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The dynamic *N*¹-methyladenosine methylome in eukaryotic messenger RNA

Dan Dominissini^{1,2,*}, Sigrid Nachtergaele^{1,2,*}, Sharon Moshitch-Moshkovitz^{3,*}, Eyal Peer^{3,4}, Nitzan Kol³, Moshe Shay Ben-Haim^{3,4}, Qing Dai^{1,2}, Ayelet Di Segni³, Mali Salmon-Divon³, Wesley C. Clark⁶, Guanqun Zheng⁶, Tao Pan⁶, Oz Solomon^{3,5}, Eran Eyal³, Vera Hershkovitz³, Dali Han^{1,2}, Louis C. Doré^{1,2}, Ninette Amariglio^{3,5}, Gideon Rechavi^{3,4}, and Chuan He^{1,2,6}

¹Department of Chemistry and Institute for Biophysical Dynamics, The University of Chicago, 929 East 57th Street, Chicago, Illinois 60637, USA

²Howard Hughes Medical Institute, The University of Chicago, 929 East 57th Street, Chicago, Illinois 60637, USA

³Cancer Research Center, Chaim Sheba Medical Center, Tel Hashomer 52621, Israel

⁴Sackler School of Medicine, Tel Aviv University, Tel Aviv 69978, Israel

⁵The Mina and Everard Goodman Faculty of Life Sciences, Bar-Ilan University, Ramat Gan 52900, Israel

⁶Department of Biochemistry and Molecular Biology, The University of Chicago, 929 East 57th Street, Chicago, Illinois 60637, USA

Abstract

Gene expression can be regulated post-transcriptionally through dynamic and reversible RNA modifications. A recent noteworthy example is *N*⁶-methyladenosine (*m*⁶A), which affects messenger RNA (mRNA) localization, stability, translation and splicing. Here we report on a new mRNA modification, *N*¹-methyladenosine (*m*¹A), that occurs on thousands of different gene transcripts in eukaryotic cells, from yeast to mammals, at an estimated average transcript

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Correspondence and requests for materials should be addressed to D.D. (dandominissini@gmail.com), G.R. (gidi.rechavi@sheba.health.gov.il) or C.H. (chuanhe@uchicago.edu).

*These authors contributed equally to this work.

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stoichiometry of 20% in humans. Employing newly developed sequencing approaches, we show that m¹A is enriched around the start codon upstream of the first splice site: it preferentially decorates more structured regions around canonical and alternative translation initiation sites, is dynamic in response to physiological conditions, and correlates positively with protein production. These unique features are highly conserved in mouse and human cells, strongly indicating a functional role for m¹A in promoting translation of methylated mRNA.

Four decades after it was first observed as an abundant constituent of the eukaryotic transcriptome, m⁶A has only recently been revealed to broadly affect mRNA metabolism^{1,2}. Consequently, a broader notion has emerged that dynamic chemical modifications of mRNA, now also encompassing pseudouridine (Ψ)^{3–5} and 5-methylcytosine^{6,7}, have vital roles in post-transcriptional gene regulation. With the epitranscriptome, as these modifications are now known, RNA joins both DNA and proteins in being subject to dynamic and reversible chemical control^{8,9}. Unlike the latter two, however, the functions of RNA chemical adducts are only now being determined. It is likely that the complement of mRNA modifications, at present much smaller than that found on transfer RNA (tRNA) and ribosomal RNA (rRNA)¹⁰, is far from complete.

Transcriptome-wide mapping of m¹A

m¹A was identified in total RNA from several species decades ago^{11,12}. It was later characterized in tRNA¹³ and rRNA¹⁴ where it has a major influence on structure and function, owing to the presence of both a methyl adduct and a positive charge under physiological conditions¹⁵. Since m¹A can rearrange to m⁶A under alkaline conditions (Dimroth rearrangement)¹⁶, its signature in mRNA might be erased by the purification and measurement process, simultaneously introducing noise in m⁶A measurements (Fig. 1a and Extended Data Fig. 1a), as earlier attempts to survey mRNA base modifications acknowledged¹⁷. To determine whether m¹A is present in mRNA and to quantify it, we performed quantitative mass spectrometry (LC-MS/MS) of pure mRNA preparations under controlled conditions (Supplementary Note 1, Extended Data Fig. 1b–d) and obtained m¹A/A molar ratios in several human and mouse cell lines and tissues, both normal and cancerous. Our results unequivocally show that m¹A is present in pure mammalian mRNA, at m¹A/A ratios ranging from approximately 0.015% to 0.054% in cell lines and up to 0.16% in tissues (Figs 1b, 5d and Extended Data Fig. 1e). Metabolic labelling of cells with deuterated methionine (d₃-Met) results in detection of d₃-m¹A in mRNA, suggesting that S-adenosyl-methionine (SAM) is the methyl donor (Extended Data Fig. 1f). Although these ratios are smaller than the m⁶A/A and Ψ/U ratios in the same samples (Fig. 1b), the absolute and fractional levels of m¹A in mRNA (see stoichiometry below) are sufficiently high to exhibit potential regulatory functions of selectively methylated transcripts.

We adapted an antibody-based approach, methylated RNA immunoprecipitation sequencing (MeRIP-seq), termed here m¹A-seq, for transcriptome-wide localization of m¹A sites and coupled it to an orthogonal chemical method based on Dimroth rearrangement to obtain high-resolution m¹A maps (Supplementary Notes 2, 3; Fig. 1c–e and Extended Data Fig. 1g–m). We applied m¹A-seq to polyA⁺ RNA from HeLa cells in two replicates and

determined the methylated positions (m^1A peaks) using the MACS peak-calling algorithm¹⁸. We identified 7,154 peaks (fold change (FC) ≥ 2 , false discovery rate (FDR) $\leq 5\%$) in 4,151 coding and 63 non-coding adequately expressed (that is, above the 1st quartile of expressed genes) gene transcripts that occurred in both replicates (Fig. 2a, Supplementary Tables 1, 4). On average, each methylated gene carries 1.4 peaks, with most of these genes (over 70%) methylated only once (Fig. 2b). In a conservative estimate, 35% of adequately expressed genes are methylated, attesting to the breadth of the phenomenon. Of note, non-coding genes are significantly underrepresented among methylated genes ($P = 1 \times 10^{-140}$, hypergeometric test). Methylated genes constitute a progressively larger fraction of genes as expression increases (Fig. 2c), and are significantly enriched in several gene ontology terms related to translation and RNA processing (Extended Data Fig. 2a, b).

As most methylated transcripts harbour a single m^1A , methylation stoichiometry can be estimated based on the change in transcript levels following immunodepletion. We used microarrays to determine relative transcript abundance in HepG2 RNA before and after depletion m^6A . Methyl groups ($-CH_3$) are in red and the positive charge (+) on with anti- m^1A antibody, and found that the average methylation level in single m^1A -containing genes is $\sim 20\%$ (Fig. 2e, Extended Data Fig. 2c, Supplementary Table 7). The presence of m^1A in a substantial fraction of mRNA transcripts suggests a potentially sizeable effect on mRNA metabolism.

m^1A associates with translation initiation sites and the first splice site

We found that m^1A peaks appear in all segments of transcripts—5' untranslated region (UTR), coding sequence (CDS) and 3' UTR—but are markedly under-represented in the latter (\sim fivefold depletion relative to chance, $P < 1 \times 10^{-200}$, hypergeometric test, Fig. 2d). m^1A -seq of two additional cell lines in replicates, HepG2 and HEK293, produced similar distribution patterns (Fig. 2b, d, Extended Data Fig. 2d–g and Supplementary Tables 1, 4). We noted that while input sequence reads have a typical distribution along transcripts, m^1A immunoprecipitation reads pile up towards their 5' ends (Extended Data Fig. 3a). To investigate this further, metagene profiles were generated in which gene segments were rescaled proportionally so that all segments of one kind appear to have the same length. This revealed that m^1A clusters sharply around the start codon to produce a remarkably similar profile in the three human cell lines (Fig. 2f). Over 50% of the peaks reside within a stretch of 300 nucleotides centred on the start codon (Extended Data Fig. 3b–g).

Ribosome profiling¹⁹, and a recently modified version that captures initiating 80S ribosomes^{20,21}, have uncovered a surprising variety of alternative translation initiation sites (TISs). We examined our data sets from this perspective and found that methylated transcripts have, on average, more alternative TISs than non-methylated ones (1.98 versus 1.59 in HeLa; 2.37 versus 1.69 in HEK293), and that the number of m^1A sites per transcript positively correlates with the number of alternative TISs per transcript (Extended Data Fig. 4a–c). Furthermore, transcripts with upstream TISs are more likely to be methylated within the 5' UTR, whereas transcripts with downstream TISs tend to be methylated within the CDS (Extended Data Fig. 4d, e). The distances of alternative TISs and respective m^1A peaks from the annotated start codon are well correlated, particularly those of upstream TISs and

5' UTR m¹A peaks, suggesting that m¹A is associated with alternative TISs as well (Extended Data Fig. 4f–h). In summary, m¹A in human mRNA seems to be associated with a subset of TISs, both canonical and alternative, in all three cell lines examined.

To gain a deeper understanding of the underlying transcript attribute or processing event that potentially guides m¹A deposition to create the observed distribution, we binned methylated transcripts based on the exon that harbours the start codon and plotted the distribution of m¹A relative to the start codon. We observed an intriguing dissociation: when the start codon is in the 2nd or 3rd exons, m¹A tends to occur closer to the transcription start site (TSS) compared to when the start codon is in the 1st exon (Extended Data Fig. 3h). Whereas 5' UTRs of transcripts with a start codon in downstream exons are on average longer (323 versus 189 nucleotides), their first splicing event tends to occur closer to the TSS (204 versus 359 nucleotides), mirroring m¹A behaviour. Therefore, we plotted the distribution of m¹A relative to the TSS in bins based on the length of the 1st exon and observed that m¹A moves away from the TSS as the length of the first exon increases (Extended Data Fig. 3i). These observations suggest that neither the TSS nor the start codon 'anchors' m¹A, but rather the first splice site does. We substantiated this by plotting m¹A relative to the nearest splice site (which is also the first splice site for 85% of m¹A peaks) and observed convergence of m¹A distribution in all start codon bins mostly upstream of the splice site, engendering the hypothesis that the first splicing reaction somehow guides m¹A deposition (Fig. 2g, Extended Data Fig. 3j). Analysis based on the smaller high-resolution m¹A trough data set recapitulated all the salient features of the methylome (Extended Data Fig. 3k–q, Supplementary Note 3 and Supplementary Table 6).

Conserved features of methylated transcripts

We next sought to identify sequence patterns that may define or associate with m¹A peak locations. GC content, a key determinant of RNA structure, is unusually high in m¹A peak areas that appear in all three transcript segments (Extended Data Fig. 5a). Several motif detection algorithms showed enrichment of GC-rich sequences with variable adenosine content, yet did not identify any specific motif *per se* (other than the 'AUG' itself which is expected to appear in peaks centred on the start codon) (Extended Data Fig. 5b). This suggests that a combination of sequence and structure features may specify methylation positions, perhaps similar to the case of m¹A deposition in tRNA^{22,23}. We were able to identify and score single-nucleotide m¹A positions based on the change in mismatch rates between (1) untreated and alkaline-treated (m¹A-to-m⁶A rearranged) immunoprecipitation sequence reads and between (2) input and immunoprecipitation (Supplementary Table 5). A sequence frequency logo of identified positions (Fig. 3a) and representative individual sequences (Extended Data Fig. 5c) demonstrate that m¹A sites are embedded in GC-rich environments. Using the m¹A trough data set we also identified a GA-rich motif that was present in 10–15% of the cases (Extended Data Fig. 3r).

