression by modulating miRNA activity.

Dynamic 3'-UTRs enriched in functional elements

During the MZT, maternally-deposited factors (Nanog, Oct4 and SoxB1) mediate activation of the zygotic genome\(^{28}\), after which different programs mediate post-transcriptional regulation of maternal mRNAs (Fig. 1a). This process is required for the embryo to transition from the maternal to the zygotic state, and is characterized by changes in the transcriptional and epigenetic landscapes, cellular motility and the cell cycle, with an extension of G1 and G2\(^{27}\).
Analysis of the structurally dynamic regions in 3'-UTRs between 4 and 6 hpf revealed an enrichment for conserved sequences (Fig. 6a and Extended Data Fig. 9a), suggesting a potential role in the post-transcriptional regulation of the maternal mRNAs. To test this hypothesis,
we performed a parallel reporter assay (RESA) to measure the regulatory role of 71 different 3'-UTR regions derived from maternal mRNAs subject to decay during the MZT (Fig. 6b). Out of these 71 regions, 53 contained dynamic RNA structures, whereas 18 showed no structural changes during MZT. These sequences, inserted into the 3'-UTR of a reporter gene, were in vitro transcribed and injected into 1-cell stage embryos. Then, the abundance of each reporter was measured by high-throughput sequencing at 2 and 6 hpf (Fig. 6b). To identify zygotic-dependent regulation, we compared the levels of each fragment in the presence and the absence of zygotic transcription (i.e. with and without alpha-amanitin). Quantification of mRNA reporters identified decay elements in the 3'-UTR of the cyclin A1 (ccna1) and nanog mRNAs and an enrichment of destabilizing elements in regions that change in structure during development compared to regions that do not change (P=0.025) (Fig. 6c, d, and Extended Data Fig. 9b-e). These results suggest that regions with dynamic structures are enriched in functional regulatory elements.

One such dynamic regulatory region is located in the ccna1 mRNA. Cyclin A1 controls the cell cycle and is actively degraded during the MZT. Time-course analyses of mRNA expression during the MZT using RNA-seq reveal that ccna1 mRNA decays in wild type embryos but not in embryos where zygotic transcription is blocked with alpha-amanitin (Fig. 6c and Extended Data Fig. 9f). The structurally dynamic region of ccna1 tested in RESA also possesses a decay activity that relies on the activation of the zygotic program (Fig. 6f). This is consistent with the regulation of ccna1 in vivo, and in the change in cell cycle length after zygotic genome activation. Another dynamic region with regulatory activity resides in the nanog mRNA, a transcription factor required for genome activation and miR-430 expression, and which is degraded after zygotic genome activation (Fig. 6e and Extended Data Fig. 9f). We predicted the secondary structure of the nanog 3'-UTR at 2 and 6 hpf, using in vivo DMS-seq accessibilities as constraints (Fig. 6g). We identified two domains with constant structure (I and III), flanking a region with differential structure (II). This domain (II) overlaps with the dynamic region that had regulatory activity in vivo (Fig. 6d, h). To test the regulatory activity of this region in the context of the 3'-UTR, we compared the regulation provided by the wild type nanog 3'-UTR to a deletion mutant reporter that disrupted the stem region (II) with a 20-nt deletion (Fig. 6g, h). The full-length nanog 3'-UTR had a decay activity similar to that of the differentially structured region tested in RESA (Fig. 6h). However, a deletion of 20 nucleotides in domain II stabilized the reporter mRNA, suggesting that this region is required to confer regulation in vivo (Fig. 6g, h). Altogether, these results demonstrate that tracking mRNA structure dynamics throughout biological transitions can reveal functional elements in 3'-UTRs.

**Discussion**

Here we examine how RNA structure dynamics change during the MZT, a universal regulatory transition in animal embryogenesis. Our study reveals that 3'-UTRs and untranslated mRNAs are more structured in the embryo than in vitro. Dynamic 3'-UTR structures in vivo correspond to novel regulatory regions controlling maternal mRNA decay. Lastly, we show that the ribosome is one of the principal factors unwinding the mRNA, and while structure does not emerge as a global regulator of translation, translation is a major driver shaping the mRNA structure landscape during early embryogenesis.

Our demonstration that untranslated mRNAs and 3'-UTRs are more structured in vivo than in vitro (Fig. 5a, b and Extended Data Fig. 6b) runs contrary to work showing that mRNAs are mostly unfolded in the cell13,15. Previous in vitro and in vivo analyses of RNA structures have been carried out in yeast, where the average 3'-UTR length is much shorter than that of vertebrate mRNAs13. In the absence of constitutive unwinding by the ribosome, both molecular crowding and RNA binding proteins are likely to favor higher-order RNA structures46,47. Remodeling of mRNA structure can also affect regulatory structural elements modulating 3'-end processing and mRNA stability48. Our data support a view where cellular factors explicitly affect 3'-UTR structures, modulating the regulatory activity of miRNAs and gene expression during vertebrate development.

Our analysis of dynamic mRNA structures identifies regulatory regions during the MZT in the cyclin A1 and nanog mRNAs, two maternal factors that are required to regulate the cell cycle and zygotic transcription, and that are rapidly degraded during the MZT. We find that the characterization of structurally dynamic regions during cellular or developmental transitions represents a means of identifying potential regulatory elements in vivo (Fig. 6b-d and Extended Data Fig. 9b, c).

Global RNA unwinding in yeast is an active process13. The cellular factors responsible
for this structural remodeling and the specific regions that are remodeled remain unknown. While the ribosome has been shown to have a helicase activity in vitro, previous experiments in exponentially growing yeast do not reveal a relationship between translation and RNA unfolding. Here, we have followed the dynamics of mRNA and show that the ribosome is a major engine for mRNA structure remodeling, promoting alternative RNA conformations for thousands of endogenous transcripts in vertebrate cells. This effect is also observed in 5'-UTRs containing translated uORFs, indicating that the ribosome helicase activity has the potential to regulate elements that rely on specific RNA conformations. Several mechanisms regulating mRNA translation, such as changes in poly(A) tail length and microRNA-mediated repression, also result in the remodeling of mRNA structure during embryogenesis.

