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Structural imprints *in vivo* decode RNA regulatory mechanisms

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Abstract

Visualizing the physical basis for molecular behavior inside living cells is a grand challenge in biology. RNAs are central to biological regulation, and RNA's ability to adopt specific structures intimately controls every step of the gene expression program¹. However, our understanding of physiological RNA structures is limited; current *in vivo* RNA structure profiles view only two of four nucleotides that make up RNA^{2,3}. Here we present a novel biochemical approach, *In Vivo* Click SHAPE (icSHAPE), that enables the first global view of RNA secondary structures of all four bases in living cells. icSHAPE of mouse embryonic stem cell transcriptome versus purified RNA folded *in vitro* shows that the structural dynamics of RNA in the cellular environment distinguishes different classes of RNAs and regulatory elements. Structural signatures at translational start sites and ribosome pause sites are conserved from *in vitro*, suggesting that these RNA elements are programmed by sequence. In contrast, focal structural rearrangements *in vivo* reveal precise interfaces of RNA with RNA binding proteins or RNA modification sites that are consistent with atomic-resolution structural data. Such dynamic structural footprints enable accurate prediction of RNA-protein interactions and N⁶-methyladenosine (m⁶A) modification genome-wide. These results open the door for structural genomics of RNA in living cells and reveal key physiological structures controlling gene expression.

Selective 2'-Hydroxyl Acylation followed by Primer Extension (SHAPE) accurately identifies flexible (single stranded) bases in RNA for all four bases. However, current methods are potentially limited by high background (>70% of RNA molecules have no modification due to single hit kinetics) and high false positive rates due to spurious reverse transcription (RT) stops⁴. We overcome these problems by developing a new SHAPE probe that permits *in vivo* SHAPE modification and subsequent selective purification of the modified RNAs.

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Contributions

R.C.S., E.T.K., and H.Y.C. conceived of the study. R.C.S., P.C., J-W.J., H.K., E.T.K. performed chemical design and synthesis. R.C.S. R.A.F., B.L., E.A.T., P.J.B. performed biological experiments. Q.C.Z., R.C.S., R.A.F., and H.Y.C. performed data analysis. R.C.S. and H.Y.C. wrote the paper with input from all authors.

We designed, synthesized, and tested a novel bifunctional chemical probe for *in vivo* RNA structure profiling genome-wide (NAI-N₃, **Fig. 1a, 1b, and Extended Data Fig 1**). NAI-N₃ adds an azide group to NAI, a cell permeable SHAPE reagent⁵. By using copper-free click chemistry, a biotin moiety is selectively and efficiently added to NAI-N₃-modified RNA, providing a stringent purification handle with streptavidin beads (**Fig. 1c, Extended Data Fig. 2**). NAI-N₃ generated identical profiles of RT stops to those obtained using our previously designed SHAPE reagent⁵. The fidelity of structural measurements was not affected by “clicking” biotin onto the NAI-N₃ nor by molecular crowding of proteins, and NAI-N₃ showed uniform modification of all bases in denatured RNAs (**Extended Data Fig. 3**). We term this new chemoaffinity structure probing methodology *In Vivo* Click SHAPE (icSHAPE); this method can also be applied to any *ex vivo* preparation of RNA with slight modifications.

icSHAPE of ribosomal RNAs in mouse embryonic stem cells (mESC) indicated that it is quantitative and accurate, reporting the known structures of 18S and 28S rRNAs (**Fig. 1d-f and Extended Data Fig. 4**). Manual structure-probing gels and deep sequencing results from icSHAPE revealed high correlations (Pearson correlation $r=0.93$, *in vivo*; **Fig. 1d and Extended Data Fig. 4**). Ribosomal RNA is known to require the cellular environment for proper folding, and differences between *in vivo* and *in vitro* icSHAPE measurements highlighted important structural elements in the intact ribosome. We mapped nucleotides of icSHAPE difference to the cryo-EM structure of the human 80S ribosome⁶ and searched for differences between the *in vivo* and *in vitro* conditions. Conserved (mouse to human) nucleotides of high icSHAPE signal *in vivo* were unpaired in the cryo-EM structure (**Fig. 1e**); conversely residues lacking icSHAPE reactivity *in vivo* were base-paired or engaged in extensive interactions that may stabilize the RNA backbone in the mature ribosome (**Fig. 1f**). Overall, these data demonstrate that icSHAPE accurately measures RNA structures, both in and outside of living cells.

We next used icSHAPE to measure RNA structural profiles of polyadenylated transcripts in mESCs and generated ~2.1 billion measurements for over 13,200 RNAs *in vivo* and *in vitro*, with high reproducibility (**Extended Data Fig. 5 and 6**). The nucleotide composition in the transcriptome, mock-treated RNA, and icSHAPE RNA are highly concordant, with a slight enrichment in NAI-N₃ samples for A's and U's (**Fig. 2a**). This enrichment is expected given their bias for being located in single stranded or loop regions⁷. icSHAPE thus affords the first complete RNA structurome of all four nucleotides *in vivo*.

icSHAPE data revealed the scale and distribution of RNA structural dynamics between *in vitro*, where folding is programmed entirely by sequence, versus *in vivo* where folding occurs in the context of the intracellular environment.⁸ Recent transcriptome-wide dimethylsulfate probing (DMS-seq), which interrogates two bases with strong bias toward adenosines (68% A's and 24% C's)^{2,4}, suggests that RNA structures are largely unfolded *in vivo*²; however sampling only two of four nucleotides could result in an incomplete picture. We quantified RNA structural dynamics using two metrics: First, we calculated the difference in reactivity between our *in vivo* and *in vitro* icSHAPE measurements, termed the Vivo – Vitro Difference (VTD, **Fig. 2b and Methods**). Adenosine residues have the largest

VTD, whereas guanosine and cytidine residues are less variable between environments (**Fig. 2c**). These observations suggest that utilizing probes that have a broader reactivity profile such as NAI-N₃ will give a more complete representation of RNA structure.

Second, we used the GINI index² to quantify the distribution of icSHAPE reactivity profiles. Structured RNAs have some bases that are reactive and some not, leading to unequal distribution and high GINI, while unfolded RNAs have most bases in a uniformly reactive conformation (low GINI). We found that RNAs are less folded *in vivo*, consistent with a prior report², but the extent of unfolding varies in degrees that distinguish different classes of RNAs (**Fig. 2d**). Protein-coding mRNAs exhibited noticeable but partial unfolding (average GINI of 0.7 *in vitro* to 0.5 *in vivo*), with the largest variation noted at 3' untranslated regions (UTR) compared to coding sequences (CDS) or 5'UTRs. In contrast, noncoding RNAs, such as pseudogenes, long noncoding RNAs, and primary microRNA precursors, retain substantially more of their RNA structure *in vivo* ($p < 2.2 \times 10^{-16}$, noncoding vs. coding, Student's *t*-test). One exception to this rule are snoRNAs, which exhibit the greatest level of increased reactivity *in vivo* among all classes of transcripts and may result from extensive rearrangements due to snoRNP binding. Most RNAs *in vivo* possess a substantial level of RNA structure beyond previous expectation based on DMS-seq². The current data suggest that RNA structural signatures *in vivo* can distinguish coding vs. structural or regulatory RNAs, consistent with prior *in vitro* studies⁹⁻¹².

The dramatically different environments that RNA experiences when inside a cell compared to *in vitro* predicts that our VTD parameter could provide insight into functionally important RNA regulatory elements. To assess this possibility, we measured the VTD for all hexamer sequences (**Fig. 2e, Supplementary Table 1**). We observed unique VTD profiles for sequence motifs driving diverse post-transcriptional processes, including translation initiation, interaction with RNA binding proteins (RBPs, e.g. Rbfox2), RNA modification (m⁶A), and microRNA seed matches^{13,14,15} (**Supplementary Table 1 and 2**). These results show that the VTD may classify RNA regulatory elements as pre-programmed or sensitive to *in vivo* remodeling. Further, distinctive VTD profiles precisely at sites of post-transcriptional regulatory motifs suggest that RNA structural dynamics may be used to monitor these regulatory events in cells.

We hypothesized that translational regulatory elements may have their icSHAPE profiles conserved between *in vivo* and *in vitro* because the Kozak sequence, important for translation initiation¹⁶, is among the most stable (low VTD) regions within mRNAs (**Fig. 2d**). RNA accessibility from -1 to -5 nucleotides upstream of the start codon plays a major role in regulating translational output^{10,17}. We used translation initiation¹⁸ and pause sites¹⁸, defined by ribosome profiling, to center our structural reactivity analysis across the transcriptome (**Fig. 3**). Canonical initiation AUG sites are indeed preceded by ~5 nts of increased accessibility, and this pattern is nearly identical to *in vitro* folded RNA (**Fig. 3a and 3b**). A similar pattern of conserved upstream accessibility also precedes noncanonical start sites at upstream open reading frames (uORF) and N-terminal truncations (**Fig. 3c**). Non-start site AUG codons are also associated with increased preceding reactivity, while noncanonical CUG start codons have a different profile, suggesting that RNA accessibility alone is not sufficient to dictate translational start sites (**Extended Data Fig. 7**). Ribosome

distinguish true binding sites from other sequence motif instances, collectively boosting prediction accuracy.