Attributes of the 5' UTR such as length, GC content, and associated minimum free energy (MFE), are known to affect the efficiency of translation initiation²⁴. 5' UTRs of methylated transcripts differ significantly from those of non-methylated ones in that their GC content is higher (median of 71% compared to 63.9%) (Extended Data Fig. 5d) and their length-

adjusted MFE (aMFE) is lower (median of $-43.5 \text{ kcal mol}^{-1}$ compared to $-38.4 \text{ kcal mol}^{-1}$) (Fig. 3b, Extended Data Fig. 5e). We employed a sliding window approach to further characterize the continuous structural landscape of the AUG window (± 150 nucleotides) in terms of GC content, MFE and experimental parallel analysis of RNA structure (PARS) score (Fig. 3c–e, Extended Data Fig. 5f, g). Both the experimental and calculated parameters agree that m¹A decorates highly structured AUG windows.

Evolutionary conservation and dynamics

Evolutionary conservation typically indicates a common fundamental biological function. We show above that m¹A is present in murine mRNA at comparable levels to those in human (Fig. 1b). Therefore, m¹A-seq was applied to RNA of mouse liver, mouse embryonic fibroblasts (MEFs) and mouse embryonic stem cells (mESC) (Supplementary Tables 2, 4). A previously unreported single m¹A position in mouse 28S rRNA was identified in an orthologous location to the known m¹A position in human 28S rRNA (Extended Data Fig. 6a–c). The mouse m¹A methylome closely resembles that of human: approximately 15% of adequately expressed genes are methylated, the majority of which carry a single m¹A (Fig. 4a), and the percentage of methylated genes increases with expression level (Fig. 4b). m¹A in mouse mRNA is non-randomly distributed along transcripts with dense clustering around translation initiation sites, mirroring the profile in human (Fig. 4c–d, Extended Data Fig. 6d–l), including the association with the nearest splice site (Fig. 4e, Extended Data Fig. 6m). Similarly, m¹A sites reside within a GC-rich context (Extended Data Fig. 7a, b), and the 5' UTRs of methylated transcripts have a higher GC content and a lower aMFE than those of all other (unmethylated) transcripts (Fig. 4f and Extended Data Fig. 7c–f).

We systematically assessed the extent of m¹A peak conservation between human and mouse and found that it extends to the gene level: out of 1,338 identified mouse liver m¹A peaks that could be mapped to adequately expressed, orthologous positions in human HepG2 cells, 441 had an m¹A peak at the orthologous position, representing 33% overall conservation ($P < 1.3 \times 10^{-310}$, Mann–Whitney *U* test) (Fig. 4g, Extended Data Fig. 7g). The 5' UTR and a 300-nucleotide window centred on the AUG codon exhibit the highest degree of conservation with 44–46% conserved peaks, compared to 3–16% in other gene segments (Fig. 4g).

We applied m¹A-seq to RNA of *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe* to further examine this mark in simple eukaryotes, and found it to be present in their mRNA transcriptomes, albeit without the characteristic mammalian pattern (Extended Data Fig. 8a–g, Supplementary Tables 3, 4). Importantly, upon transfer to a nitrogen-source deficient medium the *S. pombe* methylome exhibited noticeable changes in the identity of methylated transcripts (some transcripts gained m¹A while others lost it), providing an example of how physiological conditions dynamically shape m¹A (Extended Data Fig. 8h, i). These results further demonstrate the conservation of m¹A as a mark in eukaryotic mRNA, although the distribution pattern and functions in lower eukaryotes could be different from mammals, as also observed for m⁶A^{25,26}.

Next, using quantitative LC-MS/MS we surveyed a range of physiological perturbations that could alter the total m¹A level in mRNA of mammalian cells. We found that glucose or amino acid starvation reduced m¹A level by approximately threefold and twofold, respectively, as early as 4 h; that heat shock increased m¹A level by approximately 1.5-fold at 4 h; and that serum starvation resulted in only a modest change within the same time frame (Fig. 5a, Extended Data Fig. 9a, b). Of note, m⁶A levels measured in parallel do not change, or change slightly in the opposite direction, suggesting that the response is m¹A-specific (Fig. 5a). m¹A-seq performed after 4 h of glucose starvation or heat shock uncovered a more complex picture: while the overall number of m¹A peaks decreased (by 17%) and increased (by 23%), respectively, mirroring the LC-MS/MS measurement, still some individual positions behaved in the opposite direction (Fig. 5b, c; Extended Data Fig. 9c, d).

The m¹A levels across various tissues obtained from a single mouse also showed significant variability, with kidney and brain possessing the highest m¹A level (about threefold higher than that measured in liver, for example) (Fig. 5d). A comparison between tissues from lean (wt/wt) and obese mice (ob/ob) revealed that brains of the former contain 75% more m¹A (Fig. 5d). In sum, m¹A in mRNA is a dynamic modification in response to certain stress conditions and potentially physiological signalling, and its level varies between different tissues.

SAM-dependent methyltransferases that install m¹A on tRNA and rRNA are known in eukaryotes^{23,27}. These or related homologues could mediate mRNA m¹A methylation. FTO, the first RNA demethylase to be discovered, belongs to the AlkB family of proteins, and catalyses oxidative demethylation of m⁶A in mRNA²⁸. Other members of the AlkB family are known to perform oxidative demethylation of m¹A and m³C lesions in DNA/RNA. In particular, ALKBH3 was shown to have a strong preference for single stranded DNA/RNA, and to demethylate m¹A in tRNA and mRNA *in vitro*^{29,30}. In line with previous biochemical observations, overexpression of wild-type ALKBH3, but not of an inactive mutant, reduced m¹A levels in mRNA (Extended Data Fig. 9e). These results suggest that ALKBH3 and its homologues are likely candidates that could reverse m¹A in mRNA and contribute to its dynamic regulation.

m¹A correlates with elevated translation

Structure and sequence features of the 5' UTR and both sides of the start codon affect the scanning process and thereby the efficiency of translation initiation and early elongation^{31,32}. We show here that m¹A is a widespread and conserved post-transcriptional modification that is associated with TISs in thousands of mammalian transcripts characterized by structured 5' UTRs. The precise localization of m¹A in an area of obvious regulatory importance, coupled with its ability to restructure RNA in other cases (such as tRNA³³) and to affect RNA–protein interactions, points to possible involvement in translation initiation.

We used published ribosome profiling and proteomic data sets, obtained from the same cell lines and tissues used for m¹A profiling, to assess possible effects of m¹A on translation (see

Methods). The translation efficiency¹⁹ (TE) and ribosome release score³⁴ (RRS) of genes with a single m¹A peak in the start codon window compared to genes devoid of m¹A altogether are significantly higher (~1.2–1.4, on average) in both human and mouse cells (Extended Data Fig. 10). Furthermore, correlation of m¹A with protein levels normalized to mRNA abundance reveals a similar trend: in all cell types examined, methylated genes tend to have higher protein levels than non-methylated genes (Fig. 6a–f). The average fold-change in protein levels between methylated and non-methylated genes in all cell types—and across all gene expression bins—is ~1.7. Considering that the average fraction of methylated transcripts is ~20%, the isolated true effect of m¹A could be even higher. The contribution of m¹A remains significant even after accounting for multiple covariates that are known to affect translation (ANCOVA, Supplementary Table 8).

Discussion

N¹-methylation on RNA adenosine occurs at the Watson–Crick interface and is expected to affect RNA base pairing³⁵. In addition to an added methyl group, this modification also endows the modified adenosine with a positive charge under physiological conditions (Fig. 1a), which could markedly alter RNA structure and protein–RNA interactions. The impact of m¹A on the structure and function of tRNA is well documented, where a single m¹A can be essential for correct functional folding of a tRNA molecule³³. A single charge difference has been shown to induce ~100–1,000-fold affinity difference of protein–DNA interactions³⁶. We show here that m¹A is a modification occurring on thousands of mRNA transcripts in several human and mouse cell types, as well as on hundreds of transcripts in two yeast species. Applying m¹A-seq, a method relying on immunoprecipitation of m¹A-containing RNA fragments, we found a striking enrichment of m¹A around the start codon in human and mouse mRNA. Taking advantage of the Dimroth rearrangement, which converts m¹A (mutation prone in reverse transcription) to m⁶A (low mutation rate in reverse transcription³⁷) with alkaline treatment, we confirmed the accumulation of m¹A around the start codon at high resolution.

Transcriptome-wide mapping of m¹A revealed unique features of its distribution that strongly indicate functional roles: (1) m¹A resides close to both canonical and alternative TISs found mostly upstream of the first splice site; (2) m¹A is present in highly structured regions around the start codon; (3) m¹A positively correlates with translation efficiency and protein level; (4) the distribution pattern and m¹A peaks are highly conserved in all mouse and human cell types examined; and (5) m¹A is dynamic in response to certain stress, and potentially physiological, signals, and its level varies across mouse tissues. Together, these attributes suggest a positive and dynamic role for m¹A in translation initiation in mammalian cells. The importance of RNA structure in translation initiation is well appreciated³⁸. A positively charged m¹A may alter the secondary/tertiary structure of mRNA around TISs by blocking Watson–Crick base pairing or introducing charge–charge interactions. Alternatively, potential binding proteins may specifically recognize m¹A and facilitate translation initiation of methylated transcripts in a way analogous to the role of the m⁶A reader YTHDF1 in translation enhancement³⁹. Either way, it is tempting to speculate that m¹A can modulate the strength of a particular translation initiation site, thereby affecting the balance between alternative and canonical initiation sites.

In contrast to m⁶A, which is enriched around the stop codon and in the 3' UTR^{1,2}, m¹A preferentially associates with the start codon region in mammalian mRNA. These two marks could complement each other in controlling mRNA metabolism and translation. A single m⁶A modification usually induces ~10-fold change in either duplex stability or protein–RNA affinity, as would be expected by the presence of an extra methyl group^{40,41}. The propensity of m⁶A to cluster at 3' UTRs also enables a wider range of dynamic tuning than a single m⁶A site could offer^{1,2}. The unique chemical properties of m¹A, with both a positive charge and a methyl group, potentially allow each individual modified site to have a stronger effect than m⁶A in terms of RNA structure or protein–RNA interaction.

The discovery of this new mRNA mark that positively correlates with translation adds an exciting new dimension to post-transcriptional gene expression regulation mediated through mRNA modifications. While our paper was in proof, we were alerted to a study by Chengqi Yi and colleagues reporting similar findings on the m¹A methylome in a human cell line⁴².

METHODS

Cells and tissues

Human HeLa (cervical adenocarcinoma), HepG2 (hepatocellular carcinoma) and HEK293 (embryonic kidney) cell lines (ATCC) and primary mouse embryonic fibroblasts (MEFs) (C57BL/6, ATCC) were maintained in DMEM (Thermo Fisher Scientific) containing 25 mM glucose, 4 mM -glutamine, supplemented with 100 U ml⁻¹ penicillin, 100 µg ml⁻¹ streptomycin and 10% FBS. Mouse embryonic stem cells (mESCs) were maintained in FBS-free N2B27-based media, as previously described⁴³. Cells were routinely checked for mycoplasma contamination. Mouse tissues were obtained from wild-type and ob/ob C57BL/6 mice. For metabolic labelling, methionine-devoid DMEM was supplemented with L-methionine-(methyl-d₃) (Sigma-Aldrich).

Cell treatments

HepG2 cells were maintained as described above and subjected to the following growth conditions: (1) heat shock (43 °C for 4 h); (2) glucose starvation (glucose-depleted medium for 4 h); (3) amino acid starvation (amino acid-depleted medium, with or without 200 µM *S*-adenosyl-methionine (SAM) supplementation, for 4 h); and (4) serum starvation (serum-depleted medium for 4 h), following which they were immediately harvested in Cell Lysis Solution (5 Prime).

Yeast strains and growth conditions

Wild-type (BY4741) *S. cerevisiae* cells were grown vegetatively in YPD medium to mid-log phase and harvested. Wild-type heterothallic (Sp1) *S. pombe* cells were grown vegetatively in YES medium to mid-log phase and harvested. For induction of sporulation/arrest, Sp1 cells were washed and transferred to Edinburgh Minimal Medium lacking NH₄Cl (EMM-N) for 4 h and harvested.