RNA structure has generally been thought to have a major effect on mRNA translation rates, based on early experiments in bacteria and eukaryotes, where strong hairpin structures in mRNAs reduce translation. Here, we have demonstrated precisely the opposite effect, i.e. translation modulates mRNA structure, and mRNA structure does not have a global effect on translation during embryogenesis (Fig. 2 and Extended Data Fig. 2). While individual examples in eukaryotes show that stable 5'-UTR structures can block translation, at the level of the transcriptome, complex eukaryotic cells have efficient helicase activities that greatly diminish the global effect of 5'-UTR structure on translation. We propose that eukaryotic systems have evolved powerful helicases that, together with sophisticated mechanisms to regulate translation, such as uORFs, miRNAs, RNA-binding proteins, and longer 3'-UTRs, weaken the transcriptome-wide effects that 5'-UTR and CDS structures might have on translation. Our results provide a conceptual shift in our understanding of the relationship between mRNA structure and translation, and suggest that on a global level, translation guides structure rather than structure guiding translation.

Supplementary Information is available in the online version of the paper.

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Author Contributions JDB and AJG conceived the project. JDB performed the experiments. JDB, EMN and CEV performed data processing. VY identified regulatory element in the nanog 3'-UTR and built the reporter constructs. CT performed the KHSRP iCLIP experiment. JDB and EMN performed data analysis and, together with AJG, interpreted the results. AJG supervised the project, with the contribution of MK. JDB, EMN and AJG wrote the manuscript with input from the other authors.

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References


Extended Data Figure 1. DMS-seq controls and mRNA structure changes during the MZT. To determine how mRNA structure changes during cellular transitions in vivo, we analyzed the structure of the zebrafish transcriptome during the MZT using DMS-seq (at 2, 4 and 6 hours post-fertilization, hpf). DMS methylates accessible N1 of adenine and N3 of cytidine bases in single-stranded regions, at the end of a stem, and at base pairs flanking G-U wobble interactions, causing reverse transcriptase drop-off during reverse transcription. The truncated RNAs are then captured, analyzed by high-throughput sequencing and normalized per-transcript, providing per-nucleotide accessibility values between 0 and 1 (see Methods). Three lines of evidence indicate that we can effectively probe RNA structure during embryogenesis. First, we observed a robust and reproducible enrichment of reads mapping to A and C bases (74-83%) compared to DMS-untreated samples (43%) (c), as well as reproducible counts between biological replicates (r of 0.989) (d, e). Second, the DMS-seq accessibility profile of the *Tetrahymena* ribozyme, used as an exogenous control, was in close agreement (P=6.3e-18) with the previously reported structure (f). Third, analysis of conserved secondary structures, such as those found in *dgcr8*89, *selt1a* and *selt2* mRNAs showed that paired A and C bases exhibited lower accessibilities than single stranded regions (P=1.8e-05) (g, h). These results indicate that our DMS-seq analysis provides a robust transcriptome-wide map of RNA structure dynamics in a vertebrate embryo. a, Schematic view of in vivo and in vitro genome-wide DMS probing. Red stars represent DMS-modified nucleotides. b, Cartoon shows the DMS-seq protocol. Dashed lines highlight the second size selection capturing reverse transcription products that prematurely stop due to DMS modifications. c, Proportion of DMS-seq reads from DMS-modified nucleotides in vivo at 2 hpf (+DMS), compared to untreated (no DMS) and the composition of the whole transcriptome. In vivo DMS treatment markedly enriches A/C modified sites, and is increased when nucleotides with minimum coverage...
of 5 RT-stops/nt are considered (see Methods). **d, e**, DMS-seq data is reproducible across biological replicates at a per-transcript (**d**) and per-nucleotide (**e**) resolution *in vivo* at 2 hpf. **f**, Validated secondary structure of the *Tetrahymena* ribozyme (Rz) (adapted from Walter et al.70). In the zoomed panels (corresponding to regions P2, P9 and P4-5-6), DMS-seq accessibilities have been overlaid (from white to red) onto the known secondary structures. In the upper middle part of the figure, boxplot represents consistency between accessibilities and Rz pairing status. P-value computed using a one-sided Mann-Whitney U test. Box spans first to last quartiles and whiskers represent 1.5× the interquartile range. **g**, *in vivo* accessibility profile of the dgcr8 mRNA. Red and turquoise bars correspond to A and C bases, respectively. Accessibilities corresponding to annotated conserved secondary structures (gray rectangles) are highlighted. The detailed secondary structure is also shown, where *in vivo* accessibilities have been overlaid for A and C bases (from yellow to red), while Gs and Us are depicted in white. **h**, Box plot comparing the distribution of *in vivo* accessibilities and the pairing status of A and C bases found in three conserved secondary structures used as controls (selt1a, selt2 and dgcr8). P-value computed using a one-sided Mann-Whitney U test. Box spans first to last quartiles and whiskers represent 1.5× the interquartile range. **i**, Comparisons of per-transcript DMS-seq accessibilities between replicates (left) and across developmental stages (right). **j**, Proportion of differentially DMS-seq accessibilities between replicates (left) and across developmental stages, expressed as the percentage of differentially structured windows -relative to the total windows analysed- (black), and the percentage of mRNAs that contain these windows –relative to the total transcripts present in both stages- (gray). **k**, Proportion of differentially structured windows that increase (orange) or decrease (turquoise) in RNA structure, for each pairwise comparison, based on the directionality of their change in Gini index. The direction of change is defined with respect to the latest developmental stage, for each pairwise comparison. **l**, Correlation between the per-transcript changes in translation efficiency and accessibility for each transcript region (5'-UTR, CDS and 3'-UTR) between 2 and 6 hpf.
Extended Data Figure 2. Ribosomes unwind RNA structures in 5'-UTR and CDS, including the region surrounding the AUG. a, Cumulative distributions of global accessibilities of the 50 nucleotides upstream of the AUG, CDS or 50 nucleotides downstream of the STOP codon are shown for in vivo (top), in vitro (middle) and in vivo vs in vitro (bottom) samples at 2 hpf. mRNAs have been binned in quintiles according to their translation efficiency (TE) and colored from blue (lowest TE) to red (highest TE). P-values have been computed using a two-sided Wilcoxon signed-rank test and a one-sided Mann-Whitney U test, for comparisons within and across translational statuses, respectively. b, c, Correlation between translation efficiency and the accessibility of CDS (left) and 3'-UTR regions (right), for both in vivo (b) and in vitro (c) conditions. d, Cartoon shows the region surrounding the AUG initiation codon used to predict RNA secondary structure and highlights the Kozak region (5'-NNNNAUGNNN-3'). e, Correlation between translation efficiency and the accessibility at the AUG initiation codon for both in vivo (left) and in vitro (right) conditions. f, The RNA secondary structure and the Gibbs free energy (ΔG') were computed for each AUG initiation codon. The energy required to unfold the Kozak region (ΔΔG) was calculated by subtracting the ΔG' from the energy of the structure where the Kozak sequence is forced to be single-stranded (ΔG''). g, Correlation between translation efficiency and the energy needed to unfold and make accessible the Kozak sequence (ΔΔG) for both in vivo (left) and in vitro (right) conditions. No correlation is observed in both cases, suggesting that the structure of the Kozak sequence is not a major factor driving translation in the early embryo.
Extended Data Figure 3. Ribosomes promote alternative RNA conformations with reduced stability in the cell. 