We also identified a critical connection between RNA structure and RNA modification, a newly appreciated and pervasive mode of post-transcriptional control¹³. The most prevalent modification in mRNAs, m⁶A, occurs at GGm⁶ACU motifs near stop codons, and acts in part to control RNA splicing and stability^{22,23}. It has been hypothesized that m⁶A methylation occurs at sites that contain un-paired motifs²⁴, but no direct structural evidence has been presented to support this model. Comparison of icSHAPE signals at m⁶A-modified vs. unmodified instances of the GGACU motif in mESCs²⁵ revealed a specific structural signature, with stronger icSHAPE reactivity (consistent with unpaired RNA) at positions both surrounding and including the modified A (**Fig. 4d, Extended Data Fig. 8**). m⁶A sites in different subdomains of mRNAs or in lncRNAs have nearly identical icSHAPE profiles (**Extended Data Fig. 10**). Evaluation of all predictive features using our SVM algorithm showed that motif conservation or motif position offers some predictive value (AUC=0.617 or 0.824, respectively) as previously reported²⁴, but use of icSHAPE data (AUC = 0.846) or all features together (AUC = 0.914) improved prediction rate (**Fig. 4e**). These results show that icSHAPE structure profiles can be used to accurately predict post-transcriptional modifications on a transcriptome-wide scale.

The strong RNA structural signature at m⁶A sites may arise from the ability m⁶A to destabilize RNA helices²⁶ (represented in **Fig. 4c**) or the structural selectivity of m⁶A modification machinery for unpaired bases. In the former scenario, removal of m⁶A should cause increased basepairing (loss of icSHAPE signal) whereas the latter scenario predicts little change to RNA structural profile. To distinguish between these hypotheses, we determined the icSHAPE profile of mESCs genetically ablated for *Mettl3*²⁵, a key m⁶A methyltransferase that is required for ESC differentiation. We observed that in *Mettl3* knockout cells, canonical motif sites that lost m⁶A modification also substantially lost icSHAPE signal transcriptome-wide (**Fig. 4d**), as exemplified by key m⁶A target sites in *Nanog* mRNA (**Fig. 4f**). These results suggest that m⁶A impacts RNA structure, favoring the transition from paired to unpaired RNA. The ability to couple genetic perturbation with comprehensive, base-resolution structural maps *in vivo* is a potentially powerful approach to dissect regulators of RNA structure.

Understanding how RNA structures contribute to biological regulation opens the door to understanding a physical dimension of the transcriptome. icSHAPE bridges a gap in RNA sequencing technologies that currently lack the ability to infer a mechanistic basis of biological function. The ability to view the structural dynamics of all four RNA bases in living cells is essential to uncover specific sequence motifs underlying different modes of post-transcriptional regulation²⁷, and has enabled accurate identification and *de novo* prediction of trans-acting factor binding and chemical modification at single nucleotide resolution. In the future, viewing the RNA structure when cells are exposed to different stimuli or genetic perturbations should revolutionize our understanding of gene regulation in biology and medicine.

Methods

Methyl 2-(azidomethyl)nicotinate

1.00 g of methyl 2-methylnicotinate was dissolved in 5 mL anhydrous dichloromethane. 2.30 g trichloroisocyanuric acid was added and the resulting suspension stirred overnight at room temperature. The reaction was diluted with dichloromethane and quenched by the addition of saturated sodium bicarbonate solution. The phases were separated and the organic phase was washed once with brine, dried over magnesium sulfate, filtered, and concentrated to afford a yellow oil. NMR data was consistent with literature reports.

The crude product of the above reaction (1.09 g) was dissolved in 12 mL anhydrous N,N-dimethylformamide and 0.77 g sodium azide was added. The reaction was stirred overnight at room temperature then quenched with saturated sodium bicarbonate solution. The aqueous layer was extracted with ethyl acetate, and the organic layer washed 3 times with water and 3 times with brine. The organic layer was dried over magnesium sulfate, filtered, and concentrated to afford 0.91 g (71%, 2 steps) of a yellow oil that solidified upon standing.

^1H NMR (400 MHz, CDCl_3): 3.94 (3H, s), 4.88 (2H, s), 7.37 (1H, m), 8.29 (1H, dd, $J = 8$ Hz, 1.6 Hz), 8.76 (1H, dd, $J = 4.6$ Hz, 1.6 Hz)

^{13}C NMR (100 MHz, CDCl_3): 52.8, 54.4, 122.9, 125.0, 139.1, 152.4, 156.7, 166.0

ESI-MS (Calc M-H = 191.06): 191.98

2-(azidomethyl)nicotinic acid

0.50 gram of methyl 2-(azidomethyl)nicotinate was stirred vigorously in 10 mL of 1:1 MeOH:10% aqueous NaOH. After 10 minutes TLC indicated complete consumption of starting material. 25 mL of water were added, the crude reaction mixture was washed once with ether (10 mL) then acidified to pH 4 with 10% aqueous HCl and extracted 5 times with 50 mL ethyl acetate. The organic layer was dried over magnesium sulfate, filtered, and concentrated to afford 0.46 g (99%) of a white solid that was pure by NMR.

^1H NMR (400 MHz, $\text{DMSO}-d_6$): 4.81 (2H, s), 7.50 (1H, m), 8.28 (1H, dd, $J = 7.8$ Hz, 1.6 Hz), 8.74 (1H, dd, $J = 5$ Hz, 1.6 Hz), 13.64 (1H, br. s).

^{13}C NMR (100 MHz, $\text{DMSO}-d_6$): 53.4, 123.4, 125.9, 139.0, 151.9, 156.0, 167.1

ESI-MS (Calc M-H = 177.04): 177.05

2-(azidomethyl)nicotinic acid acyl imidazole

0.15 g 2-(azidomethyl)nicotinic acid was dissolved in 0.21 mL anhydrous dimethylsulfoxide. A solution of 0.14 g carbonyldiimidazole in 0.21 mL anhydrous dimethylsulfoxide was added dropwise, creating rapid gas evolution. The reaction was allowed to proceed for one hour and the resulting solution used as a 2M stock solution for RNA SHAPE experiments. For NMR data collection an analytical sample was prepared in dichloromethane as described above. The reaction was stirred over night and the solvent

removed in vacuum. The product was then isolated by flash column chromatography on silica (ethyl acetate).

^1H NMR (400 MHz, DMSO- d_6): 4.62 (2H, s), 7.16 (1H, dd, $J = 1.7$ Hz, 0.8 Hz), 7.60 (1H, dd, $J = 7.9$ Hz, 4.9 Hz), 7.66 (1H, m), 8.15 (2H, m), 8.84 (1H, dd, $J = 4.9$ Hz, 1.7 Hz).

^{13}C NMR (100 MHz, DMSO- d_6): 52.8, 117.9, 123.0, 127.3, 130.9, 137.7, 138.5, 151.9, 154.6, 164.7

***In Vitro* transcription and acylation of RNA**

RNA was transcribed from amplified inserts using T7 Megascript kit from Ambion, following manufacturer's protocol. In a typical *in vitro* modification protocol, RNA was heated in metal-free water for two minutes at 95°C. The RNA was then flash-cooled on ice. The RNA 3x SHAPE buffer (333 mM HEPES, pH 8.0, 20mM MgCl₂, 333mM NaCl) was added and the RNA was allowed to equilibrate at 37°C for ten minutes. To this mixture, 1μL of 10x electrophile stock in DMSO (+) or DMSO (–) was added. The reaction was permitted to continue until the desired time. Reactions were cleaned up using RNeasy columns (Qiagen) following manufactures protocol and eluted RNase-free water.

***In Vitro* manual SHAPE analysis**

^{32}P -end-labeled DNA primer (reverse primer above) was annealed to 3μg of total RNA by incubating at 95°C for two minutes followed by a step-down cooling (2°/sec) to 4°C. To the reaction first-strand buffer, DTT and dNTPs were added. The reaction was pre-incubated at 52°C for one minute, then Superscript III (2U/μL final concentration) was added. Extensions were performed for ten minutes. To the reaction, 1μL of 4M NaOH was added and allowed to react for 5 minutes at 95°C. 10μL of Gel Loading Buffer II (GLBII, Ambion, Inc.) was then added, and cDNA extensions were resolved on 8% denaturing (7M Urea) polyacrylamide gels (29:1 acrylamide:bisacrylamide, 1xTBE). All (–) lanes are those from DMSO control treated cells. In addition, all sequencing lanes are from DMSO control treated cells. cDNA extensions were visualized by phosphorimaging (STORM, Molecular Dynamics). cDNA bands were integrated with SAFA²⁸ SHAPE reactivities were normalized to a scale spanning 0 to 1.5, where 1.0 is defined as the mean intensity of highly reactive nucleotides.²⁹ RNA secondary structures were predicted using mFOLD software.³⁰

Characterization of manual SHAPE-enriched RT stops

Copper-free click chemistry of acylated RNA—In a typical reaction, acylated RNA (1pmol) was reacted with 100eq. of DIBO-biotin (Life Technologies) for two hours, at 37°C, in 1x Phosphate Buffer Saline (PBS). Reactions were extracted once with acid phenol:chloroform (pH 4.5±0.2) and twice with chloroform. RNA was precipitated with 40μL of 3M sodium acetate buffer (pH 5.2) and 1μL of glycogen (20μg/μL). Pellets were washed twice with 70% ethanol and resuspended in 10μL RNase-free water.

Enrichment of NAI-N₃ modified RNA—The following protocol was used for manual enrichment protocols used to optimize capture conditions. To 1pmol of precipitated and biotinylated RNA (in 900μL of binding buffer: 50mM Tris-HCl pH 7.0 and 1mM EDTA)

was added 50 μ L (slurry) of DYNAL MyOneC1 beads (Life Technologies) was added. The reaction mixture was then incubated for one hour at room temperature. The beads were then collected on a magnetic plate and the solution decanted. The beads were then resuspended and washed 4 times with Biotin Wash Buffer (10mM Tris-HCl, pH 7.0, 1mM EDTA, 4M NaCl, 0.2% Tween). The beads were then washed 3x with RNase-free water. To elute the purified RNA off the resin beads were incubated in 1X proteinase K buffer with 20U of proteinase K (Life Technologies), 1mM D-Biotin (Sigma Aldrich), and 20U of SUPERaseIn (Life Technologies). The reaction was permitted to run for 30 minutes at 37°C. Beads were then collected by magnet and the supernatant removed and set on ice. This was repeated twice more and elutions were pooled. Reactions were extracted once with acid phenol:chloroform (pH 4.5 \pm 0.2) and twice with chloroform. RNA was precipitated with 40 μ L of 3M sodium acetate buffer (pH 5.2) and 1 μ L of glycogen (20 μ g/ μ L). Pellets were washed twice with 70% ethanol and resuspended in 10 μ L RNase-free water.