RNA purification

Total RNA from cells in culture and mouse tissues was purified using PerfectPure RNA Cultured Cell Kit (5 Prime) and PerfectPure RNA Tissue Kit (5 Prime), respectively, and DNase-treated. Enrichment of polyadenylated RNA (polyA⁺ RNA) from total RNA was carried out using one round of GenElute mRNA Miniprep Kit (Sigma-Aldrich). Ribo-Zero Gold rRNA Removal Kit (Illumina) was used to deplete rRNA from polyA⁺ RNA before LC-MS/MS.

Dot blot assays

RNA Oligonucleotides were synthesized in-house with either m¹A, m⁶A or A at a single internal position (5'-AC(m¹A/m⁶A/A)UG-3'), spotted onto a nylon membrane (GE Healthcare) in decreasing amounts (1,000, 200, 40 and 8 pmol) and UV-crosslinked. Membranes were blocked with 5% non-fat dry milk in 1 × PBST (blocking buffer) for 1 h at 25 °C, and incubated overnight with either mouse anti-m¹A antibody (1 µg ml⁻¹, MBL) or rabbit anti-m⁶A antibody (1 µg ml⁻¹, Synaptic Systems) in 1 × PBST at 4 °C. Following extensive washing with 1 × PBST, membranes were incubated with either HRP-conjugated goat anti-mouse IgG or anti-rabbit IgG antibody (1:2,500, Thermo Fisher Scientific) in blocking buffer for 1 h at 25 °C. Membranes were washed in 1 × PBST, developed with ECL substrate (Thermo Fisher Scientific) and imaged with FluorChem imager (Protein Simple) or X-ray film. Competitive dot blots were performed on separate membranes spotted with 75 pmol of the m¹A-containing oligonucleotide by co-incubation of anti-m¹A antibody with increasing concentrations of either m¹A or m⁶A competitor mononucleoside (0, 1, 2 and 4 µM).

m¹A detection and quantitation

Purified RNA (see 'RNA purification' section above) was subjected to liquid chromatography-tandem mass spectrometry (LC-MS/MS) for detection and accurate quantitation of m¹A, essentially as previously described⁴⁴. 200–400 ng of purified RNA was digested by P1 nuclease (Wako, 2 U) in 40 µl of buffer containing 25 mM NaCl and 2.5 mM ZnCl₂ for 2 h at 37 °C. Subsequently, 5 units (1 µl) of Antarctic Phosphatase (New England Biolabs) and 1 × Antarctic Phosphatase reaction buffer were added and the sample was incubated for another 2 h at 37 °C. The sample was then filtered (0.22 µm, Millipore) and injected into a C18 reverse phase column coupled on-line to Agilent 6410 QQQ triple-quadrupole LC mass spectrometer in positive electrospray ionization mode. Quantitation was performed based on nucleoside-to-base ion transitions (268-to-136 for A; 282-to-150 for m⁶A and m¹A (retention times of 2.5 and 0.9 min, respectively); 285-to-153 for d₃-m¹A and d₃-m⁶A; 245-to-179 for Ψ; 245-to-113 for U) using standard curves of pure nucleosides and stable isotope-labelled internal standards (d₃-m¹A and d₃-m⁶A), as previously described^{45–47}.

Synthesis of stable isotope-labelled internal standards

d₃-m¹A was synthesized following a previously described procedure for the synthesis of m¹A⁴⁸, replacing CH₃I with CD₃I. d₃-m⁶A was synthesized following a previously described procedure for the synthesis of m⁶A^{49,50}, replacing CH₃NH₂ with CD₃NH₂.

m¹A-seq

Mapping of m¹A in total or mRNA was performed using m¹A-seq, which is based on the previously described m⁶A-seq protocol⁵¹ with the following modifications: RNA fragmentation was performed using RNA Fragmentation Reagents (Thermo Fisher Scientific) for 15 min at 70 °C to minimize m¹A-to-m⁶A rearrangement followed by column purification (RNA Clean & Concentrator, Zymo). Anti-m¹A antibody (MBL) was pre-coupled to Protein G Dynabeads (Thermo Fisher Scientific) and used to immunoprecipitate methylated RNA fragments for 3 h at 4 °C. Fragment elution was carried out by either digestion with Proteinase K (Sigma-Aldrich; 5 units in 5 mM Tris pH 7.5, 1 mM EDTA and 0.25% SDS for 1.5 h at 37 °C) followed by TRIzol (Thermo Fisher Scientific) extraction of the supernatant and ethanol precipitation, or by competitive elution using m¹A mono-nucleoside (Santa Cruz Biotechnology), as in m⁶A-seq. Induced m¹A-to-m⁶A rearrangement was achieved by incubating the input and immunoprecipitation fragments in an alkaline buffer (50 mM Na₂CO₃, 2 mM EDTA, pH 10.4) for 1 h at 50 °C or 60 °C followed by column purification (RNA Clean & Concentrator, Zymo) (Extended Data Fig. 1m). Immunoprecipitated RNA fragments and comparable amounts of input were subjected to first-strand cDNA synthesis using the NEBNext Ultra RNA Library Prep Kit for Illumina (New England Biolabs). Sequencing was carried out on Illumina HiSeq2500 according to the manufacturer's instructions, using 10 pM template per sample for cluster generation, TruSeq SR Cluster kit v3 (Illumina), TruSeq SBS Kit v3-HS (Illumina) and TruSeq Multiplex Sequencing primer kit (Illumina). Summary of read numbers for each replicate can be found in Supplementary Table 4.

Validation of m¹A-seq

Reads were aligned to the 28S ribosomal RNA sequences of human (RefSeq NR_003287.2), mouse (RefSeq NR_003279/NCBI X00525) and *S. cerevisiae* (SGD RDN25-2, ID: S000006485.) using Bowtie2⁵² with local alignment option (--local). Enrichment was calculated as the coverage ratio of immunoprecipitation to input. Mismatch rate was assessed using mpileup tool of the SAMtools software package (version 0.1.18)⁵³ for calling variants over the entire 28S transcript. The frequency of non-adenosine bases at each adenosine position and the coverage of each base in the transcript were reported.

RNA expression level

Reads per kilobase of transcript per million mapped reads (RPKM) expression levels of RefSeq genes were calculated using HTseq-count⁵⁴ and R package edgeR⁵⁵. Fragments per kilobase of transcript per million mapped reads (FPKM) values were calculated by the CUFFLINKS tool (version 2.2.1)⁵⁶. Only genes whose expression level was above the first quartile were considered adequately expressed and used for downstream analyses. Adequately expressed genes (RPKM) were divided into 10 expression bins for correlation with methylation and the fraction of methylated genes in each bin was calculated.

Peak calling

Adaptors and low quality bases were trimmed from raw sequencing reads using cutadapt⁵⁷. Reads were aligned to the relevant genome (human-hg19, mouse-mm10, *S. pombe*-

ASM294v2.29, and *S. cerevisiae*-S288C with UTR data downloaded from http://genie.weizmann.ac.il/pubs/PARS10/pars10_catalogs.html) using Tophat2 (version 2.0.12)⁵⁸. Peaks enriched in immunoprecipitation over input experiments were identified using MACS2 (version 2.1.0.20140616)¹⁸. MACS2-identified peaks were intersected with a database of exons of the relevant genome (RefSeq annotation). Peaks were allocated to the feature containing the segment with which they share the largest overlap. Peaks falling in intergenic sequences or having an overlap shorter than 50 nucleotides were excluded from further analyses. For each cell type only peaks identified (FC ≥ 2 or FC ≤ 4 , as indicated, FDR ≤ 0.05) in replicates were considered. Common human peaks were defined as peaks independently identified in both HeLa and HepG2 RNA. Negative peaks were identified by switching the immunoprecipitation and input samples.

Coverage analysis

The coverage of unique reads at each nucleotide position at the transcriptomic level, in immunoprecipitation and input reads, was calculated and reported. Only transcripts in which at least one nucleotide exceeded a minimum depth of 10 reads in immunoprecipitation were used in further analysis for reduction of background noise. For whole-transcript coverage plots, each transcript was divided into 100 bins of equal length, and the median coverage for each bin across all transcripts was calculated.

m¹A stoichiometry

HepG2 mRNA was subjected to immunoprecipitation with anti-m¹A antibody in 2 biological replicates. Input and immunodepleted (unbound sup) RNA samples (75 ng each) were hybridized to PrimeView human gene expression microarrays (Affymetrix) and expression levels were determined. Genes expressed below the first quartile (as determined by mRNA-seq, see above) were set as the minimal intensity limit (blank), and only genes with intensities above this threshold were considered. To correct for technical loss of RNA during the different steps of the procedure, the average ratio of sup/input intensity was calculated for never-methylated genes (considering only genes that were unmethylated in all m¹A-seq experiments) and used for ratio correction of methylated genes carrying one m¹A peak. Fractional methylation level per gene was calculated as $1 - (\text{corrected sup/input ratio})$.

Identification of m¹A-induced mismatches in m¹A peaks

Identification of sequence variants in m¹A peaks due to misincorporation at the m¹A position during reverse transcription was carried out by determining the base composition at each position within peak regions using bam-readcount (<https://github.com/genome/bam-readcount>). Identified adenosine positions (according to the encoding DNA strand) were then filtered to exclude known genomic polymorphism sites (dbSNP version 141) and identified A-to-I editing sites (Radar database). Variants were considered if the mismatch rate was greater than 0.1 and the overall coverage was greater than 20 reads. For HepG2 samples the fold enrichment of mismatch rate in untreated over rearranged sequence reads was calculated.

Trough identification

To identify points of lower coverage inside peak areas, we converted the bam files of the immunoprecipitation libraries to BigWig files and subjected them to PeakSplitter⁵⁹ with options of $-valley$ 0.65 and $-cutoff$ 30. The trough was identified as the minimal point ± 10 nucleotides, between the summits of identified sub-peaks. Only troughs reoccurring in 3 biological repeats were considered.

Bidirectional conversion of genomic and transcriptomic coordinates

Genomic coordinates of all 5' UTRs, CDS and 3' UTRs of RefSeq transcripts were downloaded from the University of California, Santa Cruz genome browser (UCSC) table browser in BED format (hg19 and mm10 for human and mouse, respectively). Transcripts with IDs corresponding to more than one genomic location were discarded, as were transcripts of non-coding genes. Custom, in-house, awk and perl scripts were then used to convert transcriptomic coordinates to genomic coordinates and vice versa.

Definition of peak middle point

MACS2-generated genomic m¹A peak coordinates were converted to transcriptomic coordinates. Neighbouring peaks (at a distance of up to 10 nucleotides) on the same transcript were merged using mergeBed from the BEDTools package⁶⁰. This allowed the unification of peaks overlapping more than one exon that were divided by MACS2 due to the presence of introns in genomic coordinates. The middle of the resulting transcriptomic peak was defined as peak middle and referred to as m¹A peak position in further analyses.

Metagene profiling

The relative position of each peak's middle point was located in the corresponding transcript segment (5' UTR, CDS or 3' UTR). Each segment was assigned a value corresponding to its average length fraction out of the overall transcript length. The corresponding relative peak position was then deduced within each segment and plotted along a normalized metagene profile with respect to the TSS, AUG start codon and stop codon positions. The distribution of m¹A peaks with respect to the canonical AUG start codon, TSS, first and nearest splice sites, and stop codon was also drawn in actual nucleotide distances.

Sliding window analysis of AUG start codon window

Structural features of a 300-nucleotide window centred on the AUG start codon were compared between methylated and non-methylated RefSeq genes. GC content was calculated as the average percent in a sliding window of 3 nucleotides. Local free energy (G) was calculated in a 30-nucleotide consecutive jumping window (in steps of 10 nucleotides) using the RNAfold tool in the ViennaRNA package⁶¹ with default parameter settings.

Minimum free energy (MFE) calculation

The 5' UTR sequence of the predominant isoform of each gene (the highest expressed isoform in the input sample) was used to calculate GC content, MFE, length-adjusted MFE

(aMFE) and MFE density (MFE_{den}) using the ViennaRNA package⁶¹ and as previously described⁶².