**a**, Cumulative distributions of 5'-UTR, CDS and 3'-UTR global accessibilities in PatA-treated and untreated samples. *P*-values have been computed using a two-sided Wilcoxon signed-rank test. 

**b**, Arc plots of DMS-seq guided RNA secondary structure predictions of high ribosome occupancy regions found in the *nsun2* and *ddit4* genes, for *in vivo* (yellow) *in vitro* (green), untreated (yellow), PatA-treated (purple) or CHX-treated (blue) samples. Each arc represents a base pair interaction. 

**c**, Pairwise comparisons of Gibbs free energy differences (∆∆G; untreated-*in vitro*, PatA-*in vitro* and CHX-*in vitro*) of windows with high ribosome footprint densities (gray) and without ribosome footprints (white). Positive ∆∆G indicates a less stable structure in the first condition (untreated, PatA or CHX), compared to *in vitro*, whereas negative ∆∆G indicates a more stable structure in the first condition, compared to *in vitro*. Note the decrease in stability for structures formed by ribosome-rich regions (gray) in the untreated and CHX-treated samples, but not in the ribosome-free PatA-treated sample or in regions without ribosome (white). Violin plot features a kernel density estimation and lines represent the quartiles of the distribution. *P*-values have been computed using a two-sided Wilcoxon signed-rank test and a one-sided Mann-Whitney *U* test.
Whitney $U$ test, for comparisons across probing conditions for a same group and between groups within a specific condition, respectively. d, Cartoon representation depicting the impact of CHX, along with its corresponding cumulative distribution of CDS accessibility ratios (untreated/CHX) for highly and lowly translated mRNAs at 2 hpf. Note the minor increase in accessibility in lowly translated mRNAs following CHX treatment (blue arrow). P-value was determined using a one-sided Mann-Whitney $U$ test. e, Arc plots of DMS-seq guided RNA secondary structure predictions of the full-length ctcf, maea and nsun2 mRNAs using SeqFold40. The CDS region has been shadowed in gray.

Extended Data Figure 4. uORF translation remodels 5’-UTR structure. a, Schematic view of the 5’-UTR RNA structure remodeling that occurs upon translation of upstream open reading frames (uORF). b, Cumulative distributions of in vivo (top) and in vitro (bottom) uORF accessibilities for highly (red) and lowly (blue) translated uORFs. $P$-values were determined using one-sided Mann-Whitney $U$ tests. c, Distributions of Ribo-seq reads (top panel) and accessibility differences (in vivo – in vitro) (bottom panel) for the cdc25 5’-UTR, which contains two translated uORFs (highlighted in gray). Increased accessibility in vivo is shown in red, whereas decreased accessibility in vivo is shown in blue. d, Arc plot depicting the predicted cdc25 5’-UTR secondary structure, guided by either in vivo (yellow) or in vitro (green) DMS-seq accessibilities. Each arc represents a base pair interaction.
Extended Data Figure 5. Poly(A) tail length-dependent mRNA structure remodeling depends on translation. a, Comparison of per-region accessibilities (CDS and 3'-UTR) and per-transcript poly(A) tail length, at 2 hpf (left panels) and 4 hpf (right panels). b, Comparison of CDS accessibilities and per-transcript poly(A) tail lengths in PatA-treated (left) and in vitro refolded samples (right) from 2 hpf embryos.
Extended Data Figure 6. Differences between in vivo and in vitro RNA structures are not uniformly distributed along transcripts. a, Cumulative distributions of global 5'-UTR, CDS and 3'-UTR accessibilities in vivo and in vitro. Only transcripts with a minimum of 85% coverage of As and Cs and with at least 10 reads in average per As and Cs for both in vivo and in vitro DMS-seq experiments are shown. P-values were computed using two-sided Wilcoxon signed-rank tests. b, Distribution of differentially structured windows along the transcripts, comparing in vivo and in vitro conditions. Windows with increased structure in vivo are depicted in orange, whereas those with decreased structure in vivo are shown in turquoise. Each transcript region (5'-UTR, CDS, 3'-UTR) has been normalized by its length, as well as by total number of windows analyzed in each region. Transcripts have been binned into quintiles based on their translation efficiency, which has been determined using ribosome profiling data.
Extended Data Figure 7. DMS-seq signal provides information on the RNA conformation favored upon the binding of two different trans-factors. While a high DMS-seq signal implies a single-stranded conformation, a low DMS-seq signal signifies either an RNA-RNA or RNA-protein interactions involving the Watson-Crick face of the base\(^7\). To test whether DMS probing is mainly analyzing RNA-RNA pairing versus protein footprinting in vivo, we analyzed the structure over two types of regions, one bound by the RNA binding protein KHSRP identified by iCLIP and another occupied by the Ago2/miR-430 complex. We find that KHSRP-bound 3'-UTR regions are more accessible in vivo, compared to control regions from the same transcripts, and comparable to in vitro, suggesting that KHSRP binding helps to maintain a single-stranded RNA conformation in the cell. In addition, decreased accessibility was observed at miR-430 seed regions when the Ago2/miR-430 complex is present, but not in the flanking sequences nor in absence of miR-430 expression, therefore measuring seed-miRNA interactions. Analyses from two different types of trans-factors (KHSRP and Ago2/miR-430 complex) suggest that our approach measures the RNA structure promoted by those factors rather than their footprints limiting DMS’s access to RNA. a, Sequence logo of KHSRP identified using iCLIP. b, Cartoon depicting the binding status of KHSRP to its target in vitro and in vivo. c, Comparison of the accessibility ratio (in vivo / in vitro) of KHSRP-bound regions (cyan) and matching unbound controls (gray), suggesting that KHSRP binding helps to maintain a single-stranded RNA conformation in the cell. KHSRP binding sites were determined using iCLIP experiments (see Methods) and limited to those found within 3'-UTRs. Unbound control regions originate from 3'-UTR targeted by KHSRP, overlap with less than 25% of the binding site and possess the same length than the binding site. P-value computed using one-sided Mann-Whitney U test. Box spans first to last quartiles and whiskers represent 1.5× the interquartile range. d, Schematic representation of the impact of KHSRP binding on RNA structure in vivo keeping bound-RNA in a single-stranded conformation. e, Cartoon depicting the binding status of the Ago2/miR-430