Dot Blot analysis of enriched NAI-N₃ modified RNA—Hybond N+ membranes (GE) were pre-incubated in 1xPBS. Precipitated RNA was dissolved in 100 μ L of 1xPBS. RNA was added to the Hybond membrane and crosslinked using 254nm UV light. The Hybond membrane was washed three times with 1xPBS. To the membrane was added NorthernLights Streptavidin NL493 (in PBS-Tween20) for visualization. After incubation, the membrane was washed 3x in 1xPBS-Tween20. The membrane was dried and imaged by phosphorimaging (STORM, Molecular Dynamics).

Tissue culture and *In Vivo* SHAPE modification

Mouse embryonic stem cells (v6.5 line) were grown on gelatinized dishes in serum and LIF. Unmodified total RNA was extracted by removing media, washing once in room temperature 1xPBX, and adding 2mL (10cm dish) or 7mL (15cm dish) of Trizol directly to the cells. Subsequent RNA extract was performed using the miRNeasy mini- or midi-column and protocol (Qiagen) as recommended by the manufacture. *In Vivo* modification of cellular RNAs was performed as described previously.⁵ Briefly, cells were rinsed once on the plate in room temperature 1xPBS, decanted, scraped in 1xPBS, and collected into a 15mL tube. Cells were pelleted at room temperature and resuspended in 450 μ L of 1xPBS. 50 μ L of 10x electrophile stock in DMSO (+) or DMSO (–) was added drop-wise, immediately mixed by inversion, and incubated at 37°C on end-over-end rotation for 20 minutes. Reactions were pelleted for 1min at 4°C at 10,000 rpm and resuspended in 500 μ L of 1xPBS. Samples were then transferred to 15mL tubes with 2-7mL of pre-aliquoted Trizol and RNA extracted as described above.

Methods to ensure titrated hit-kinetics of RNA modification—We titrated NAI-N₃ for single-hit kinetics that are comparable to those routinely used in chemical probing of RNA structure. For example, we obtained nearly identical secondary structure for 5S rRNA as previously reported with single hit regime.⁵ After NAI-N₃ modification and biotin pulldown, we retrieved approximately 10-20% of the input RNA as modified RNA, consistent with the expected Poisson distribution of single-hit modification.

icSHAPE deep sequencing library preparation

RNA preparation—DMSO (mock) or NAI-N₃ (experimental) modified total RNA was used as input for the deep sequencing library preparation. Before library preparation input RNA should be modified (or mock treated) under *In Vitro* or *In Vivo* conditions as described above. For “total RNA” libraries, no additional processing was needed. For “poly-A selected” samples, 200µg of total RNA was used per poly-A purist column (Ambion, Inc.), which should yield ~2µg of enriched RNA. Poly-A selection was performed a total of two times using the same poly-dT beads (“double poly-A selection”). The NAI-N₃ sample may have lower yield post purification so additional starting material could be required.

NAI-N₃ biotinylation and RNA fragmentation—All RNA samples (NAI-N₃ and DMSO treated) are processed through a copper-free “click” reaction. RNA is brought to 97µL in 1xPBS and 1µL of SUPERaseIn and 2µL of 185mM DIBO-Biotin are added. Samples were mixed by brief vortexing and then incubated at 37°C for 2 hours in a Thermomixer (Eppendorf). Reactions were stopped by adding 350µL of Buffer RLT (Qiagen) and then 900µL of 100% ethanol (EtOH). Each RNA sample was processed by passing over a RNeasy Mini column (Qiagen), two 500µL washes with Buffer RPE (Qiagen), one no-buffer spin to dry the column, and finally two 50µL elutions in RNase-free water (final 100µL). Samples were then frozen for 5 minutes on dry ice and concentrated to 9µL using a lyophilizer (Labconco). Concentrated RNA samples (9µL) were then moved to 0.5mL PCR tubes for fragmentation. Samples were heated to 95°C for 90 seconds and then 1µL of 10x RNA Fragmentation Reagent (Ambion, Inc.) was added and samples were placed back at 95°C for 70-90 seconds. Reactions were quenched by adding 1µL of RNA Fragmentation Stop Solution (Ambion, Inc.) and moved to ice. RNA was cleaned up by adding 35µL of Buffer RLT and 100µL of 100% EtOH and purified using RNeasy Mini columns as above. Samples are then concentrated with a lyophilizer to 5µL.

RNA end repair, RNA ligation, and RNA size selection—To resolve the 3'-end phosphate generated by the fragmentation process T4 PNK is used. To each 5µL sample 2µL of 5x PNK buffer (350mM Tris-HCl pH 6.5, 50mM MgCl₂, 25mM DTT), 1µL SUPERaseIn, and 2µL of T4 PNK (NEB) is added, mixed by flicking, and incubated at 37°C for 1 hour. After end-repair samples are moved directly to 3'-end ligation by adding 1µL of 50µM 3' Adaptor, 1µL of 10x T4 RNL2tr buffer (NEB), 1.5µL of T4 RNL2tr K227Q (NEB), 1µL of 100mM DTT, and 8µL of 50% PEG8000. Mix samples by flicking and incubate at 16°C overnight.

Ligation Note: NAI-N₃ samples must use “3'-Adaptor-3'ddc” (/5rApp/AGATCGGA AGAGCGGTTTCAG/3ddC/) while DMSO samples must use “3'-Adaptor-3'Biotin” (/5rApp/AGATCGGAAGAGCGGTTTCAG/3Bio/). The “click” chemistry will label only the NAI-N₃ modified RNAs in the NAI-N₃ pool of transcripts with a biotin moiety, thus allowing the selective purification of structurally informative molecules. The DMSO samples are not capable of “click” chemistry and every molecule in this pool is desired for sequencing so addition of a biotin moiety must happen in an unbiased fashion. Thus, DMSO samples have a 3'Biotin modification added specifically to their 3' Adaptor to allow for downstream processing in parallel of the DMSO and NAI-N₃ samples.

After the overnight ligation 30 μ L of water, 185 μ L of Buffer RLT, and 400 μ L of 100% EtOH is added to each sample and purified using RNeasy Mini columns as above. Samples are concentrated to 5 μ L using a lyophilizer and 5 μ L of GLBII is added and stored on ice. To size select the RNA samples a mini 6% TBE PAGE gel with 7M Urea is cast and pre-run to 50W for 8 minutes. Samples are loaded without prior heating and the PAGE gel is imaged using a 1:10,000 dilution of SybrGold (Life Technologies). RNA is visualized on a BlueBox (Clare Chemical) and fragmented RNA ranging between 20-120nts (40-140nts with the 3' Adaptor ligated) are excised with a scalpel. Gel slices are crushed through a 0.75mL tube nested in a 2mL tube by centrifugation and 300 μ L of Crush Soak Buffer (500mM NaCl, 1mM EDTA) is added with 3 μ L of SUPERaseIn. RNA is eluted overnight at 4°C on rotation.

RT, streptavidin capture, cDNA elution, and cDNA size selection—RNA samples are purified away from residual PAGE using 0.45 μ m Spin-X columns (Corning) and the 300 μ L elutions are transferred to silicionized 1.5mL tubes (Fisher Scientific, used in all subsequent steps). RNA is precipitated by adding 30 μ L of 3M sodium acetate buffer (pH 5.2), 0.8 μ L of GlycoBlue (Ambion, Inc.) and 1mL of 100% EtOH. Samples are frozen for 1 hour on dry ice, spun at max speed (15,000 rpm) for 1 hour at 4°C, washed with 800 μ L of ice-cold 80% EtOH, decanted, air-dried and then resuspended in a 0.5mL PCR tube with 11.5 μ L of water. To the RNA samples add 1 μ L of 10 μ M RT primer (/5phos/ DDDNNAACCNNNN AGATCGGAAGAGCGTCGTGAT/iSp18/GGATCC/iSp18 / TACTGAACCGC, /5phos/ = 5' phosphate, D=A/T/G, /iSp18/ = 18carbon PEG spacer) and 1 μ L of 10mM dNTPs. Heat the samples to 70°C for 5 minutes and then cool slowly to 25°C (2deg/second) and hold at 25°C for one minute. After primer annealing add 0.5 μ L of SUPERaseIn, 1 μ L 100mM DTT, 4 μ L of 5x First Strand Buffer and 1 μ L of SuperScript III (Life Technologies). cDNA extension occurs for 3 minutes at 25°C, 7 minutes at 42°C, and finally at 52°C for 15 minutes. After cDNA extension do not raise samples above 37°C to avoid denaturing conditions.