PARS-seq and PARS scores calculations

Parallel analysis of RNA structure (PARS) was performed as previously described⁶³ on HepG2 polyA⁺ RNA. Reads were mapped to the human transcriptome (UCSC known canonical genes) using Bowtie2⁵² with local alignment option (--local). Only uniquely aligned reads were used for analysis. Read start positions were counted for each base in the transcriptome, as the read start is +1 to the nuclease cleavage site and therefore informative regarding local RNA 2D structure⁶⁴. Only reads with more than 5 consecutive bases matching the transcript at the start of the read were counted. Read counts were normalized to the respective library size. PARS-scores were then calculated based on the following equation:

$$\text{PARS-score}(i,j)=\log_2(V1'_{ij}/S1'_{ij})$$

$V1'_{ij}$ and $S1'_{ij}$ are the normalized read-start counts at base (i) in transcript (j) for V1 or S1, respectively.

PARS scores at the region around the start-codon (± 150 nucleotides from the AUG) were averaged and used for a metagene analysis.

Motif search

To search for a motif, 101-nucleotide sequences in the AUG start codon window, centred on m¹A peaks in HeLa and mouse liver, were used. Compatible background sequences were used to prevent skewing of the results due to over-representation of known motifs in the region of the start codon, such as Kozak sequences and the AUG. The findMotifs program from the HOMER package⁶⁵ was then run using the settings -rna -len 6,7,8,9,10 -noweight. The same peak areas were used with MEME⁶⁶ for *de novo* motif enrichment analysis, with the settings -dna -minw 6 -maxw 10 -evt 0.01 -maxsize 1,000,000. A similar analysis was performed on m¹A peaks falling in the CDS with background sequences randomly generated from non-methylated exons. 199 sequences of 20 nucleotides, surrounding the mutations identified above, were used to create a frequency plot of the nucleotides surrounding the mutation sites using WebLogo⁶⁷.

Comparison of m⁶A and m¹A profiles

m¹A and m⁶A peaks^{1,42,68} were intersected using BEDTools⁶⁰. Similarity between sets was also evaluated by comparing the metagene profiles of the two modifications.

TIS correlation

TISs data was downloaded from <http://www.ncbi.nlm.nih.gov/sra?term=SRA160745> (ref. 21) and genomic locations were converted to transcriptomic locations (as described above). For each alternative TIS, the closest m¹A peak was identified and the distance of both from the canonical AUG start codon was calculated. Correlation of alternative TISs with the

number of m¹A peaks per gene was carried out by dividing the genes containing alternative TISs to bins according to the number of peaks per gene.

Gene ontology (GO) enrichment

Methylated gene RefSeq IDs were uploaded to DAVID Bioinformatics Resources (<http://david.abcc.ncifcrf.gov>) and functional enrichment analysis was performed using all adequately expressed genes as background. Resulting enriched GO terms were restricted to fold enrichment ≥ 1.5 , Bonferroni corrected $P \leq 0.005$.

Human–mouse conservation

m¹A mouse liver peaks were converted to the homologous coordinates in the human hg19 genome, using the LiftOver tool of the UCSC genome browser⁶⁹. To assess the significance of these results we chose at random an equal number of 200 bp sized areas from within mouse genes that have a human orthologue. The areas were chosen from all gene segments in the same proportion as observed in the original data set. Randomly chosen areas were similarly converted and intersected with HepG2 m¹A peaks. To calculate the statistical significance the randomization was repeated 1,000 times and Mann–Whitney U test was used.

ALKBH3 overexpression

Plasmids encoding wild-type and mutant (D193A) FLAG-tagged human ALKBH3 were transfected into 293 Freestyle cells using polyethylenimine (PEI; Polysciences, Inc.) at a ratio of 3 μ l PEI per 1 μ g plasmid DNA. After 24 h, cells were pelleted and washed once with cold PBS before being lysed in Cell Lysis Solution (5 Prime). mRNA was purified and digested for LC-MS/MS analysis as described in the ‘RNA purification’ and ‘m¹A detection and quantitation’ sections above.

Translation efficiency (TE) and ribosome release score (RRS)

Sequencing data was downloaded: for HEK293, mouse liver and MEFs - SRA160745²¹, <http://www.ncbi.nlm.nih.gov/sra?term=SRA160745>; for mESCs - GSE30839¹⁹, <http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE30839>. Reads were aligned and FPKM values were calculated (see above). For TE calculation, the Ribo-Seq FPKM of the most highly expressed transcript of each gene was divided by the mRNA-Seq FPKM of the same transcript. Only transcripts with FPKM and Ribo-Seq FPKM above first quartile were considered. TE data for mESCs was downloaded as-is from the original paper¹⁹. TE and RRS for HeLa were calculated using our published data⁴⁴. RRS values were calculated using mRNA-Seq and Ribo-Seq reads as described previously^{34,42}.

Protein abundance

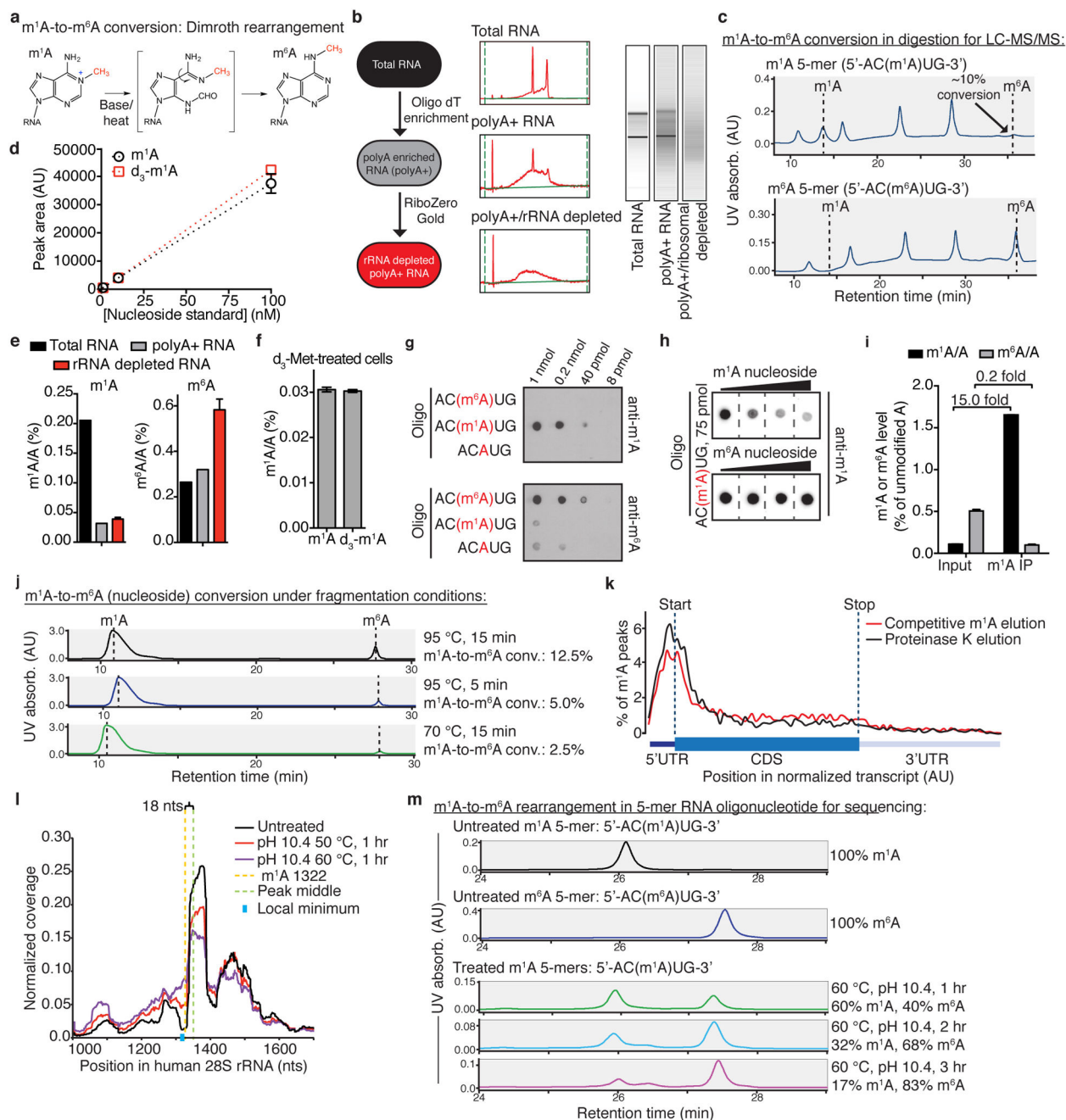
All protein abundance data analysed in this study was downloaded from the MOPED database (Multi-Omics Profiling Expression Data, <https://www.proteinspire.org>): HeLa, HepG2, HEK293 (Experiment ID: Geiger_MCP_2012)⁷⁰, and mouse liver (Experiment ID: Baylor_mouse_liver_profiling), or from Supplementary Data (mESCs⁷¹ and MEFs⁷², corrected as described⁷³). For all cell types, excluding mESCs, we used the normalized

abundances as provided. For mESCs, raw intensities were divided by each leading protein molecular weight. Protein abundance data was then \log_2 transformed and subjected to an analysis of variance (ANOVA) with mRNA expression (FPKM) percentile bins for start m¹A and non-m¹A genes. For MEFs, the published mRNA expression data was used along with the corresponding protein abundance. Genes with mRNA expression below the lowest expressed gene containing m¹A or with FPKM < 0.2 were discarded.

Analysis of covariance (ANCOVA)

ANCOVA was calculated for genes methylated at the AUG start codon window (start m¹A) versus unmethylated genes (non-m¹A) as a fixed factor against \log_2 protein abundances, while cleaning the contribution of the following covariates: \log_2 mRNA expression, TIS Kozak score⁷⁴, MFE calculated by RNAfold and divided by 5' UTR length, G-quadruplex motif in 5' UTR^{75,76}, 5' TOP motif in 5' UTR from UTRdb (<http://utrdb.ba.itb.cnr.it/>), number of conserved miR sites (PCT score < 0) from TargetScan⁷⁷, and number of u/dTISs and u/dAUGs in human and mouse form QTI-seq²¹. The remaining factors, % GC Content in 5' UTR, length of 5' and 3' UTRs, and number of exons in 5' and 3' UTRs were downloaded and calculated from the UCSC genome browser (hg19 and mm10). We considered the most expressed RefSeq genes in our data for unique transcript-associated parameters and gene symbols for general features. Variables were examined for linearity with protein output and for normal distribution. Log transformations were employed where appropriate. Tolerance was kept above 0.2 for all variables.