\(m\)
complex to its target in various conditions. f, Comparison of accessibility ratios (across conditions and time points) of miR-430 seeds (cyan) and matching controls (gray). miR-430 seeds correspond to any 8- and 7-mers miR-430 binding sites found in the 3'-UTR of miR-430 targets (see Methods). Control regions of 8-nt were randomly chosen within the miR-430 seed containing 3'-UTRs (see Methods). MiR-430 seed and control accessibilities were calculated by averaging the accessibility of A and C bases found in each 8-nt sequence. P-values computed using one-sided Mann-Whitney U tests. Box spans first to last quartiles and whiskers represent 1.5× the interquartile range g, Meta-analysis of average accessibilities at 2 and 4 hpf (top panels) and nucleotide content (bottom panels) at each position of miR-430 seed (n=74, left panels) and control (n=75, right panels) regions. Each region corresponds to a 100-nt window centered on a 8-nt miR-430 seed or control sequences highlighted in cyan and gray, respectively. Dotted lines correspond to the average accessibility of the metaplot for each developmental stage. Error bars denote s.e.m.

Extended Data Figure 8. In vivo specific 3'-UTR structures impact miRNA activity and gene expression. a, b, In vivo and in vitro predicted secondary structures and stabilities (∆G) of the 200-nt region centered on the miR430 target site found in non miR-430 targets (rab33ba, zgc:55733 and pgk1; a) and miR-430 targets (fam171a1, znf706 and rtkn2a; b). Regions complementary to the miR-430 seed and its complementary binding region within the 3'-UTR are shown in cyan and red, respectively. Changes in mRNA abundance during the MZT of the endogenous transcripts in wild type conditions (black) or in conditions where miR-430 activity is inhibited by a tinyLNA complementary to the miR-430 seed (green) are shown for each gene. MiR-430 targets are characterized by a gain in stability at 6 hpf when miR-430 activity is reduced (-miR-430) (b).
Extended Data Figure 9. Dynamic 3'-UTR structures reveal decay elements during the MZT. a, Enrichment of conserved sequences in 3'-UTR regions with a dynamic structure between 4 and 6 hpf (6v4), compared to all 3'-UTR regions analyzed. Dynamic structures correspond to those with a KS test P-value <0.05. Box spans first to last quartiles and whiskers represent 1.5× the interquartile range. P-value computed using a one-sided Mann-Whitney U test. b, Decay activity of regions with a dynamic structure (left column, red) and a static structure (right column, green) originated from the same 3'-UTR, calculated from the RESA experiment. c, Decay (blue) and stabilizing (dark orange) activities of all the 200-nt 3'-UTR regions with sufficient read coverage from the RESA experiment (see Methods). Genes labeled in red correspond to those that have regions with a dynamic structure during the MZT, whereas genes labeled in green correspond to those with no structural change. Note the enrichment of regions with a dynamic structure among the decay elements (blue). d, Per-transcript KS test P-value profile (top) and conservation (bottom) for the igf2bp3 mRNA, comparing 4 and 6 hpf. The profiles have been computed by analyzing 100-nt sliding windows throughout the transcript. Examples of conserved 3'-UTR regions with a dynamic structure (red) and without structural change (green) are highlighted. P-values between 1 and 0.2 are shown in gray, between 0.2 and 0.05 are shown in yellow and <0.05 are shown in red. The horizontal dotted gray line indicates P-value equal to 0.05. The vertical black lines delineate the AUG and STOP codons of the CDS. e, Quantification of the decay activity for the dynamic (red) and non-changing (green) structures found in the igf2bp3 3'-UTR, as calculated by RESA. Data are represented as the mean ± SD. Student t-test P-values are indicated as ** <0.01. f, Genome tracks of RNA-seq experiments representing mRNA levels of the ccna1 (top) and nanog (bottom) transcripts at 2, 4 and 6 hpf in wild type (left) and alpha-amanitin (right) conditions. Alpha-amanitin inhibits zygotic genome activation, highlighting the implication of zygotic factors in the clearance of ccna1 and, to a lesser extend, nanog mRNAs.
Methods

Data Availability. Raw reads that support the findings of this study will be publicly available in the Sequence Read Archive under SRP114782 upon acceptance in a peer-reviewed journal. Meanwhile, we will be glad to share our raw data with anyone upon request.

Zebrafish maintenance. Wild-type zebrafish embryos were obtained through natural mating of TU-AB strain of mixed ages (5-18 months). Mating pairs were randomly chosen from a pool of 60 males and 60 females allocated for each day of the month. Fish lines were maintained following the International Association for Assessment and Accreditation of Laboratory Animal Care research guidelines, and approved by the Yale University Institutional Animal Care and Use Committee (IACUC).

In vivo and in vitro DMS modification. For in vivo DMS modification, 150 embryos at the specified stage were transferred to 5 mL eppendorf tubes containing 400 µL of system water from the fish facility. 100% DMS (Sigma-Aldrich) was diluted in 100% ethanol to obtain a 20% DMS stock solution. The DMS stock solution was used to generate a master mix containing 6% DMS and 600 mM Tris-HCl pH 7.4 (AmericanBio) in system water from the fish facility. The master mix solution was immediately mixed vigorously and 200 µL was added to each tube to reach a final concentration of 2% DMS and 200 mM Tris HCl pH 7.4. Embryos were incubated at room temperature for 10 min with occasional gentle mixing. The DMS solution was then quickly removed from the tubes and the embryos were flash frozen in liquid nitrogen. Frozen embryos were thawed and actively lysed with 800 µL of TRIzol (Life Technologies) supplemented with 0.7 M β-mercaptoethanol (Sigma-Aldrich) to quench any remaining trace of DMS. After 2 min incubation, TRIzol was added to reach a final volume of 4 mL and total RNA extracted following the manufacturer’s protocol. Poly(A)+ mRNAs were isolated using oligo (dT)25 magnetic beads (New England BioLabs) following the manufacturer’s protocol and eluted in 20 µL of water.