MyOneC1 streptavidin beads for cDNA capture and NAI-N₃ modified RNA enrichment are prepared (40 μ L slurry per sample) by washing 3x in 1mL of Biotin Bind Buffer (100mM Tris-HCl pH 7.0, 10mM EDTA, 1M NaCl) and resuspending the beads in 40 μ L Biotin Bind Buffer and 1 μ L SUPERaseIn per reaction. After the reverse transcription reaction completes 40 μ L of pre-washed beads are added to each sample, mixed by flicking, and incubated at room temperature for 45 minutes. After streptavidin capture samples are washed at room temperature serially with 4 \times 100 μ L of Biotin Wash Buffer, 2 \times 100 μ L 1xPBS and finally moved to 1.5mL tubes. cDNA is eluted by adding 1 μ L RNaseA/T1 cocktail (Ambion, Inc) 1 μ L RNaseH (Enzymatics), 12.5 μ L 50mM D-Biotin, 5 μ L 10x Elution Buffer (500mM HEPES, 750mM NaCl, 30mM MgCl₂, 1.25% Sarkosyl, 0.25% Na-deoxycholate, 50mM DTT), 30.5 μ L water and incubating at 37°C for 30min in a Thermomixer at 800 rpm. Samples are mixed with 1 μ L 100% DMSO, heated to 95°C for 3 minutes, placed on a magnet, and the 50 μ L cDNA elution moved to a new tube. The elution is repeated once (total of two times and final of 100 μ L). cDNA is processed by adding 1mL of Buffer PNI and purifying over a MiniElute columns (Qiagen) following manufactures protocol and eluting twice in 15 μ L of Buffer EB (final 30 μ L). cDNAs are concentrated using a lyophilizer

to 5 μ L and equal volume of GLBII is added. Size selection of cDNAs is performed as was done for the RNA size selection. 6% PAGE gel pre-running is critical to achieve denaturing conditions as well as heating the samples to 95°C for 3 minutes prior to PAGE separation. cDNAs are selected for insert sizes of ~20-120nts (~85-205nts with RT primer extension) and depending on the input material amount the libraries may be invisible at this step. Gel slices are crushed as above, 300 μ L of Crush Soak Buffer is added and cDNAs are eluted at 50°C overnight on rotation.

cDNA circularization, library qPCR, library size selection, sequencing PCR—

Purification of eluted cDNA is performed as above for RNA elution. After cDNA precipitation samples are resuspended in 16 μ L of water, 2 μ L of 10x CircLigaseII Buffer, 1 μ L of CircLigaseII (Epicentre) and moved to 0.5mL PCR tubes. cDNA circularization takes place at 60°C for 120 minutes in a PCR machine. Circularized cDNA is purified by adding 200 μ L of Buffer PNI and processing as above using MiniElute columns, eluting the cDNA twice in 14 μ L (final ~27 μ L). Samples are initially amplified in a 60 μ L qPCR reaction (27 μ L cDNA, 30 μ L 2x Phusion HF Master Mix, 0.75 μ L of 10 μ M P3_short primer (CTGAACCGCTCTTCCGATCT), 0.75 μ L of 10 μ M P5_short primer (ACACGACGCTCTTCCGATCT), 0.72 μ L of 25x SybrGold). The qPCR machine is programmed as follows: 98°C for 1 minute, 98°C for 15 seconds, 62°C for 30 seconds, 72°C for 45 seconds). After qPCR amplification samples are purified with 600 μ L of Buffer PNI and MiniElute columns as above. Library DNA is eluted twice in 15 μ L (total 30 μ L) and concentrated using a lyophilizer to less than 5 μ L. A second 6% TBE 7M Urea PAGE gel selection is performed as above to remove any PCR dimer products and all short qPCR primers. Gel slices are crushed as above and eluted overnight at 50°C on rotation. Purification of library DNA is performed as above post PAGE gel elution and after precipitation resuspended in 19 μ L of water. A final library PCR amplification is performed for three cycles in 40 μ L reactions (19 μ L library DNA, 0.5 μ L of 10 μ M P3_solexa primer (CAAGCAGAAGACGGCA TACGAGATCGGTCTCGG CATTCCTGCTGAACCGCTCTTCCGATCT), 0.5 μ L of 10 μ M P5_solexa primer (AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACA CGACGCTCTTCCGATCT)) and cleaned up using Agencourt AMPure XP beads (Beckman) according to manufactures protocol and eluted the library in 20 μ L of water. Final library material was quantified on the BioAnalyzer High Sensitivity DNA chip (Agilent) and then sent for deep sequencing on the Illumina HiSeq2500 machine for 1 \times 100bp cycle run.

iCLIP and data analysis

The iCLIP method was performed as described before with the specific modifications below³⁶. v6.5 mESCs were grown as described above and UV-C crosslinked to a total of 0.3J/cm². Whole cell lysates were generated in CLIP lysis buffer (50mM HEPES, 200mM NaCl, 1mM EDTA, 10% Glycerol, 0.1% NP-40, 0.2% TritonX-100, 0.5% N-lauroylsarcosine) and briefly sonicated using a probe-tip Branson sonicator to solubilize chromatin. Each iCLIP experiment was normalized for total protein amount, typically 2mg, and partially digested with RNaseA (Affymetrix) for 10 minutes at 37°C and quenched on ice. Immunoprecipitations of HuR were carried out with Protien G Dynabeads (Life Technologies) and anti-HuR antibody (3A2, Santa Cruz) for 3 hours at 4°C on rotation.

Samples were wash sequentially in 1mL for 5min each at 4°C: 2x high stringency buffer (15mM Tris-HCl pH7.5, 5mM EDTA, 2.5mM EGTA, 1% TritonX-100, 1% Na-deoxycholate, 120mM NaCl, 25mM KCl), 1x high salt buffer (15mM Tris-HCl pH7.5, 5mM EDTA, 2.5mM EGTA, 1% TritonX-100, 1% Na-deoxycholate, 1M NaCl), 1x NT2 buffer (50mM Tris-HCl pH7.5, 150mM NaCl, 1mM MgCl₂, 0.05% NP-40). 3'-end RNA dephosphorylation, 3'-end ssRNA ligation, 5' labeling, SDS-PAGE separation and transfer, autoradiograph, RNP isolation, ProteinaseK treatment, and overnight RNA precipitation took place as previously described³⁶. The 3'-ssRNA ligation adaptor was modified to contain a 3'biotin moiety as a blocking agent. The iCLIP library preparation was performed as described previously³⁶. Final library material was quantified on the BioAnalyzer High Sensitivity DNA chip (Agilent) and then sent for deep sequencing on the Illumina HiSeq2500 machine for 1×75bp cycle run. iCLIP data analysis was performed as previously described.³⁶

RNA Structure Analysis

Sequencing, reads mapping, and data quality control—We generated four replicates for each library (DMSO PolyA, NAI PolyA *in vivo* and *in vitro*). We performed single-end sequencing on Illumina's HiSeq sequencer and obtained approximately 200-million to 600-million raw reads for each replicates, totaling 3.9-billion reads. We collapsed these reads to remove PCR duplicates (only reads that have identical sequences including barcode region are regarded as duplicates). Collapsed reads were then subjected to barcode removal and primer and linker trimming by using Trimmomatic³¹. We mapped trimmed reads to the mouse transcriptome of the Ensembl annotation (build GRCm38.74³²) by using Bowtie2³³. For reads that can be mapped to multiple locations of the transcriptome, we evenly distribute them to up to ten random hits. Finally, we obtained 2.1 billion mapped reads in total. We define the -1 positions of the 5-prime the mapped reads as RT stops, which correspond to modified positions in the NAI libraries, and intrinsic modified (or fragmentation) positions in the DMSO libraries. We defined RT stop coverage as the number of times a base is mapped as a RT stop.

We calculated the expression level of all transcripts in the mouse transcriptome in terms of RPKM. The correlations of transcript expression value (RPKM>0.1) in different replicates are very high (in the range of 0.96 to 1.00). We constructed the background base density profile for each transcript as the sequencing depth of each base in the DMSO libraries. We also calculated the correlation of RT stops for each transcript in different replicates. As shown in Extended Data Figure 5, the correlation is high for most transcripts if we limit to transcripts of average RT stop coverage higher than 2 and regions of background base density higher than 200. So for each library (DMSO poly-A, NAI poly-A *in vivo* and *in vitro*) we combine all four replicates into one for the following analysis.

Reactivity score calculation and construction of structural profile—We performed a 5%-5% normalization for each transcript (i.e., The mean of the RT stops of the second top 5% bases, excluding the 32 bases at the beginning and 32 bases in the end of the transcript, will be normalized to 1. All RT stops will be normalized proportionally).

We defined reactivity score as the subtraction of background RT stops (DMSO libraries) from RT stops of the modified NAI libraries, and then adjusted by the background base density:

$$R = (RT_stop_{NAI} - \alpha RT_stop_{DMSO}) / background_base_density_{DMSO}$$

The score is then scaled into the range of [0, 1], after removing the outliers by 90% Winsorization (the top 5th percentile is set to 1 and the bottom 5th percentile is set to 0). We trained the parameter α on the ribosomal RNA structures, and set it to 0.25 to maximize the correlation of reactivity score R determined by deep sequencing and reactivity score measured in low throughput gel shift experiments.

For each transcript, we defined its structural profile as the vector of base-resolution reactivity scores from the beginning to the end. The valid structural profile of a transcript is limited to regions of RT stop coverage higher than 2 and background base density higher than 200. Finally, we obtained valid structural profiles for, respectively, 19,347 and 13,281 transcripts from *in vivo* and *in vitro*, polyA-selected RNA libraries, among which the majority are messenger RNAs (Extended Data Figure 6).

Metagene analysis of translation sites, pause site, m⁶A sites and protein binding sites—We calculated metagene structure profile around different functional sites by averaging all valid reactivity score R : (1), 10 nts upstream and downstream of the RNA methylation m⁶A site, as determined in by our lab previously³⁴; (2), 25 nts upstream and downstream of the translation pause site determined in the same ribosome profiling experiment; (3), 25 nts upstream and downstream of the RNA methylation m⁶A site, (4) 25 nts upstream and downstream of the binding sites of RNA binding protein Rbfox2³⁵ and HuR (Extended Data Figure 8 and 9. See details below).