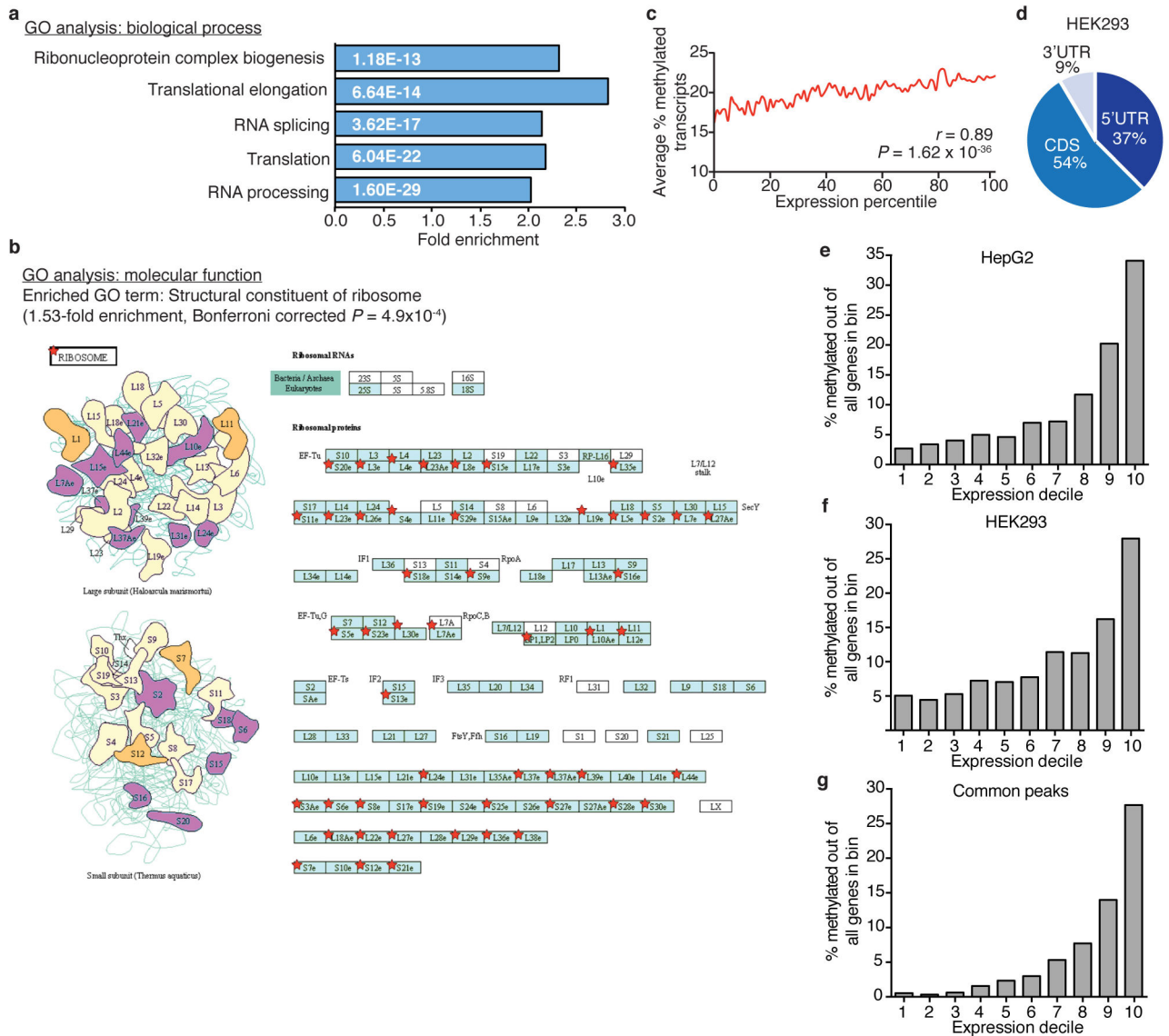
Extended Data



Extended Data Figure 1. Experimental conditions for detection, quantitation and sequencing of m¹A in mRNA

a. Schematic illustration of m¹A-to-m⁶A rearrangement (Dimroth rearrangement) under alkaline conditions at elevated temperatures. **b.** mRNA purification scheme before LC-MS/MS m¹A quantitation (left). Corresponding RNA electrophoresis profiles obtained by Agilent 2100 Bioanalyzer (middle and right). **c.** Monitoring m¹A-to-m⁶A rearrangement levels during sample preparation for LC-MS/MS. Synthetic 5-nucleotide long RNA

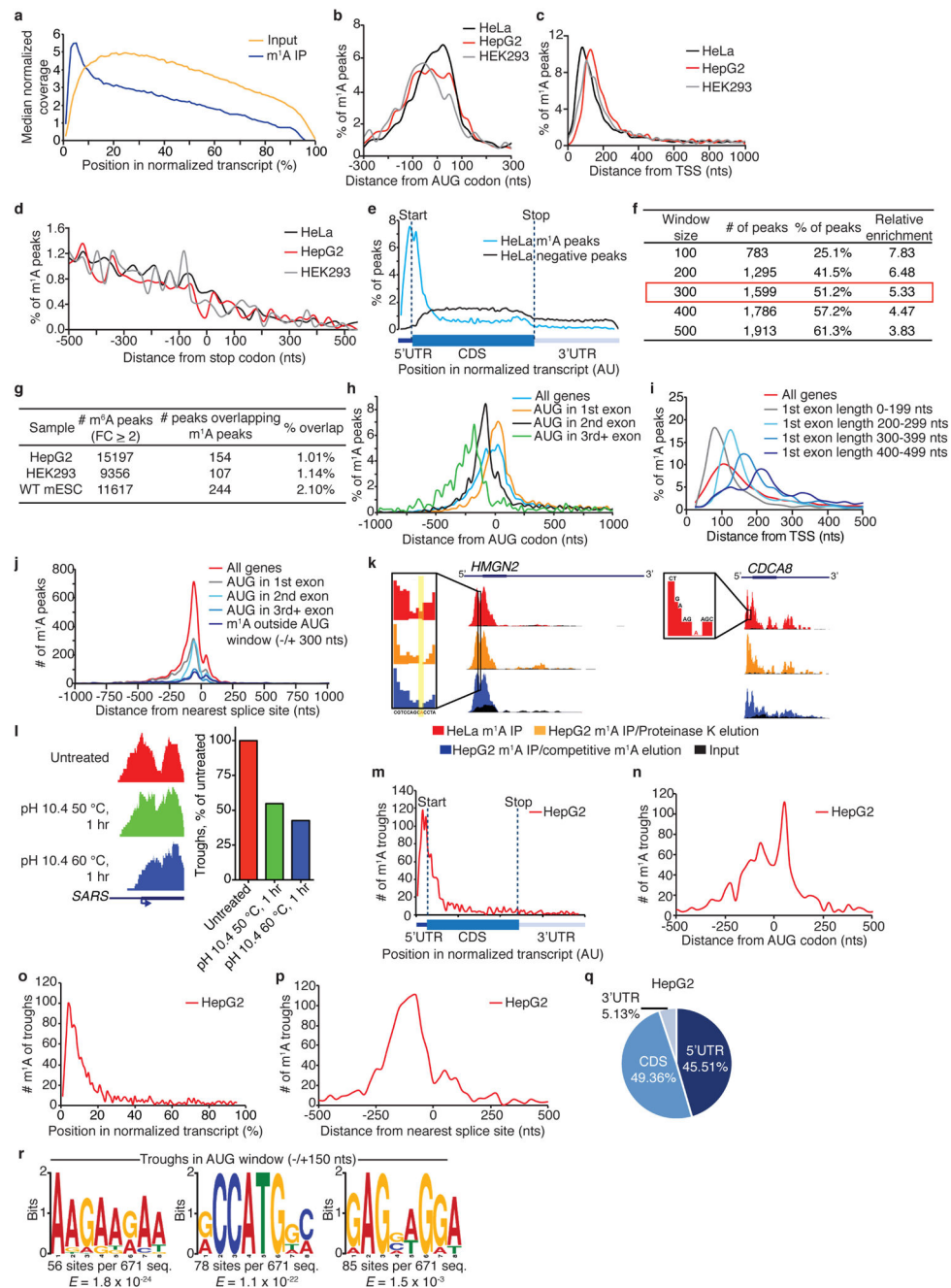
oligonucleotides containing m¹A (upper panel) or m⁶A (lower panel) were digested to mononucleotides, dephosphorylated (see methods) and analysed by HPLC-UV. Only minimal (<10%) m¹A-to-m⁶A rearrangement was observed (arrow). **d**, Standard curves for m¹A and deuterium-labelled m¹A (d₃-m¹A) demonstrate very similar detection sensitivity in LC-MS/MS. Mean values ± s.e.m. are shown, *n* = 3. **e**, The change in m¹A/A and m⁶A/A molar ratios (%) during the purification scheme outlined in **b**. Mean values ± s.e.m. are shown, *n* = 3. **f**, LC-MS/MS of mRNA isolated from HepG2 cells labelled with deuterated methionine (d₃-Met) for 24 h detects d₃-m¹A, suggesting *S*-adenosyl-methionine (SAM) is the methyl donor. Mean values ± s.e.m. are shown, *n* = 3. **g**, Dot blots demonstrating high anti-m¹A antibody specificity. Increasing amounts (indicated above the top blot) of synthetic RNA oligonucleotides containing m¹A, m⁶A or unmodified A residues were spotted onto a membrane and probed with either anti-m¹A or anti-m⁶A antibodies. Anti-m¹A antibody detects m¹A and does not exhibit cross-reactivity with m⁶A or A (upper blot); anti-m⁶A antibody demonstrates low, yet detectable, cross-reactivity with both m¹A and A (lower blot). For blot source data, see Supplementary Fig. 1. **h**, Competitive dot blots were performed on separate membranes spotted with 75 pmol of synthetic m¹A-containing RNA oligonucleotide. Whereas increasing concentrations of free m¹A mononucleoside progressively attenuate anti-m¹A binding, increasing concentrations of free m⁶A mononucleoside do not. For blot source data, see Supplementary Fig. 1. **i**, Quantitative LC-MS/MS demonstrates m¹A enrichment—and m⁶A depletion—following immunoprecipitation (IP) with anti-m¹A antibody compared to total RNA input. Mean values ± s.e.m. are shown, *n* = 3. **j**, Monitoring m¹A-to-m⁶A rearrangement levels under different RNA fragmentation conditions for use in m¹A-seq. Pure m¹A mononucleoside in 1×fragmentation buffer (see Methods) was subjected to the conditions specified to the right of the chromatograms and directly analysed by injection to HPLC-UV for rearrangement to m⁶A. **k**, Comparison of competitive m¹A elution and Proteinase K elution of immunoprecipitated m¹A-containing RNA fragments from anti-m¹A-coupled magnetic beads shows that the two elution modes are equivalent. **l**, Identification of the known m¹A site in position 1322 of human 28S rRNA validates the accuracy of m¹A-seq and the use of peak middle points as a close approximation for m¹A sites. Partial m¹A-to-m⁶A rearrangement increases the coverage around this site. **m**, Conditions for induced m¹A-to-m⁶A rearrangement of RNA oligonucleotides that maintain RNA integrity for use in m¹A-seq. A synthetic 5-nucleotide long RNA oligonucleotide of the sequence 5'-AC(m¹A)UG-3' was subjected to various base/heating conditions (indicated to the right of the chromatograms) and directly analysed by injection to HPLC-UV for rearrangement to 5'-AC(m⁶A)UG-3'. Incubation at pH 10.4, 60 °C for 1 h results in rearrangement to m⁶A in 40% of oligonucleotides; longer incubation times result in increased rates of rearrangement. Chromatograms of untreated RNA oligonucleotides appear above and mark the expected retention times.