For in vitro DMS modification, embryos were collected and total RNA extracted as mentioned above, omitting DMS and β-mercaptoethanol. For each replicate, 20 µg of total RNA was resuspended in RNA folding buffer (57 mM Tris HCl pH 7.4 and 114 mM KCl) in a final volume of 44 µL. RNA samples were incubated at 65 °C for 2 min and slowly brought back to 28 °C over ~45 min in a heating block. Then, 5 µL of 100 mM MgCl2 was added to each tube and samples were incubated at 28 °C for 3 min. RNA was modified by adding 1 µL of 25% DMS, previously diluted in ethanol (1:3 dilution), to reach a final DMS concentration of 0.5%. The reaction was incubated at 28 °C for 10 min in a PCR machine and quenched by the addition of 59 µL of stop solution (3 M β-mercaptoethanol, 508 mM sodium acetate and 15 µg glycolblue). After a 5 min incubation at room temperature, modified RNA samples were ethanol precipitated by the addition of 220 µL ethanol. Poly(A)+ mRNAs were isolated as mentioned above.

All DMS-seq experiments (in vivo, in vitro, untreated, PatA-treated, and ChX-treated at 2 hpf, as well as 4 and 6 hpf) were performed in biological duplicates from embryos originated from different crosses and collected on different days.

Translation inhibitor treatments. PatA reduces the levels of functional eIF4F initiation complex, which is essential for cap-dependent translation51, by trapping elf4A onto mRNAs and ectopically enhancing its RNA helicase activity36,37 (Fig. 3a). Cycloheximide (CHX) binds to the E-site of the 60S ribosome subunit, inhibiting ribosomal translocation during translation elongation38. For DMS-seq and ribosome profiling experiments, 16-cell stage embryos were bathed in either 10 µM pateamine A (PatA) or 50 µg/mL cycloheximide (CHX). Embryos were collected once untreated embryos from the same clutches reached 64-cell stage (2 hpf) (~45 min following the addition of inhibitors). For DMS-seq samples, 150 embryos were collected in a 5 mL tube for each condition (untreated, PatA and CHX) and DMS- treated. Translation inhibitor concentrations were maintained during the DMS modification step. For ribosome profiling samples, 55 embryos per-condition and per-replicate were collected and flash frozen in liquid nitrogen.

DMS-seq library preparation. DMS-seq experiments were performed as previously described by Rouskin et al13, with minor changes. Briefly, DMS treated poly(A)+ RNA samples were denatured at 95 °C for 2 min and fragmented at 95 °C for 1.5 min in 1X RNA fragmentation buffer (Zn2+ based from Ambion). The reaction was stopped by adding 0.1 volume of a 10X Stop solution (Ambion). One volume of formamide loading dye (95% formamide, 10 mM EDTA, 0.025% bromophenol blue and 0.025% xylene cyanol) was added to fragmented RNAs, and samples were separated in a 10% TBU (Tris borate 8 M urea) PAGE. Fragments were visualized by blue light (Clare Chemical Research), and RNAs of 60-70 nucleotides were excised. Gel pieces were shredded and gel extracted in 300 µL of nuclease free water (AmericanBio) at 70 °C for 10 min with vigorous shaking. Eluted RNA was purified using Spin-X tube filters (Sigma-Aldrich) and ethanol precipitated by adding 33 µL 3 M sodium acetate, 1.5 µL glycolblue and 700 µL ethanol. RNA samples were dissolved in 6 µL of nuclease-free water, and 3’ phosphates were removed by incubating the samples at 37 °C for 1 h with 1 µL 10X PNK buffer (NEB), 1 µL SUPERase Inhibitor (Ambion) and 2 µL of T4 PNK enzyme (NEB). Samples were then directly ligated to 1 µg of our in-house barcoded 3’ adapters, either /5rApp/ NNcacaACTGTAAGGCACCATCAAT/3ddC/ or /5rApp/ NNagagACTGTAAGGCACCATCAAT/3ddC/ (IDT DNA) —in-house barcode depicted in lower case— by adding 1 µL 0.1 M DTT, 6 µL 50% PEG, 1 µL 10X ligase2 buffer, 2 µL T4 RNA ligase2, truncated K227Q (NEB), and incubated at 25 °C for 1.5 h. Each replicate was ligated to a different barcode set, and therefore, replicates could be pooled following ligation. 20 µL of formamide dye was added to each tube, and ligated products were run in a 10% TBU
PAGE for ~45 min, visualized by blue light and separated from unligated adapters by gel extraction as described above. Reverse transcription was performed in 20 µL at 52 °C for 45 min using Transcriptor High Fidelity cDNA Synthesis Kit (Roche). The samples were then extracted using the RNeasy Mini Kit (Qiagen) and purified using the QIAcube robot (Qiagen), followed by ethanol precipitation. The RNA samples were then treated with DNase I (NEB) and purified by gel electrophoresis.

For RNA-seq experiments, 75 ng of *Saccharomyces cerevisiae* RNA was added as spike-in to 175 µL of the remaining lysate supernatant, followed by extraction of total RNA using TRIzol. Total RNA samples were then sent to the Yale Center for Genome Analysis and strand-specific TruSeq Illumina RNA sequencing libraries were constructed. Prior to sequencing, samples were treated with Epicentre Ribo-Zero Gold according to the manufacturer’s protocol.

**RNA-seq time course experiments.** To quantify changes in mRNA abundance and to measure the impact of zygotic factors and miR-430 activity, we performed an RNA-seq time course experiment in wild type conditions, in the presence of alpha-amanitin (to inhibit zygotic transcription activation), and in the presence of an miR-430 inhibitor. To inhibit transcription activation of the zygotic genome, embryos were injected with 2 ng of alpha-amanitin (aAm) (Sigma Aldrich), which is an inhibitor of RNA polymerase II. To inhibit miR-430 activity, embryos were injected with 1 nanoliter of 10^8 M of tiny locked nucleic acid (tinyLNA), complementary to the seed region of miR-430 (5'TAGCAGTCT-3' (Exiqon)). ~25 embryos were collected per-stage and per-condition. Embryos were lysed in TRIzol and spiked with 75 ng of *Saccharomyces cerevisiae* RNA for normalization purposes. Total RNA was extracted, sent to the Yale Center for Genome Analysis and strand-specific TruSeq Illumina RNA sequencing libraries were constructed for each sample. Prior to sequencing, samples were treated with either Epicentre Ribo-Zero Gold, to deplete ribosomal RNA, or oligo dT beads, to enrich in poly(A)+ RNA, according to manufacturer’s protocol.