In the analysis of differential profiles of icSHAPE reactivity scores around m⁶A and negative control sites, we retrieved a set of target m⁶A sites that have icSHAPE reactivity scores in both wild type and *Mettl3* knock out cells, and defined a set of similar number of non-methylated m⁶A site with the same sequence motifs (GGACU). For both wild type and knock out cells, we calculated the profiles of average reactivity scores in target sites and negative controls and subtracted the latter from the former scores to define the differential icSHAPE profiles.

We calculated *in vivo* and *in vitro* metagene structure profile separately. For each transcript functional sites and their flanking regions, we demand a stringent R score for thousands of transcripts being compared. We generated roughly the same number of negative controls for each set of functional sites. And whenever a sequence motif exists for a functional site, we use that motif in generating the negative control. For example, the same sequence motif GGACU is used to scan the transcriptome and negative controls are randomly selected from the hits excluding regions that are close to a true m⁶A site.

The HuR iCLIP experiments are performed and clusters of binding sites are determined with the pipeline as described previously.³⁶ Threshold 9 (at least 9 unique RT stops are each genomic coordinate) is used to filter for the true binding sites. The highest peak and its

flanking 50 nucleotides in each cluster were retrieved and used to call sequence motifs by using HOMER³⁸, with random sequences of 50 nucleotides from the same set of transcripts as background. The motifs are used as the anchor point in calculating metagene profiles and also used to generate negative controls, using the same protocol as the m⁶A negative control generation described above.

VTD (in vivo and in vitro structure difference) analysis—We defined and calculated the VTD profile of a transcript by subtracting its valid in vitro structural profile from the in vivo one. We calculate the average VTD profiles for all 4096 possible hexamers in our structurome. The overall VTD score of each hexamer is defined as the average score across the 6 bases of the hexamer.

We retrieved sequence motifs of important functional sites, including Kozak sequences (GCCGCC), m⁶A sites (GGACU), miR-290 family 6-mer seed matches (GCACUU, complementary to the seeds), and Rbfox2 binding sites (UGCAUG), and highlighted their VTD scores on the VTD histogram of all hexamers. For sites with ambiguity, for example, m⁶A sites, we took the average of all hexamers that contain GGACU.

We also compiled a resource (Supplementary Table 1 and 2) of VTD scores for all RNA protein binding motifs studied by RNAcompete experiments³⁹, and all mouse microRNA 6-mer seed matches from miRBase⁴⁰. In addition to the VTD scores, for every motif or seed match, we asked three questions by using permutation test: 1), whether the absolute value of the motif (or seed match) VTD is significantly less than a random hexamer, i.e., represents a stable region; 2), whether the motif (or seed match) VTD is significantly smaller than a random hexamer, i.e., represents a region that is more structured in vivo; and 3), whether the motif (or seed match) VTD is significantly bigger than a random hexamer, i.e., represents a region that is more structured in vitro.

Structure-based prediction of m⁶A sites and protein binding sites—We constructed a set of SVM models⁴¹ for the prediction of m⁶A sites and protein binding sites using structural profiles, genomic locations, conservations and their combinations.

The structural profile is limited to the range from the -10 to the +10 position of the m⁶A site or the motifs of the protein binding sites. We used in vivo and in vitro reactivity score separately and jointly in making predictions. We also retrieved a set of genomic features for the prediction of m⁶A sites and protein binding sites, including whether the site is in the 5'UTR, CDS, or 3'UTR, whether it is at the last exon, whether it is at the largest exon, the distance to start codon, the distance to stop codon, the distance to 5' of the splicing junction, etc. In addition, we retrieved the UCSC 60-way phastCons conservation score⁴² for nucleotides in the range from the -10 to the +10 position of the m⁶A site or the motifs of the protein binding sites.

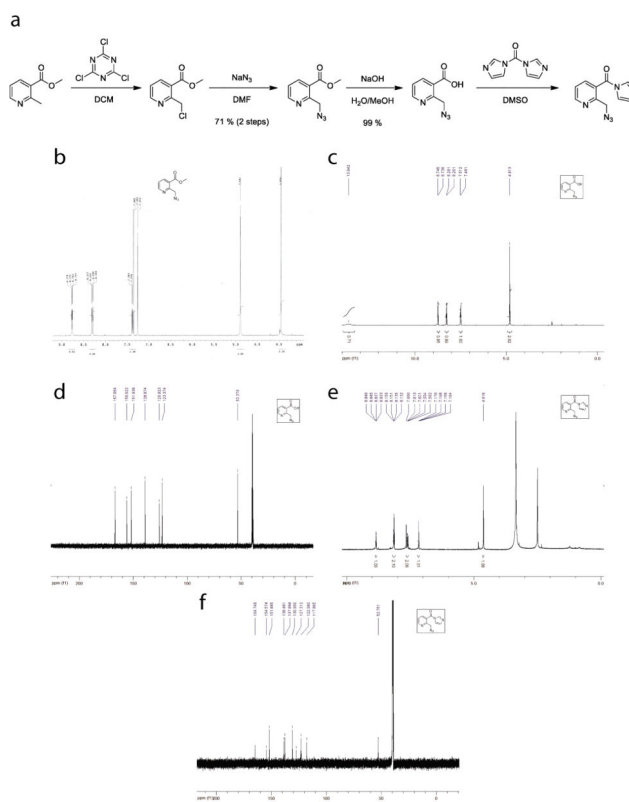
We used the same set of positive and negative controls and the best predictor is selected by using a parameter-searching tool coming along with the *libsvm* package (<http://www.csie.ntu.edu.tw/~cjlin/libsvm/>). We used a five-fold cross validation and calculate the

AUC (area under curve) of the ROC (Receiver operating characteristic) curve to evaluate the performance of the predictors (Extended Data Figure 8).

Sequencing Datasets and Source Code

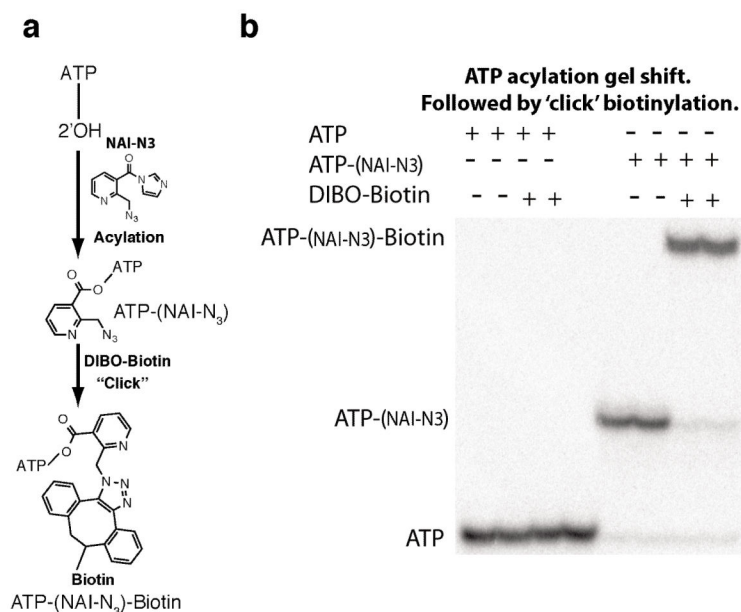
All genomic dataset are deposited on the Gene Expression Omnibus under the accession number GSE64169. Source code used for the icSHAPE analysis is deposited freely available at: <https://github.com/qczhang/icSHAPE>

Extended Data



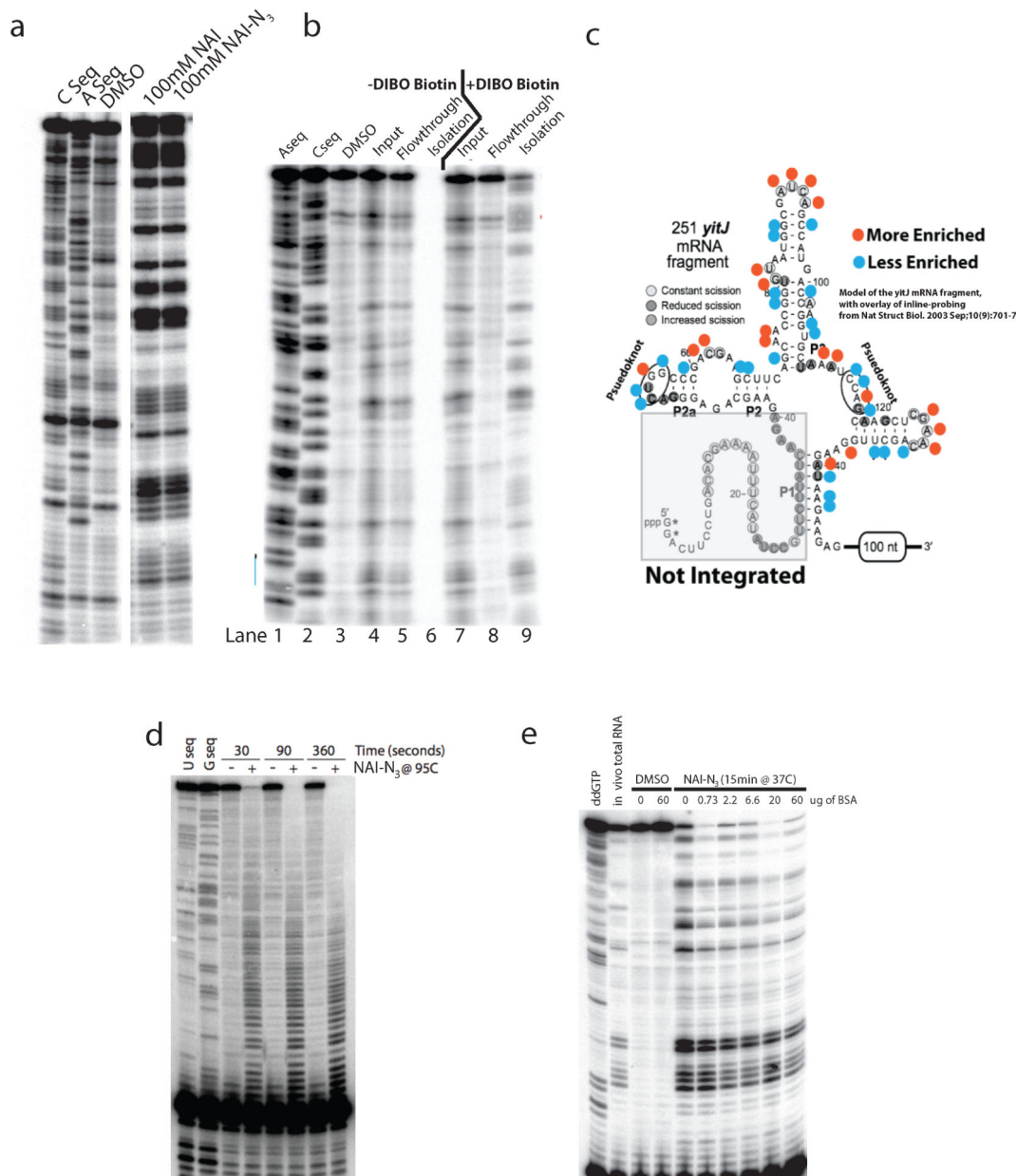
Extended Data Figure 1. Chemical synthesis of NAI-N₃