Extended Data Figure 2. Characterization of human m¹A-methylated genes

a, Gene ontology (GO) analysis of methylated HeLa genes relative to all adequately expressed genes (above the 1st quartile) reveals enrichment of biological processes related to translation and RNA metabolism. Fold-enrichment and P values are indicated for each category. **b**, GO analysis of molecular functions reveals enrichment of structural constituents of the ribosome. Scheme based on an illustration obtained from DAVID bioinformatics website of the KEGG human ribosome pathway. Red stars indicate methylated genes in the pathway. Colouring of the boxes and ribosome constituents is according to KEGG pathway maps showing interacting proteins and hyperlinks to gene entries that can be reached through http://www.genome.jp/kegg-bin/show_pathway?hsa03010. **c**, The average fraction of methylated transcripts (stoichiometry) increases with gene expression level. r and P values are indicated, Pearson correlation. **d**, Pie chart presenting the fraction of m¹A peaks in each of three non-overlapping transcript segments (5' UTR, CDS and 3' UTR) in HEK293

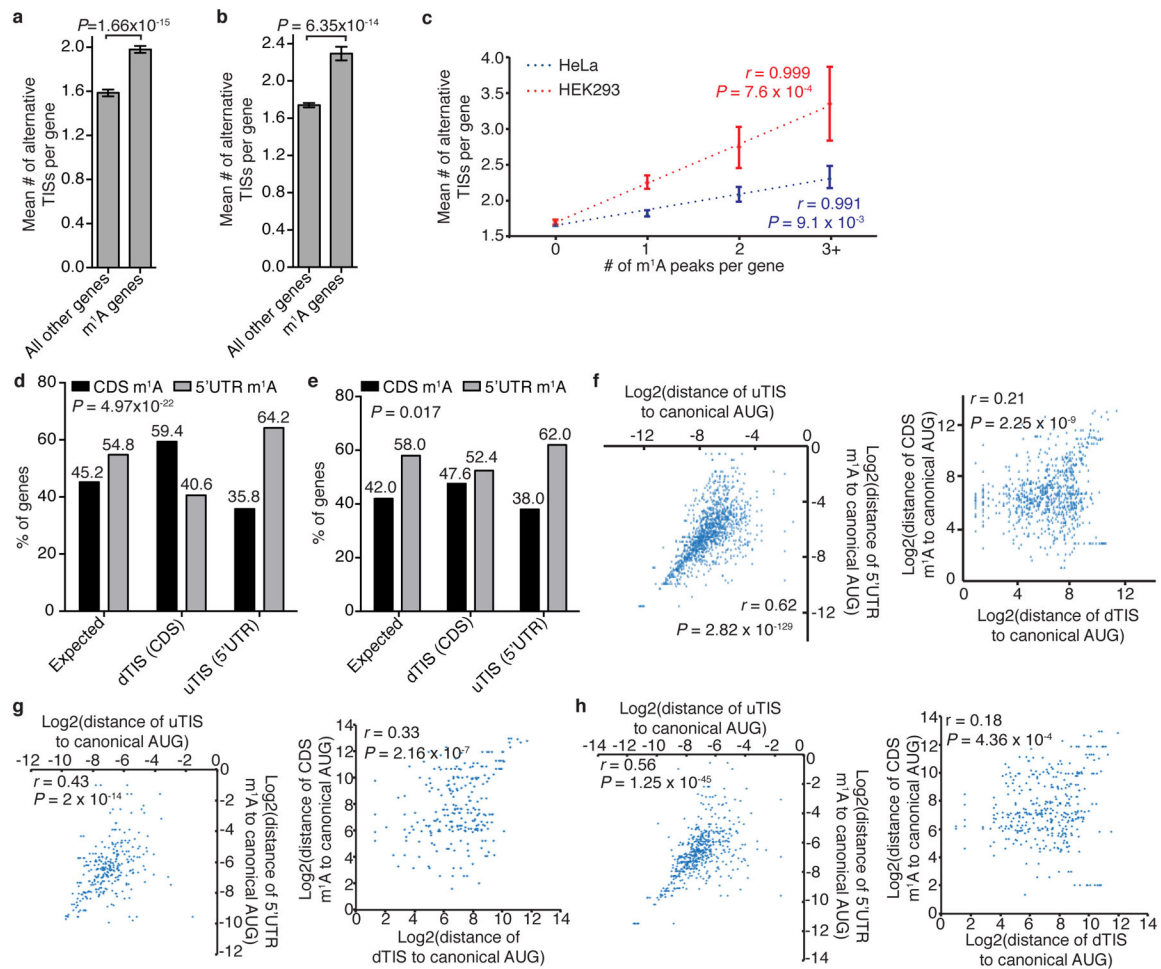
cells. **e–g**, The fraction of methylated genes increases with gene expression levels in HepG2 (**e**), HEK293 (**f**) and common human peaks (see Methods) (**g**).



Extended Data Figure 3. Metagenome analyses of human m¹A peaks

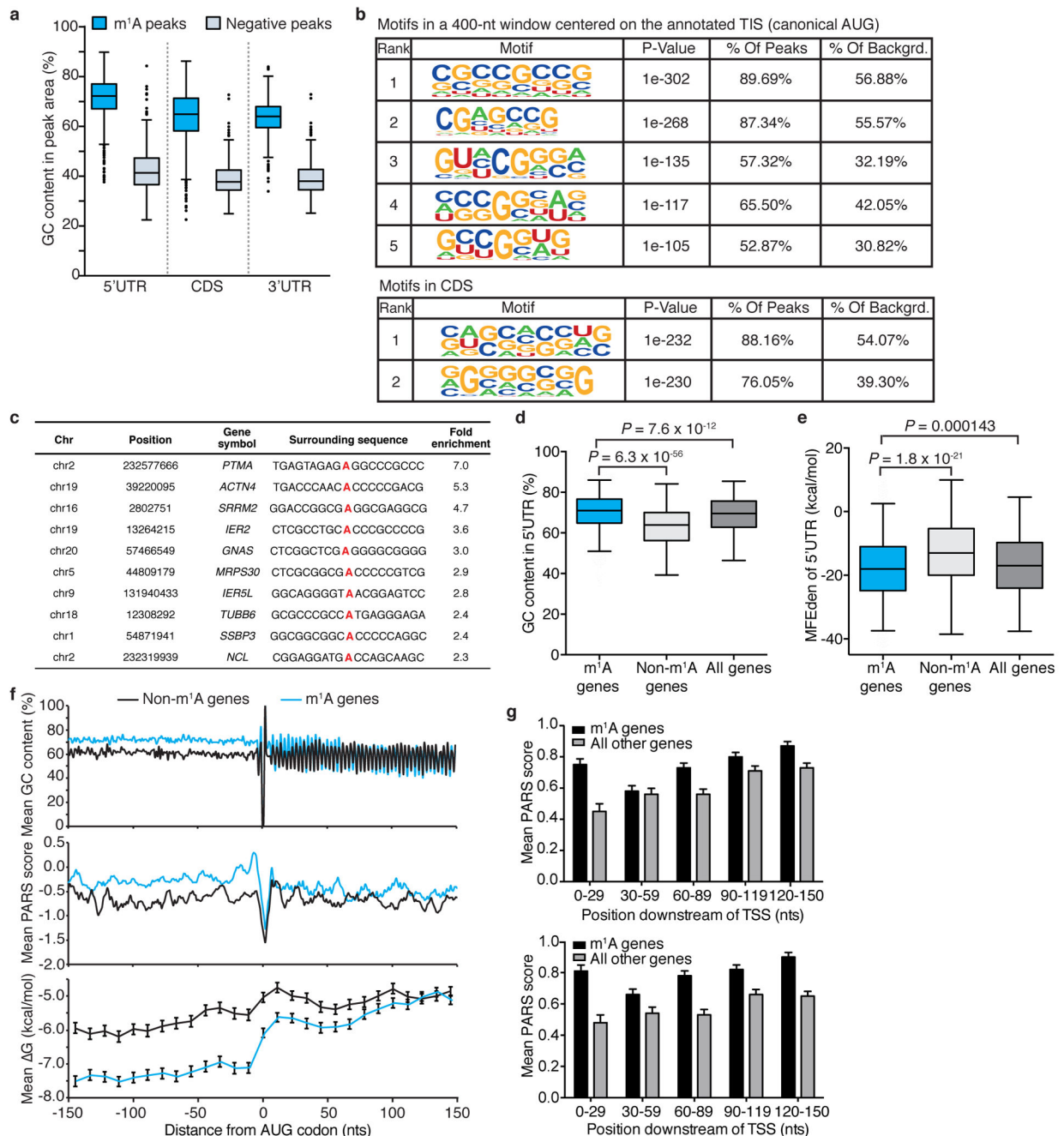
a, Metagenome profiles demonstrating sequence coverage along a normalized gene transcript. Sequence reads of m¹A immunoprecipitation and input in HeLa cells are indicated in blue and orange, respectively. **b–d**, Metagenome profiles of m¹A peak distribution in a non-normalized window centred on the AUG start codon (**b**), extending downstream from the

transcription start site (TSS) (**c**) and centred on the stop codon (**d**), in the indicated human cell types. **e**, Metagene profiles comparing the distribution of m¹A peaks (blue) to that of negative peaks (black) along a normalized transcript composed of three rescaled non-overlapping segments illustrated below, in HeLa cells. **f**, Table demonstrating m¹A peak enrichment in growing windows centred on the AUG start codon in HeLa cells. Enrichment is calculated as number of peaks in the window divided by window size (nucleotides). **g**, Table summarizing the overlap between m⁶A and m¹A peaks in HepG2, HEK293 and WT mESCs. m⁶A peaks are sourced from Dominissini *et al.*¹, Linder *et al.*⁶⁷ and Geula *et al.*⁴², respectively. m¹A peaks are from the current study. **h, i**, Metagene profiles of m¹A peak distribution in a non-normalized window centred on the AUG codon (**h**) and extending downstream from the TSS (**i**). Peaks are sorted by the exon containing the AUG codon and the length of the first exon, respectively. **j**, Metagene profiles of m¹A peak distribution in a non-normalized window centered on the nearest splice site. Peaks are sorted by the exon containing the AUG codon in that gene. **k**, m¹A-induced reverse transcription (RT) arrests produce typical m¹A peaks characterized by a central region of reduced coverage with a local minimum (m¹A trough) Examples are shown. **l**, m¹A-to-m⁶A rearrangement results in a reduced number of identified m¹A troughs. Higher rates of rearrangement further reduce the number of identified m¹A troughs (right panel). Example is shown (left panel). **m**, Metagene profile of m¹A trough distribution along a normalized transcript composed of three rescaled non-overlapping segments illustrated below, in HepG2 cells. **n**, Metagene profile of m¹A trough distribution in a non-normalized window centred on the AUG start codon in HepG2 cells. **o**, Metagene profile of m¹A trough distribution along a normalized transcript in HepG2 cells. **p**, Metagene profile of m¹A trough distribution in a non-normalized window centred on the nearest splice site in HepG2 cells. **q**, Pie chart presenting the fraction of m¹A troughs in each of three non-overlapping transcript segments (5' UTR, CDS and 3'UTR) in HepG2 cells. **r**, MEME motifs identified in 100-nucleotide windows centred on m¹A troughs that lie within the AUG start codon window (± 150 nucleotides) in HepG2 cells.