**Read sequencing and mapping.** DMS-seq, ribosome profiling and RNA-seq samples were sequenced on Illumina HiSeq 2000/2500 machines producing single-end 76 nucleotide reads. This yielded over 1 billion reads, 200 million reads and 100 million reads for DMS-seq, ribosome profiling, and RNA-seq respectively. Sequencing samples are summarized in Extended Data Table 1.

Following the library preparation protocol for DMS-seq and ribosome profiling samples, raw reads contained the following features: NNNN-insert-NN-adapter where the 6N (NNNN+NN) sequence composes the Unique Molecular Identifier (UMI), “barcode” is the sample 4-mer in-house barcode and adapter is the 3'-Illumina adapter. The UMI was used to discard PCR duplicates and count single ligation events. The barcode was used to mark individual replicate following the 3'-adapter ligation step. Base calling was performed using CASAVA-1.8.2. The Illumina TruSeq index adapter sequence was then trimmed by aligning its sequence, requiring a 100% match of the first five base pairs and a minimum global alignment score of 60 (Matches: 5,
Translating reads were demultiplexed based on the sample’s in-house barcode, the UMI was clipped from the 5′- and 3′-ends and kept within the read name, for marking PCR duplicates. Reads were then depleted of rRNA, tRNA, snRNA, snoRNA and mscRNA, using Ensembl87 annotations, as well as from RepeatMasker annotations, using strand-specific alignment with Bowtie2 v2.2.45. The remaining reads were aligned to the zebrafish Zv9 genome assembly using STAR version 2.4.2a56 with the following non-default parameters: --alignEndsType EndToEnd --outFilterMultimapNmax 100 --seedSearchStartLmax 15 --sbfdscore 10 --outSAMattributes All. Genomic sequence indices for STAR were built including exon-junction coordinates from Ensembl 78. Only reads of unique UMI were kept at each genomic coordinate for DMS-seq and ribosome profiling experiments. For ribosome profiling samples, reads were also aligned to the 256 sequences composing the 4 partially degenerated 28 nucleotides RNA spike-ins using Bowtie2 v2.2.4.

Raw reads from RNA-seq experiments were processed using the same pipeline, omitting the adapter trimming, barcoding demultiplexing and UMI clipping steps. The filtered reads were aligned onto Zebrafish Zv9 and Saccharomyces cerevisiae r64d1d1 genome assemblies using STAR, with the same parameters as described above. For both species, STAR genomic sequence indices were built including exon-junction coordinates from Ensembl 78.

**DMS-seq profiles and accessibilities.** Per-transcript profiles were computed using uniquely mapped reads overlapping at least 10 nucleotides with the transcripts’ annotation. Each read count was attributed to the nucleotide in position –1 of the read’s 5′-end within the transcript coordinate, to correct for the fact that reverse transcription stops one nucleotide prior to the DMS-modified nucleotide. To determine read distributions for each nucleotide (Extended Data Fig. 1c), only transcripts with a minimum of 100 counts were considered. In the same figure, ‘Transcriptome’ represents the frequency of each nucleotide for the same subset of transcripts (Extended Data Fig. 1c).

Accessibilities were calculated following the 2%/8% rule58, i.e., by normalizing the read counts proportionally to the most reactive A and C bases within the region after the removal of outliers. More specifically, the 2% most reactive A and C bases were discarded and each position was divided by the average of the next 8% most reactive A and C bases. Accessibilities greater than 1 were set to 1, and accessibilities for G and T were set to 0.

**Calculating translation efficiency.** Translation efficiency was calculated by dividing the Ribo-seq RPKM by the RNA-seq RPKM (RPKM; Read Per Kb and per Million reads of spike-in) for each coding sequence, excluding the first and last three codons (effective CDS) as described in Bazzini et al.52. Using the effective CDS of each transcript allows computation of translation efficiency from actively translating ribosomes.

For ribosome profiling samples, the effective CDS annotation was shifted 12 nucleotides upstream to position each read at the ribosome P-site location. RPKMs were computed by summing the effective CDS counts, including reads matching up to five times in the genome (each mapping site counting 1/n, n = number of mapping sites), and normalizing with the effective CDS length and the total number of reads mapped to the spike-in. For RNA-seq, RPKMs were calculated using the same pipeline to obtain ribosome profiling RPKMs, with the following differences: the effective CDS annotation remained unshifted, counts from reads overlapping the effective CDS by a minimum of 10 nucleotides were added, and the total number of reads mapped to yeast RNAs was used as normalization spike-in. Finally, translation efficiency was computed by dividing ribosome profiling RPKMs by RNA-seq RPKMs for each transcript effective CDS. For each sample, this analysis was performed by combining reads from both replicates.

**Identification of conserved RNA structures.** Conserved RNA structures (Extended Data Fig. 1g, h) were identified using the RNAz software57 on the UCSC Dan. rerio multiz 8-way whole genome alignment (http://hgdownload.cse.ucsc.edu/goldenPath/ danRer7/multiz8way), with default parameters.

**mRNA accessibility analysis.** To investigate global mRNA structure, the average accessibility of 5′-UTR, CDS and 3′-UTR was individually calculated and compared across conditions and/or between translation statuses. For this aim, we first identified the major transcript isoform of each gene, by selecting the one with the highest DMS-seq read density and with UTRs longer than 75 nucleotides (total of 11,404 mRNAs). If no isoform met the minimum UTR length criteria, no isoform was selected for that gene. Transcripts with average DMS-seq count per A/C equal to or greater than 5 (except when specified differently), and with counts covering at least 85% of A/C in each region (UTRs and CDS) were selected for further analysis. When the analysis was performed on full-length UTRs (Fig. 4c and Extended Data Fig. 1f, 2b, 2c, 3a, 5a and 6a), DMS-seq data was used to determine the final UTR lengths. Specifically, UTR length was extended one nucleotide at a time, starting at 76, until the A/C counts coverage fell under 85%. The longest 5′- and 3′-UTRs with at least 85% coverage was then associated to this transcript. Accessibilities were calculated within each region (5′-UTR, CDS and 3′-UTR) of each transcript as described above and averaged (for A/C bases only) to obtain a single value per region. For all global accessibility mRNA region analyses (cumulative plots), statistical significance between highly (red) and lowly (blue) translated mRNAs within the same condition was calculated using a one-sided Mann-Whitney U test while statistical significance between the same subset of mRNAs, but across two different conditions, was...
calculated using a two-sided Wilcoxon signed-rank test.