a, Synthetic scheme for NAI-N₃. **b**, ¹HNMR of Methyl 2-(azidomethyl)nicotinate **c**, ¹HNMR of 2-(azidomethyl)nicotinic acid **d**, ¹³CNMR of 2-(azidomethyl)nicotinic acid **e**, ¹HNMR of 2-(azidomethyl)nicotinic acid acyl imidazole **f**, ¹³CNMR of 2-(azidomethyl)nicotinic acid acyl imidazole.



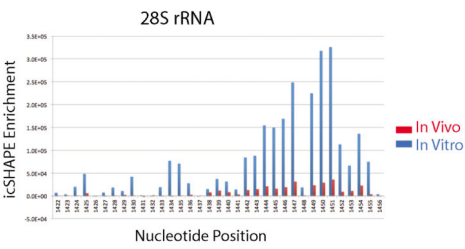
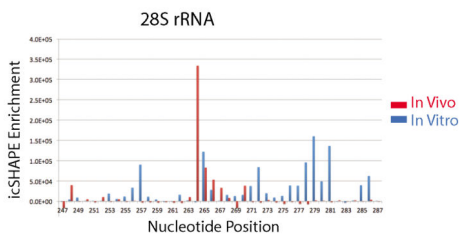
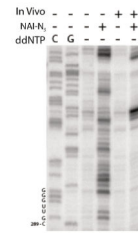
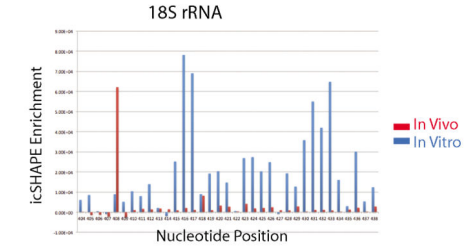
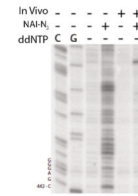
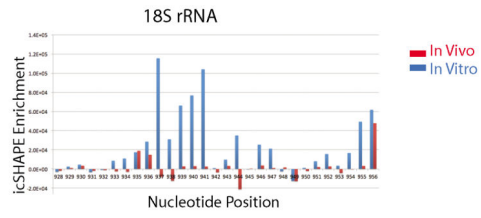
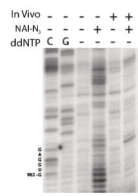
Extended Data Figure 2. NAI-N₃ is a novel RNA acylation reagent that enables RNA purification

a, Chemical schematic of RNA acylation and copper-free 'click' chemistry utilizing NAI-N₃ and dibenzocyclooctyne-biotin conjugate. **b**, ATP acylation gel shift showing ATP acylation and copper-free 'click' chemistry utilizing NAI-N₃ and dibenzocyclooctyne-biotin conjugate.

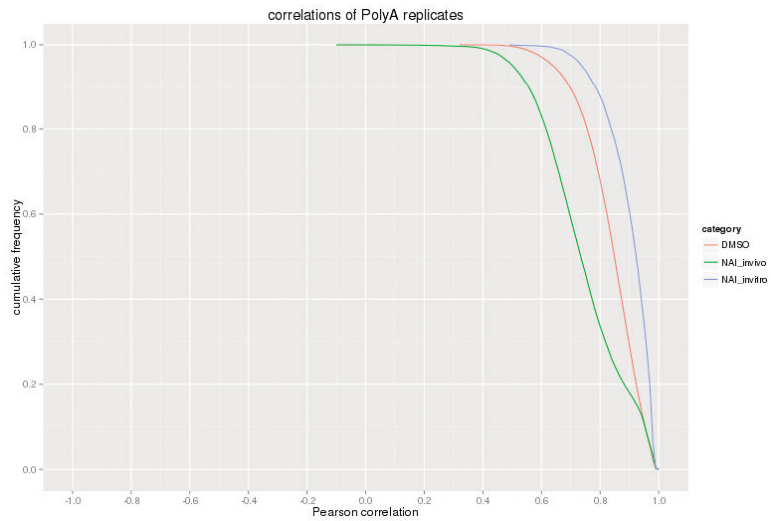


Extended Data Figure 3. NAI-N₃ is a novel RNA acylation reagent accurately reads out RNA structure

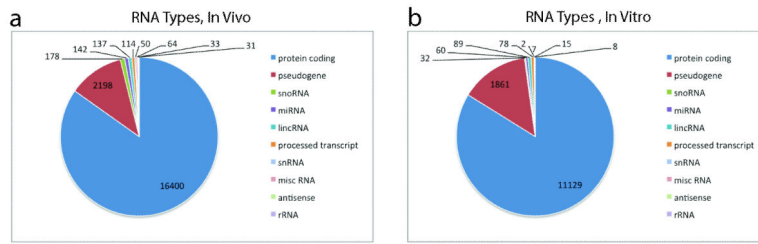
a, Comparative denaturing gel of NAI and NAI-N₃ RNA acylation. **b**, Denaturing gel analysis of cDNAs that originate from the biotin-purification protocol (**Extended Data Figure 1**) **c**, Secondary structure of the SAM-I Riboswitch with enriched residues highlighted in orange and depleted residues highlighted in blue. **d**, Denaturing gel analysis of denatured RNA probed with NAI-N₃ shows even coverage of 2'-hydroxyl reactivity when RNA is unfolded. **e**, Protein titration with BSA, demonstrating no difference in the SHAPE pattern as a function of protein concentration.



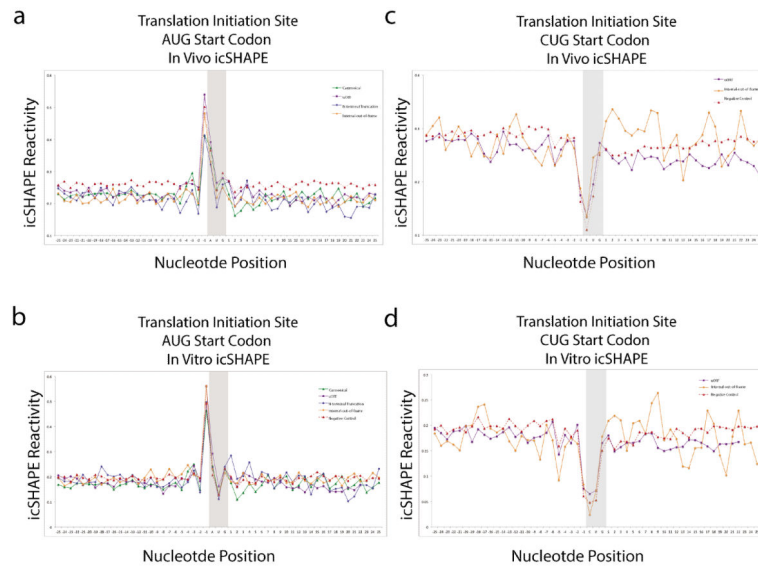
Extended Data Figure 4. icSHAPE is capable of reproducing RNA acylation profiles obtained by manual RNA modification experiments
 icSHAPE (right panels) profiles of ribosomal RNA and compared them to those obtained by manual SHAPE (left panels).