Extended Data Figure 4. Correlation between m¹A sites and TISs in human

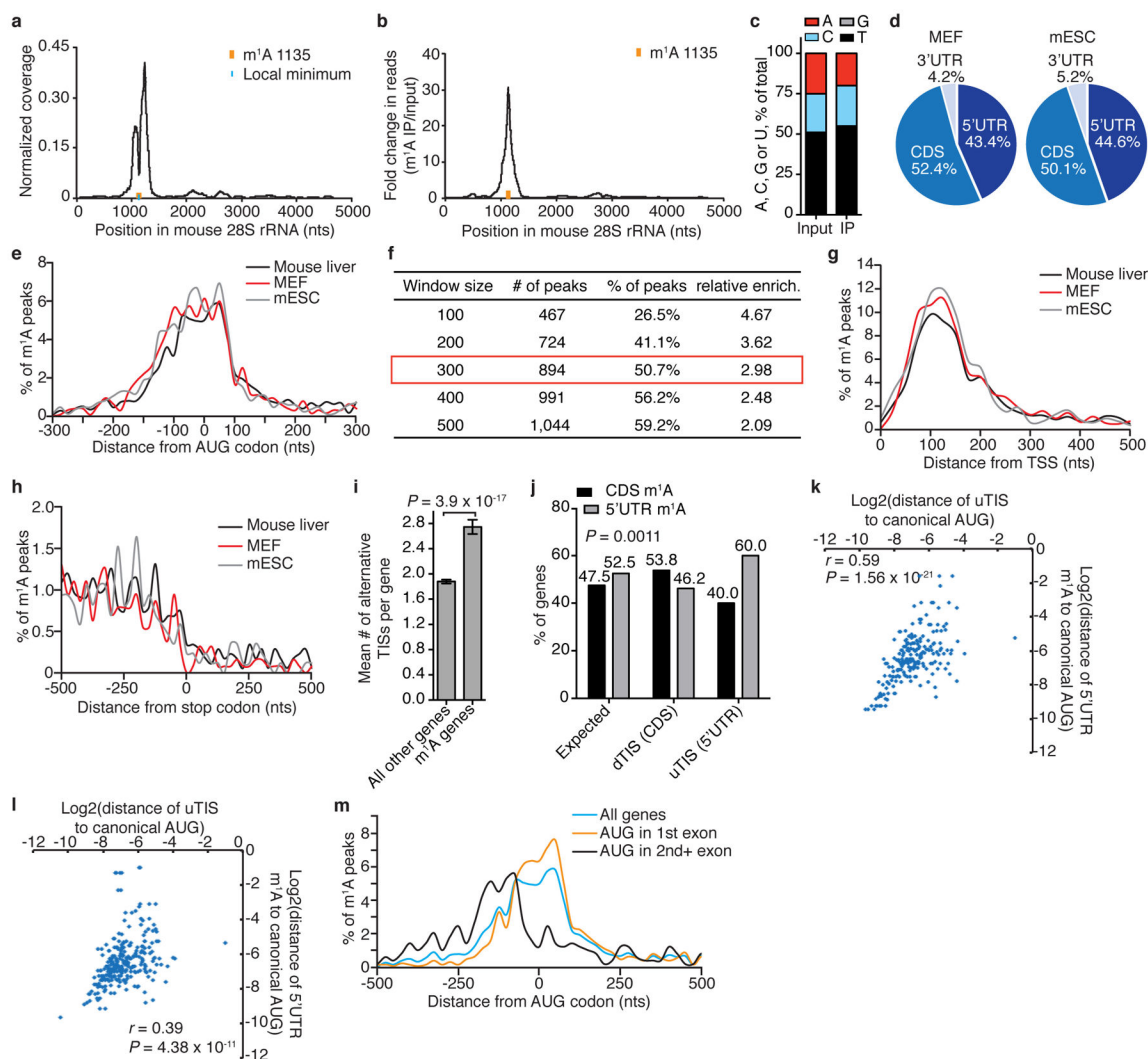
a, b, The mean number of alternative TISs in methylated (m¹A) versus all other (unmethylated) genes in HeLa (**a**) and HEK293 (**b**) cells. Mean values \pm s.e.m. are shown. P values are indicated, Mann–Whitney U test. **c**, The mean number of alternative TISs per gene as a function of the number of m¹A peaks per gene in HeLa and HEK293 cells. Mean values \pm s.e.m. are shown; r and P values are indicated, Pearson correlation; regression line is drawn. **d, e**, The percentage of genes with upstream (=5' UTR) or downstream (=CDS) m¹A sites out of all genes that have either downstream or upstream TISs, compared to the expected percentage in HeLa (**d**) and HEK293 (**e**) cells. P values are indicated, χ^2 test. **f–h**, Scatter plots showing the correlation between the locations (log₂) of alternative TISs and m¹A peaks with respect to the canonical AUG start codon (0) in HeLa (**f**), HEK293 (**g**) and HepG2 (**h**) cells: left, upstream TISs (uTIS) and 5' UTRs m¹A peaks; right, downstream TISs (dTIS) and CDS m¹A peaks. r and P values are indicated, t -test.



Extended Data Figure 5. Structure and sequence features of m¹A sites in human mRNA

a, m¹A peaks have a significantly higher GC content compared to negative peaks in all three transcript segments: 5' UTR, CDS and 3' UTR. Box limits represent 25th percentile, median and 75th percentile, whiskers represent 2.5 and 97.5 percentiles, and dots indicate outliers. $P = 1.5 \times 10^{-278}$, $P = 8.2 \times 10^{-259}$ and $P = 3.3 \times 10^{-271}$, respectively, t -test. **b**, Motifs identified in 400-nucleotide windows centred on the canonical AUG start codon in genes with m¹A peaks in this window (upper table), or around m¹A peaks located in the CDS, outside the AUG start codon window (lower table). **c**, Examples of adenosines around m¹A

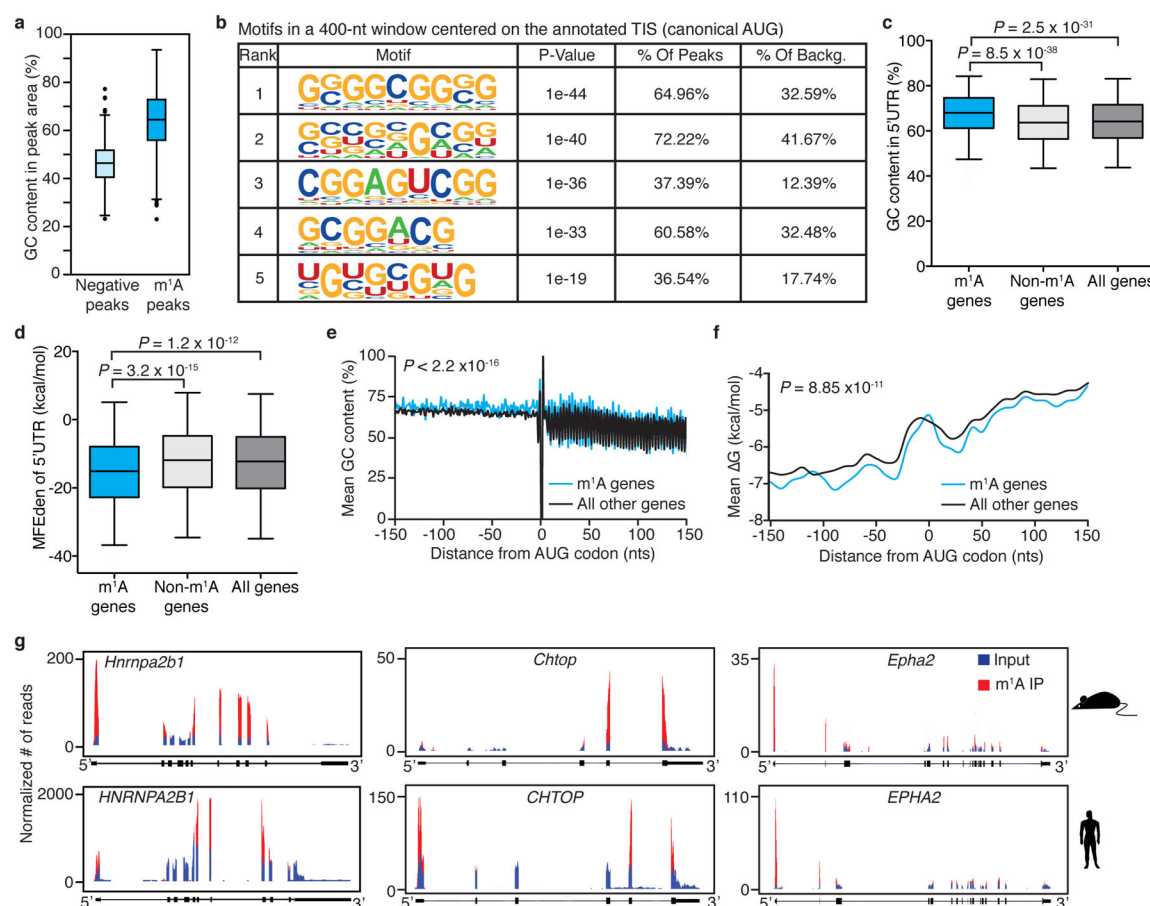
peak middles with increased mismatch rates. Fold-enrichment values are the ratios of mismatch rates in untreated relative to rearranged samples. The top ten highest fold enrichment samples are shown. **d, e**, GC content (**d**) and minimum free energy density (MFEden) (**e**) of 5' UTRs of methylated (m¹A), unmethylated (Non-m¹A) and all genes. Box limits represent 25th percentile, median and 75th percentile, whiskers represent 2.5 and 97.5 percentiles. *P* values are indicated, *t*-test. **f**, Mean GC content (upper panel), PARS score (middle panel) and free energy ($-G$, lower panel) in a 300-nucleotide window centred on the start codon of commonly methylated genes relative to non-methylated genes (see Methods). Error bars represent s.e.m. **g**, The PARS scores of methylated compared to all other genes in HepG2 cells in a 150-nucleotide window extending downstream from the TSS, calculated in 30-nucleotide sliding windows. Each plot represents data from an independent PARS experiment. Error bars represent s.d.



Extended Data Figure 6. Features of the mouse m¹A methylome (part 1)

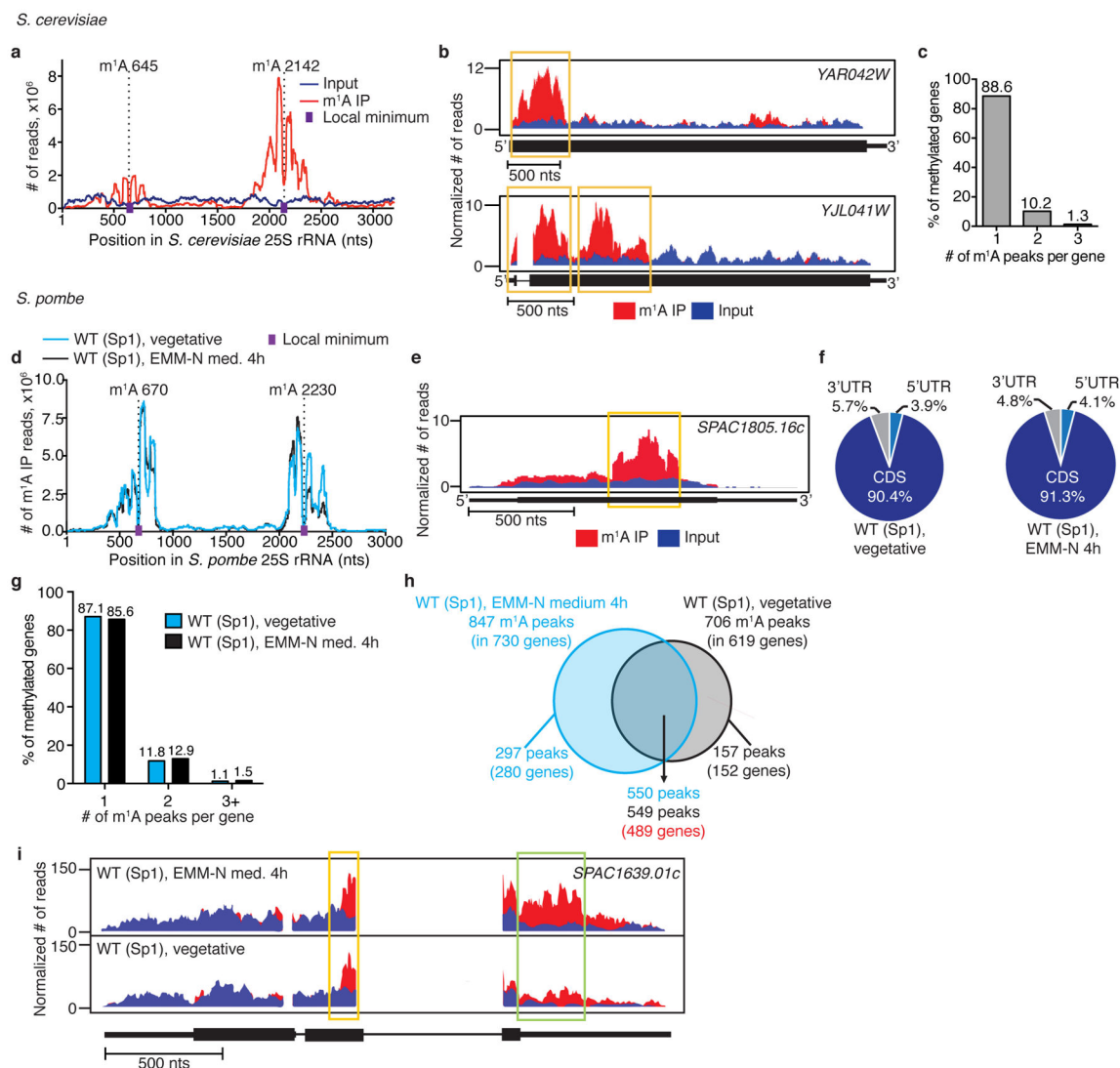
a, Detection of an m¹A site at position 1135 of mouse 28S rRNA, in mouse liver m¹A immunoprecipitation. A drop in sequence read coverage can be seen at the methylated

position. **b**, Fold-enrichment (immunoprecipitation over input reads) identifies an m¹A peak. **c**, High mismatch rate at the identified m¹A 1135 in mouse 28S rRNA. **d**, Pie charts presenting the fraction of m¹A peaks in each of three non-overlapping transcript segments (5' UTR, CDS and 3' UTR) in the indicated mouse cell types. **e**, Metagene profiles of m¹A peak distribution in a non-normalized window centred on the AUG start codon in the indicated mouse cell types. **f**, Table showing m¹A peak enrichment in growing windows centred on the AUG start codon in mouse liver. Enrichment is calculated as the number of peaks in the window divided by window size (nucleotides). **g**, **h**, Metagene profiles of m¹A peak distribution in a non-normalized window extending downstream from the TSS (**g**) and centred on the stop codon (**h**) in the indicated mouse cell types. **i**, The mean number of alternative TISs in methylated (m¹A) versus all other (unmethylated) genes in MEF cells. Mean values \pm s.e.m. are shown; *P* value is indicated, Mann–Whitney *U* test. **j**, The percentage of genes with upstream (=5' UTR) and downstream (=CDS) m¹A sites out of all genes that have either downstream or upstream alternative TISs, compared to the expected percentage in MEF cells. *P* value is indicated, χ^2 test. **k**, **l**, Scatter plots showing the correlation between the locations (\log_2) of upstream TISs (uTIS) and 5' UTR m¹A peaks relative to the canonical AUG start codon (0) in mouse liver (**k**) and MEF (**l**) cells. *r* and *P* values are indicated, *t*-test. **m**, Metagene profiles of mouse m¹A peak distribution in a non-normalized window centred on the AUG codon. Peaks are sorted by the exon containing the AUG codon in that gene.