Definition and selection of differentially translated mRNAs. To select lowly and highly translated mRNAs, transcripts were binned in quintiles based on translation efficiency. The quintile with the lowest translation efficiency was labeled as 'low translation' and the one with the highest translation efficiency was labeled as 'high translation'.

Sliding windows analysis. To decipher which regions of the transcript were significantly changing in RNA structure across conditions and developmental stages, per-transcript DMS-seq counts were subdivided into overlapping 100-nt sliding windows, offset by one nucleotide. Only windows with minimum coverage of 250 counts were considered for further analysis. Windows with statistically significant changes in RNA structure across conditions were identified using the Kolmogorov-Smirnov test, a non-parametric test sensitive to differences both in location and shape of the empirical cumulative distribution functions of two samples. The Gini index, which has been previously used to compare RNA structures due to its ability to capture inequalities in a given distribution, was used to identify windows with increased or decreased RNA structures (Fig. 1c). All analyses were performed separately for each replicate, and P-values and Gini indices were combined using Fisher's method (R package 'metap') and averaged, respectively. Differential structured windows between two conditions were defined as those that were significant (combined P-value<0.05 in the KS-test, and that had an average Gini fold change across replicates greater than 1.1 or smaller than 0.9 (Fig. 1c). Metaplots of the distribution of differentially structured windows along the transcript were normalized by the region lengths and by the total number of windows analyzed in each region (5'-UTR, CDS, 3'-UTR).

Sequence conservation analysis. To determine the sequence conservation, we employed PhastCons regional conservation scores derived from the Multiz alignment of 8 vertebrate species, which included 5 fish species (ftp://hgdownload.cse.ucsc.edu/goldenPath/danRer7/phastCons8way/).

uORF analysis. A single transcript isoform per gene was selected as described above except for the minimal UTR length, which was set to 12 nucleotides. We then used the zebrafish uORFs dataset from Johnstone et al., which contains location and translation efficiency information, to identify lowly and highly translated uORFs at 2 hpf as described above for mRNAs. Per-uORF DMS-seq accessibility profiles were calculated by applying the 2%/8% rules mentioned above, for uORFs longer than 150 nucleotides. For shorter uORFs, DMS accessibilities were computed using 150 nucleotide regions starting at the uORF AUG initiation codon. Global uORF accessibilities were derived as described above. uORFs containing less than 10 A/C bases were removed from this analysis.

Poly(A) tail length and miR-430-mediated repression analyses. Poly(A) tail length datasets were obtained from Subtelny et al., who developed a technique, called poly(A)-tail length profiling by sequencing (PAL-seq), to measure the poly(A) tail length of transcripts from various organisms, including 2, 4 and 6 hpf zebrafish embryos. Only transcripts with at least 50 poly(A) tags were considered in our poly(A) tail length analyses (Fig. 4b, c and Extended Data Fig. 5). To analyze the impact of miR-430-mediated repression on mRNA structures (Fig. 4d, e), a set of 483 mRNAs targeted by miR-430 was selected using the following criteria: i) minimum 2-fold decrease in mRNA level between 2 and 6 hpf (6/2); ii) minimum 1.5-fold increase in mRNA level between 6 hpf tlnLNA and 6 hpf wild type (6tlnLNA/6wt); and iii) minimum of 5 reads on average per A/C base in DMS-seq experiments. For comparison, a random set of 483 mRNAs was selected among all the remaining transcripts with enough DMS-seq coverage. For genes with multiple transcript isoforms, the one with UTRs greater than 50-nt in length and with the highest coverage in our DMS-seq experiment at 2 hpf was selected. If no transcript isoform met those criteria for a specific gene, no transcript from that gene was included in the analysis.

RNA secondary structure analyses. Predicted RNA secondary structures were computed using either the Fold executable of the RNA structure package (version 5.6) or the SeqFold software. For the Fold software, the default parameters were used except for the temperature parameter, which was set to 301.15 Kelvin (28 °C). DMS-seq accessibilities of A and C bases were inferred as soft constraints using the −dms option. The Gibbs energy of folding (ΔG) of each predicted RNA structure was calculated using the efm2 executable with default parameters except for the temperature parameter, which was set to 301.15 Kelvin (28 °C). Arc plots representing RNA secondary structure were drawn using the R-chie software with default parameters. The analysis of the RNA structure surrounding AUG initiation codons was performed using 100-nt windows centered on each AUG with a minimum DMS-seq depth of 5 reads in average per A and C bases (Fig. 2e and Extended Data Fig. 2d-g). The analysis of the impact of ribosomes on local RNA structure was limited to 100-nt windows located in CDS regions and with a minimum DMS-seq depth of 5 reads in average per A and C bases (Fig. 3f and Extended Data Fig. 3b, c). Windows with ribosome footprints correspond to those with an average of ribo-seq reads between replicates greater than 7. Windows without ribosome footprints correspond to those with an average of ribo-seq reads lower than 1 and found in mRNAs with translation efficiency lower than 1.65. Each subset contains a total of 728 non-overlapping windows covering 332 and 312 different genes for windows with and without ribosome footprints, respectively (Extended Data Fig. 3c).

For the SeqFold analysis, only transcripts with more than 7 reads in average per A/C bases in all four tested conditions (untreated, in vitro, PatA-
treated or CHX-treated) were considered, which resulted in a set of 1,143 transcripts. DMS-seq accessibility of A and C bases was inferred following the same protocol as for SHAPE accessibilities and energies of folding were directly retrieved from the SeqFold output. To determine the global similarity of predicted RNA structures across conditions, transcript Gini indices were computed from the SeqFold outputs using the reldist library in R. Principal Component Analysis (PCA) was then used to reduce the dimensionality of the dataset. A biplot of the PCA loadings (i.e., conditions/treatments) of the two first principal components was used to determine which conditions were most similar. The same transcriptome-wide PCA analysis was done separately for CDS and 3′-UTR regions. The cumulative proportion of variance explained by the two first components was: 92% (PC1) and 3.1% (PC2) in CDS regions, and 94.5% (PC1) and 2.3% (PC2) in 3′-UTR regions.