Extended Data Figure 5.
RT stops measured by icSHAPE is very well correlated in different library replicates

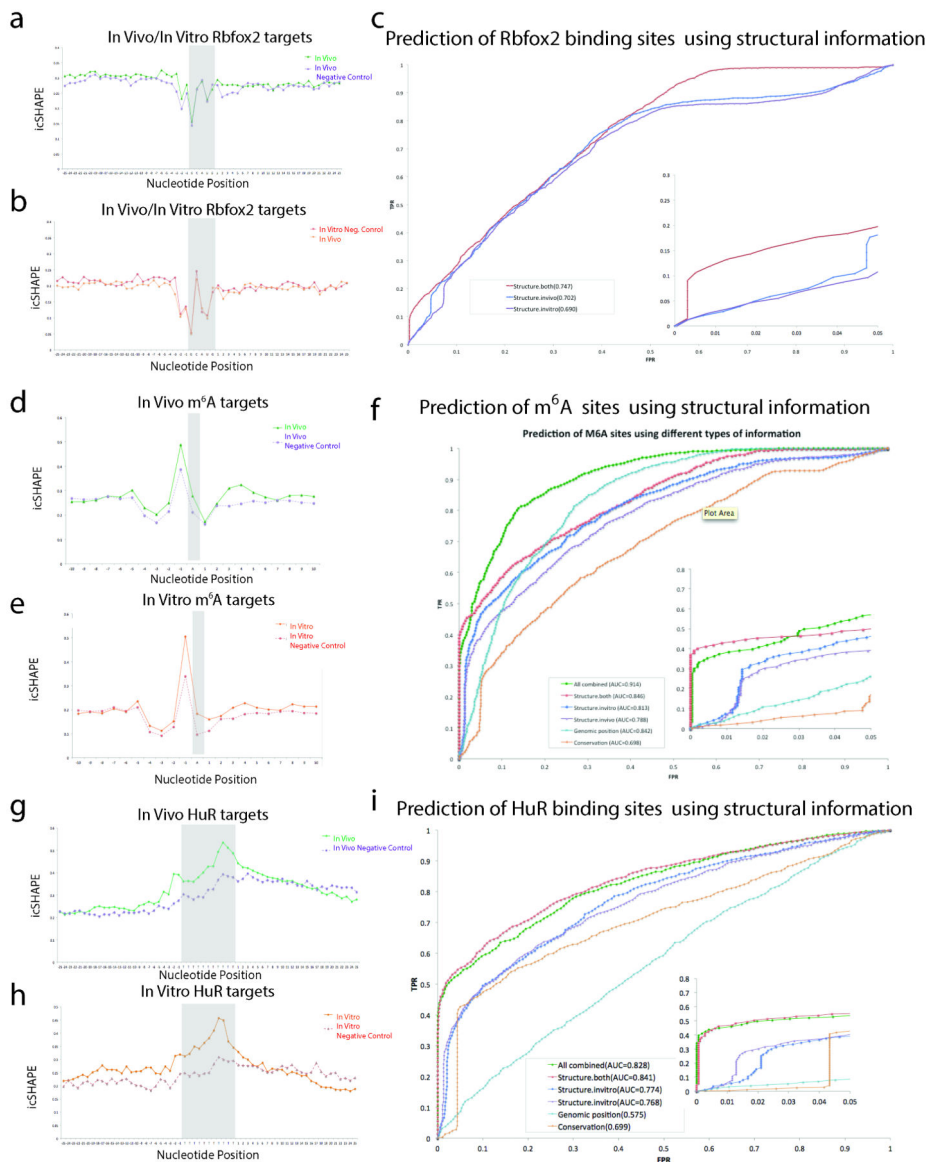


Extended Data Figure 6. icSHAPE is capable of measuring the RNA structure profiles of thousands of RNAs, simultaneously
a, The RNAs represented in polyA selected RNA, *in vivo*. b, The RNAs represented in polyA selected RNA, *in vitro*.



Extended Data Figure 7. Non-AUG start codons are associated with preceding reactivity, and Non-AUG start codons have a different profile, suggesting that RNA accessibility alone is not a sufficient to drive translation

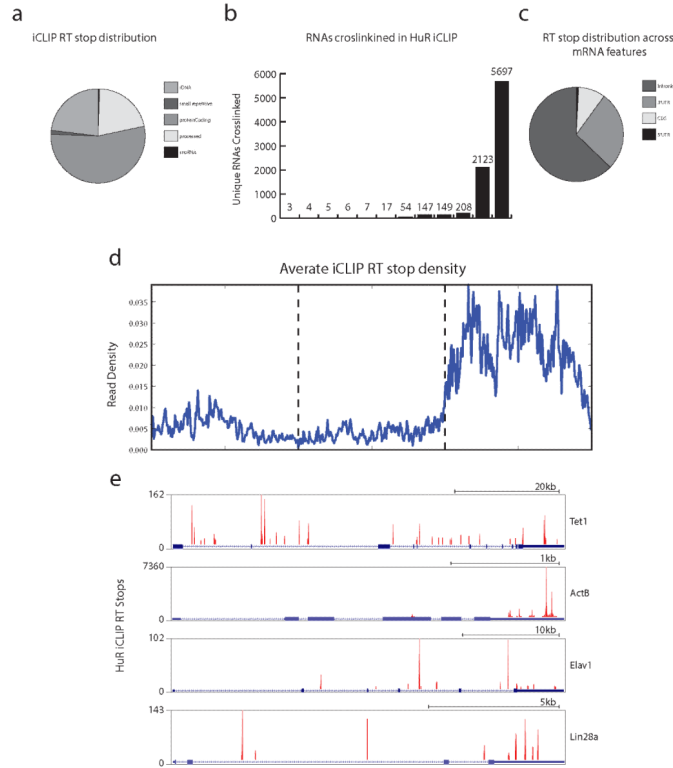
a, icSHAPE profile at AUG start codons, *in vivo*. **b**, icSHAPE profile at AUG start codons, *in vitro*. **c**, icSHAPE profile at CUG start codons, *in vivo*. **d**, icSHAPE profile at CUG start codons, *in vitro*



Extended Data Figure 8. icSHAPE can be used to predict posttranscriptional regulatory elements

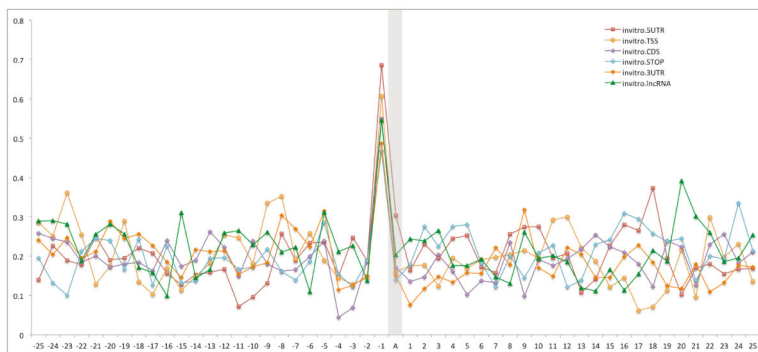
a, icSHAPE profile at Rbfox2 targets, *in vivo*. **b**, icSHAPE profile at Rbfox2 targets, *in vitro*. **c**, ROC curve of Rbfox2 RNA-protein interactions, predicted using icSHAPE profiles. **d**, icSHAPE profile at m⁶A targets, *in vivo*. The negative control is the set of motif instances that are not m⁶A modified **e**, icSHAPE profile at m⁶A targets, *in vitro*. **f**, ROC curve of m⁶A RNA modification sites, predicted using icSHAPE profiles. **g**, icSHAPE profile at HuR

targets, *in vivo*. **h**, icSHAPE profile at HuR targets, *in vitro*. **i**, ROC curve of HuR RNA-protein interactions, predicted using icSHAPE profiles.



Extended Data Figure 9. iCLIP analysis of HuR in mESCs

a, global binding preference of the RBP HuR in mESCs as represented by RT stops across the mouse transcriptome (mm9). HuR mainly binds protein coding, processed, and ribosomal RNAs. **b**, number of unique RNA transcripts bound by HuR. **c**, HuR RT stops distributed across protein coding transcript functional domains. HuR prefers intronic and 3'UTR regions. **d**, Metagene analysis of all HuR bindings sites. Each mRNA region (5'UTR, CDS, or 3'UTR) was scaled to a standard unit width RT stop density across all bound protein coding genes was plotted revealing a clear enrichment for 3'UTR regions in mature protein coding transcripts. **e**, Individual mRNA binding events of HuR to genes important for mESC biology including Tet1, β -Actin, Elav1 (HuR itself), and Lin28a. Discrete binding sites are observed focused in 3'UTR and intronic regions.



Extended Data Figure 10.

m⁶A-associated RNA structure features are preserved, independent of their position along the RNA transcript

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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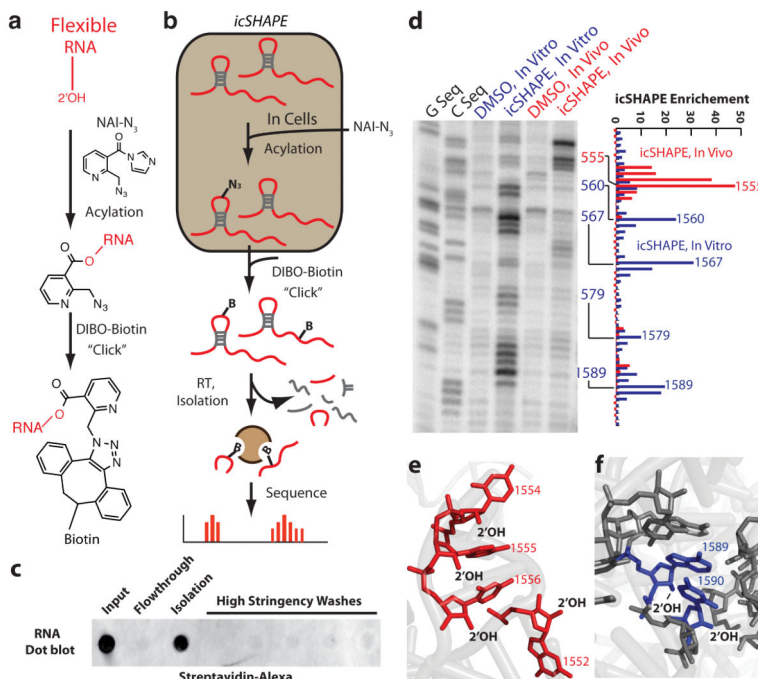


Figure 1. icSHAPE is a Novel and Robust Method for Measuring RNA Structure
a, Chemical scheme for the preparation of acylated RNA, which can be purified by biotin-streptavidin purification. **b**, Schematic of icSHAPE modification and purification steps to generate a sequencing library. **c**, Dot blot of biotin-modified RNA from icSHAPE through streptavidin affinity isolation. **d**, Denaturing gel electrophoresis of icSHAPE on rRNA from mESCs. The corresponding icSHAPE profile, generated from deep sequencing is to the right. **e**, Pymol representation of rRNA, corresponding to regions of icSHAPE that are more reactive *in vivo* (PDB3J3D). **f**, Pymol representation of rRNA, corresponding to regions of icSHAPE that are more reactive *in vitro*.