**miR-430 site structural analysis.** To assess the impact of RNA structures on microRNA activity, we selected 18 different endogenous miR-430 binding sites with DMS-seq coverage equal to or greater than 5 reads per A/C bases and obtained their decay activity from Yartseva et al.\(^45\). The stability (minimum free energy, \(\Delta G\)) of the structure formed by the 120-nt region centered on the miR-430 seed was calculated in different conditions (\textit{in silico, in vitro} and \textit{in vivo} at 2 hpf, prior to miR-430 expression), and the correlation between the RNA structure stability and decay strength was determined using Spearman correlations (Fig. 5d). The predicted secondary structure and stability of a 200-nt region centered on the miR-430 seed were computed for additional endogenous miR-430 sites found in maternal mRNA 3′-UTRs (Fig. 5e, i and Extended Data Fig. 8). For each maternal mRNA containing miR-430 sites, decay patterns were analyzed in the wild type condition (with miR-430, WT) or when miR-430 activity was inhibited (-miR-430) by using a tiny LNA antisense to the miR-430 seed region (Fig. 5g, h, k, l). In vivo, the impact of RNA structures on miR-430-mediated repression was quantified at 24 hpf using ImageJ. MiR-430 activity was calculated as the ratio GFP/DsRed for both genetic backgrounds (wild type and MZmir430). Wild type GFP/DsRed was then normalized by the GFP/DsRed ratio from MZmir430, obtaining miR-430-mediated repression values for each reporter. Reported values come from three biological replicates (Fig. 5h, l).

Finally, to determine the impact of Ago2/miR-430 complex binding on the RNA accessibility of the recognition site, we analyzed the per-nucleotide accessibilities of 100-nt windows centered on individual miR-430 seeds (8- and 7-mers) found in the 3′-UTR of miR-430 targets (identified as described above), for each developmental stage and condition (Extended Data Fig. 7e-g). For comparison, per-nucleotide accessibilities of randomly chosen 100-nt windows within the same set of 3′-UTRs were used. Global seed and control accessibilities were calculated by averaging the accessibility of each A and C bases found in the 8-nt sequence located in the center of each window (Extended Data Fig. 7f). Only windows with a minimum of 5 reads in average per A and C bases were considered.

**RNA-element selection assay.** To quantify the regulatory activity of maternal mRNA 3′-UTR sequences during the MZT, we used an RNA-element selection assay (RESA) as described in Yartseva et al.\(^45\) with minor changes (Fig. 6b). Maternal 3′-UTR regions of 200 nucleotides showing changes in RNA structure during the MZT (based on KS-test across stages, \(P<0.05\)) or no change (as controls, \(P>0.05\)) were amplified from cDNA of 2 hpf embryos (see Extended Data Table 2 for primer sequences). In total, 74 dynamic and 25 control regions were amplified and inserted in the 3′-UTR of a GFP reporter by assembly PCR with Phusion High Fidelity DNA polymerase (NEB, cat. No. M0530), and purified on agarose gel. The final library constructs consisted of an Sp6-promoter, a GFP coding sequence, a 3′-UTR with various inserts flanked by part of the Illumina 5′- and 3′-adapter sequences, and a SV40 polyadenylation signal. The library was \textit{in vitro} transcribed with an Sp6 mMessage mMachine kit (Thermo Fisher, cat. No. AM1340) and the resulting RNA library was injected in 1-cell stage embryos, with or without the presence of alpha-amanitin. Alpha-amanitin (aAm) is an inhibitor of RNA polymerase II that blocks the activation of the zygotic genome. Approximately 25 injected embryos were collected at 2 and 6 hpf (with and without aAm), and total RNA was extracted with TRIzol reagent according to the manufacturer’s protocol. Poly(A)+ mRNA was selected with NEB oligo d(T)\(_{25}\) magnetic beads (NEB, cat. no. S1419S) following the manufacturer’s protocol. Poly(A)+ mRNA was recovered with ethanol precipitation and dissolved in 11 µL of water. Concentrated poly(A)+ mRNA was reverse-transcribed with a reporter specific primer (5′-CATCAATGTATCTTTATCATGTCTGGATC-3′) with SuperScript III. Illumina adapters were added and the library amplified with ~20 circles of Phusion PCR with the forward primer (5′-
To identify the regions bound by KHSRP during early embryogenesis, we performed iCLIP experiments as described in Huppertz et al. with minor changes. Embryos at sphere stage (4 hpf) were collected and irradiated with 254 nm UV light to induce crosslinking (embryos were snap frozen and stored in batches to isolate RNA-protein complexes. Briefly, 200 μl Protein G Dynabeads were added to 50 μg antibody in lysis buffer. Beads were incubated for an hour, washed three times with lysis buffer and added to the lysates. As a control, a parallel experiment was performed without the addition of antibody. Subsequent steps in the iCLIP protocol were performed as described in Huppertz et al., except for the 3'-adaptors containing in-house barcode, the reverse transcription primer and PCR primers, which were identical to the ones used for the DMS-seq library preparation. Twenty cycles of amplification of cDNA were used for final library construction. Barcoded PCR-amplified libraries were size-selected in a 6% TBE gel and combined for Illumina sequencing. Both the KHSRP pull-down and the no-antibody control were performed in triplicates. Raw reads from iCLIP experiments were processed using the same pipeline as for the DMS-seq experiments.

To define the binding preference of KHSRP, bound windows were defined for each 6-mer with at least 10 reads and extended 5 nucleotides upstream and 10 nucleotides downstream. For each bound window, the frequency of each 6-mer was computed. These frequencies were normalized by the average 6-mer frequencies observed in the no-antibody controls, which were calculated as mentioned above except for the read depth requirement, which was set to 2 instead of 10 reads to compensate for differences in sequencing depth. To build the logo representation, the top 10 most frequent normalized 6-mers was aligned using MAFFT with the following parameters: --reorder --lop -10 --lexp -10 --localpair --genafpair --maxiterate 1000. Multiple sequence alignment positions with more than 50% gaps were trimmed on both ends. Nucleotide frequencies were calculated using the final trimmed alignment, and represented as logo (Extended Data Fig. 7a).

iCLIP peaks were called by testing scanning 30-nt windows using Poisson’s law. Overlapping significant windows (P<0.01) were then merged to obtain the peaks. To get robust Poisson parameter estimates, only transcripts with a minimum of 20 positions with at least 1 read were analyzed. Final KHSRP peaks correspond to overlaps of peaks found in at least 2 out of 3 replicates. To analyze the RNA structure of KHSRP-bound and -unbound regions, we defined bound regions as 100-nt windows centered on the middle point of each KHSRP peak. For each bound region, an unbound control within the same 3' UTR and overlapping with less than 25% with the bound region was selected. The accessibility of each region was computed by averaging the accessibility of A and C bases. Finally, only transcripts and regions with a minimum of 5 reads in average per A and C bases were considered (Extended Data Fig. 7c).