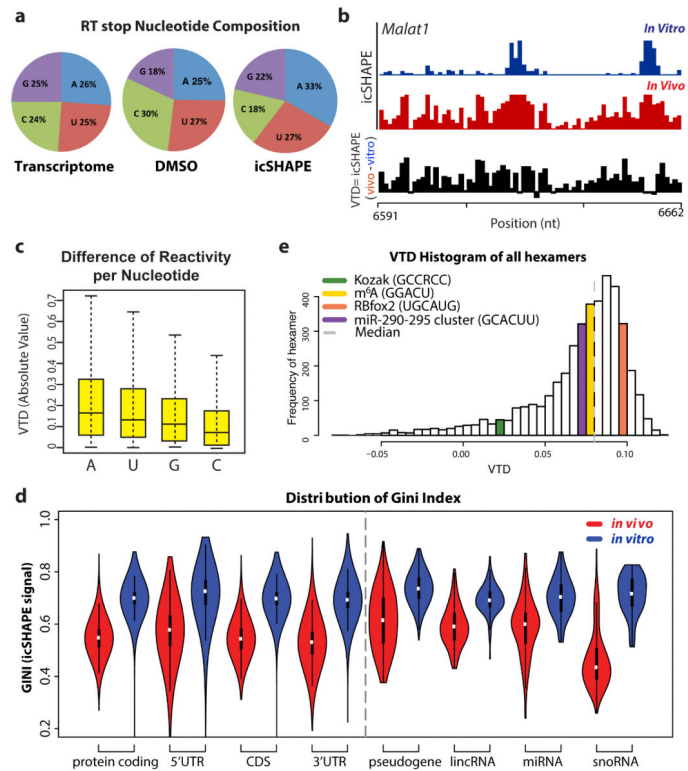


Figure 2. icSHAPE Reveals Unique Structural Profiles for Nucleobase Reactivity and Posttranscriptional Interactions

a, RT stop distribution for the transcriptome, DMSO control, or icSHAPE libraries. **b**, icSHAPE data track and VTD calculation of the MALAT1 RNA (chr19:5796010-5796081). icSHAPE data are scaled from 0 (no reactivity) to 1 (max reactivity). **c**, The VTD distribution for icSHAPE libraries. **d**, GINI index of icSHAPE data *in vivo* vs. *in vitro*. **e**, The distribution of VTD profiles for all hexamer motifs across the transcriptome. Bin locations for several motifs for post-transcriptional regulation are highlighted.

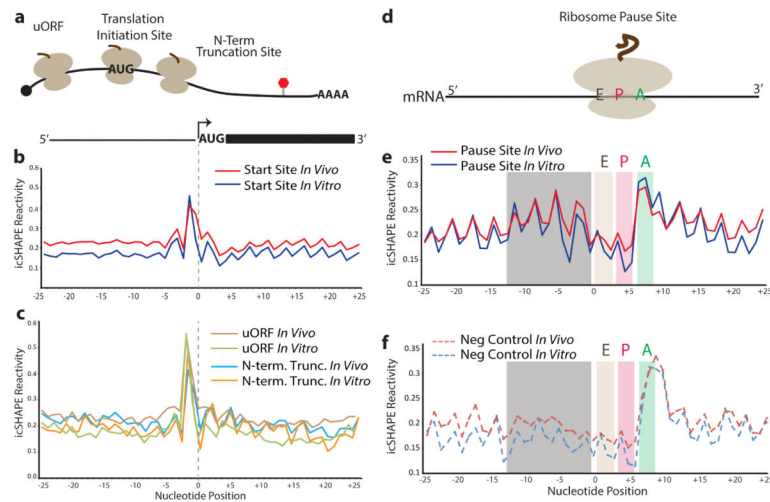


Figure 3. icSHAPE Reveals Structural Profiles Associated with Translation

a, Cartoon representation of ribosomes translating an mRNA. The uORF initiation site is represented by ribosome initiation upstream of the canonical start codon. The canonical start position is demarcated by “AUG”. The N-terminal truncation is represented as a ribosome initiating to the 3’-end of the canonical start “AUG”. **b**, icSHAPE profile at canonical start codon position. **c**, icSHAPE profiles at uORF and N-terminal truncation sites. **d**, Cartoon representation of a paused ribosome and its corresponding A-P-E sites. A: acceptor; P: peptidyl-transferase; E: exit. **e**, The icSHAPE profile at ribosome pause sites. **f**, icSHAPE profile at negative control sites for pause sequences. Gray box highlights a region of structural difference upstream of true pause sites vs. controls.

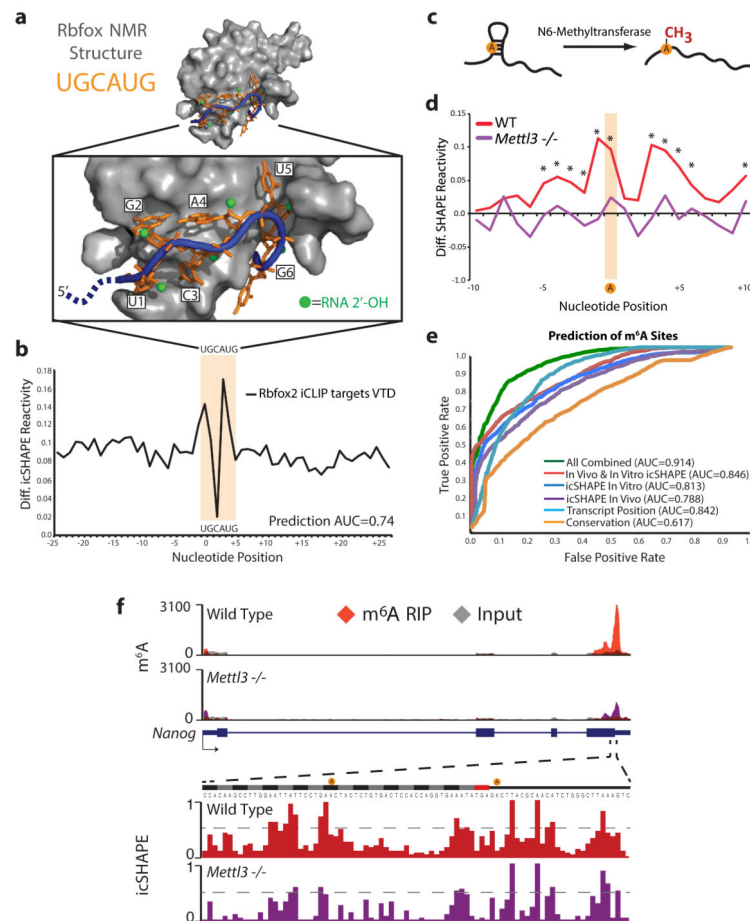


Figure 4. icSHAPE Dynamics Reveals and Predicts Post-transcriptional Interactions

a, Structure of Rbfox1-RNA interaction, highlighting the RNA-protein interface. The RNA is shown with a blue backbone and orange bases; each 2'hydroxyl is green (PDB:2ERR). **b**, The differential icSHAPE profile at Rbfox2 target mRNAs measured *in vivo* vs. *in vitro* maps precisely to the Rbfox binding sites. **c**, Model of interplay between m⁶A and RNA structure. **d**, Differential icSHAPE signal for m⁶A methylated vs. non-methylated sites with the same underlying sequence motif, both *in vivo*. icSHAPE signal from unmodified sites are subtracted from m⁶A-modified sites; asterisks (*) indicate positions with significant differences ($p < 0.05$, FDR < 0.05). Data from wild type and *Mettl3* KO mESCs are plotted for comparison. **e**, ROC curve for prediction of m⁶A sites, incorporating icSHAPE profiles. **f**, Effect of m⁶A on RNA structure on *Nanog* mRNA. Top: location of *Mettl3*-dependent m⁶A sites (highlight in yellow); m⁶A-RIP data from Batista et al²⁵. Bottom: icSHAPE profile of WT and *Mettl3* KO cells.

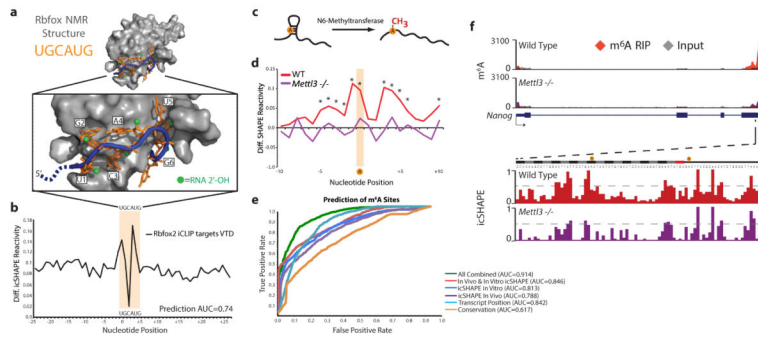


Figure 5.