Extended Data Figure 7. Features of the mouse m¹A methylome (part 2)

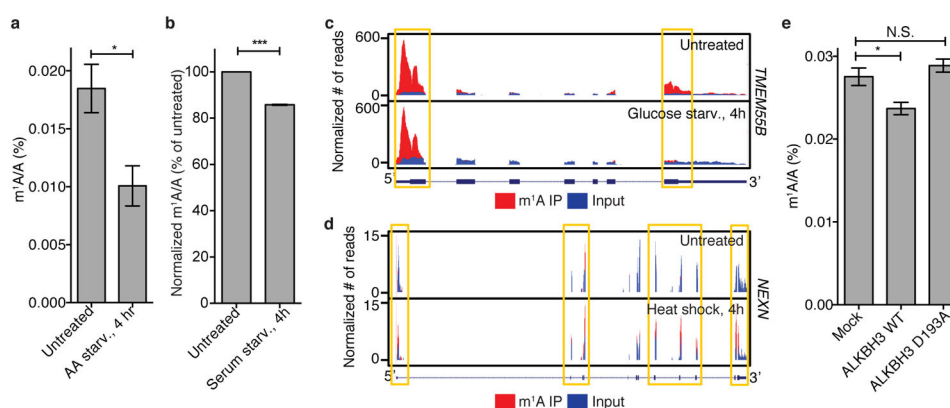
a, Mouse m¹A peaks have a significantly higher GC content compared to negative peaks. Box limits represent 25th percentile, median and 75th percentile, whiskers represent 2.5 and 97.5 percentiles, and dots indicate outliers. $P = 4.4 \times 10^{-175}$, t -test. **b**, Motifs identified in 400-nucleotide windows centred on the canonical AUG start codon in genes with m¹A peaks in this window in mouse liver. **c**, **d**, GC content (**c**) and MFEden (**d**) of 5' UTRs of methylated (m¹A), unmethylated (Non-m¹A) and all genes. Box limits represent 25th percentile, median and 75th percentile, whiskers represent 2.5 and 97.5 percentiles. P values are indicated, t -test. **e**, **f**, A sliding window profile of mean GC content (**e**) and mean ΔG (**f**) in a 300-nucleotide window centred on the canonical AUG start codon in methylated (m¹A) genes compared to all other genes in mouse liver. P values are indicated, Kolmogorov–Smirnov test and t -test. **g**, Representative plots of human-mouse orthologous genes with conserved m¹A peaks. Plot format as in Fig. 2a.



Extended Data Figure 8. The yeast m¹A methylome

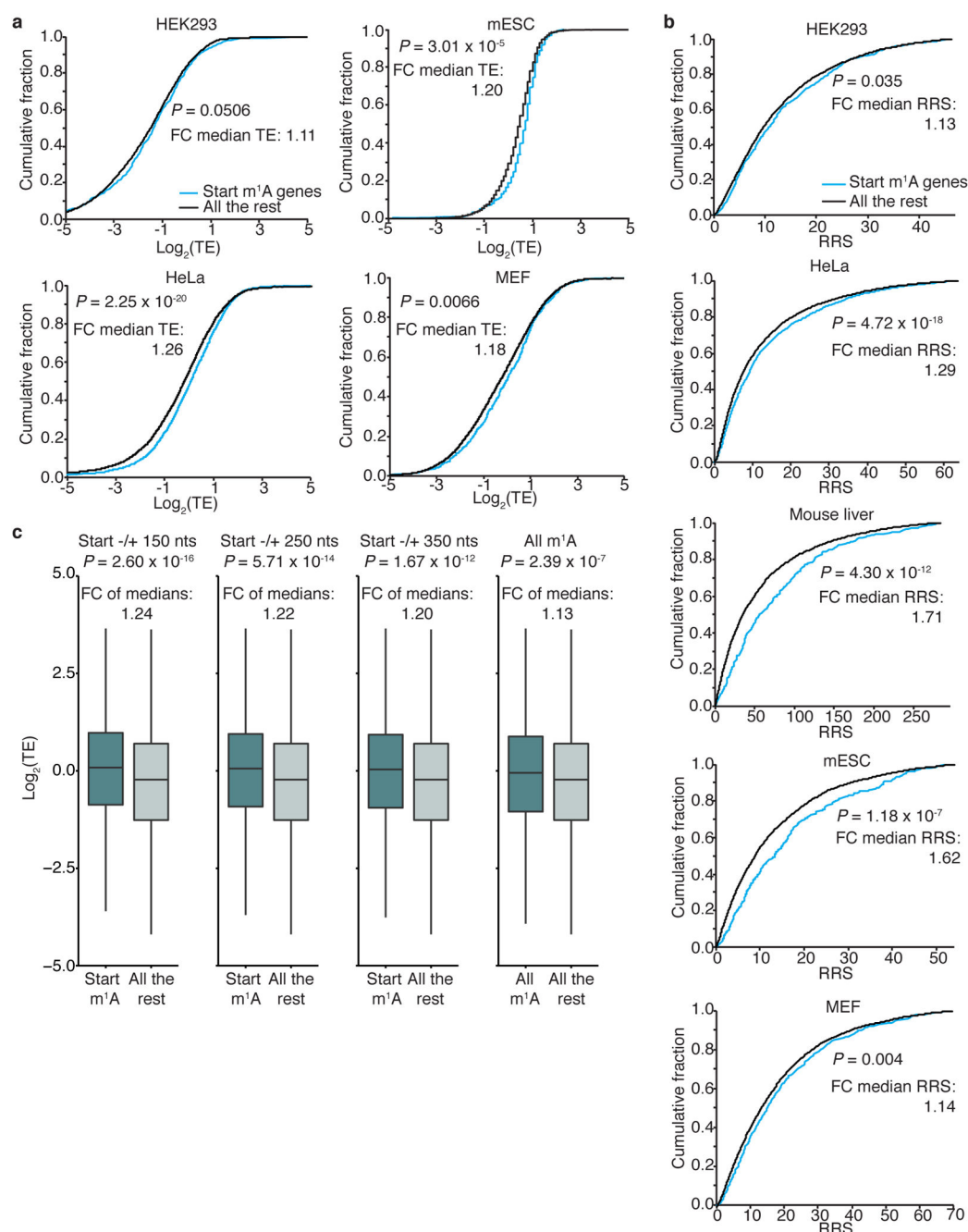
a, m¹A-seq identifies the known m¹A sites (645 and 2142) in *S. cerevisiae* 25S rRNA. A drop in sequence read coverage (indicated by purple dots) occurs around the methylated positions (indicated by dashed lines). **b**, Representative plots of methylated transcripts in *S. cerevisiae*. Plot format as in Fig. 2a. **c**, The percentage of methylated genes that carry 1, 2 or 3 m¹A peaks per gene in *S. cerevisiae*. Out of 843 m¹A peaks (FC 4, FDR 5%) in 778 genes, most (88.6%) are methylated only once. Unlike in mammals, m¹A is distributed across the coding transcriptome without an apparent preferred location. **d**, m¹A-seq identified the known m¹A sites (670 and 2230) in *S. pombe* 25S rRNA. A drop in sequence read coverage (indicated by purple dots) occurs around the methylated positions (indicated by dashed lines). **e**, Representative plot of a methylated transcript in *S. pombe*. Plot format as in Fig. 2a. **f**, Pie charts presenting the fraction of m¹A peaks in each of three non-overlapping transcript segments (5' UTR, CDS and 3' UTR) in *S. pombe* Sp1 strain under the indicated conditions. Under vegetative growth, we identified 706 m¹A peaks (FC 4, FDR 5%) in 619 gene transcripts, most of which (90.4%) distributed along the CDS. Four

hours after transfer to a nitrogen-source deficient ‘sporulation’ medium, 157 out of the vegetative state m¹A peaks were no longer detected, and 297 new peaks appeared. Importantly, transcripts that harbour differential peaks were adequately expressed (above the 1st quartile) in both conditions. **g**, The percentage of methylated genes that carry 1, 2, or 3 +m¹A peaks per gene in *S. pombe* Sp1 strain under the indicated conditions. **h**, Venn diagram representing differential and shared m¹A peaks in *S. pombe* Sp1 strain under the indicated conditions. **i**, Representative plots of a differentially methylated transcript in *S. pombe* Sp1 strain under the indicated conditions. Yellow box, conserved peak; green box, differential peak.



Extended Data Figure 9. m¹A in mRNA is a dynamic modification

a, b, LC-MS/MS quantification of m¹A in mRNA of untreated and amino acid (AA)-starved (**a**) or serum-starved (**b**) HepG2. Mean values \pm s.e.m. are shown, $n = 3$, * $P < 0.05$, *** $P < 0.001$, unpaired t -test. **c, d**, Representative plots of differentially methylated transcripts in untreated and glucose-starved (**c**) or heat shock-treated (**d**) HepG2 cells. Plot formats as in Fig. 2a. **e**, LC-MS/MS quantification of m¹A in mRNA of 293F cells overexpressing WT FLAG-ALKBH3 or an inactive mutant (D193A), presented as percentage of unmodified A. Mean values \pm s.e.m. are shown, $n = 3$, * $P < 0.05$, NS, $P > 0.05$, one-way ANOVA with Dunnett's multiple comparison test.



Extended Data Figure 10. m¹A around the start codon correlates with higher translation efficiency (TE)

a, Cumulative distribution of $\log_2(\text{TE})$ in genes methylated in a 300-nucleotide window centred on the start codon compared to all other genes, in the indicated human and mouse cell types. P values (t -test) and fold-changes (FC) of median TE values (Start m¹A genes/All the rest) are indicated. **b**, Genes methylated in a 300-nucleotide window centred on the AUG start codon have a higher ribosome release score (RRS = $\text{TE}[\text{CDS}]/\text{TE}[3' \text{ UTR}]$) compared to all other genes in the indicated cell types. RRSs, which are 'normalized' to ribosomal drop-off in the 3' UTR, are in line with TE scores. P values (Mann-Whitney U test) and

fold-changes (FC) of median RRS values (Start m¹A genes/All the rest) are indicated. **c**, Genes methylated in different start codon window sizes have higher TE compared to all other genes, in HeLa cells. When considering all m¹A genes, including those methylated outside the start codon window, the effect is reduced. *P* values (*t*-test) and fold-changes (FC) of median TE values (Start m¹A genes/All the rest) are indicated. Box limits represent 25th percentile, median and 75th percentile; whiskers extend from the limit to the highest and lowest value within 1.5 IQR (interquartile range).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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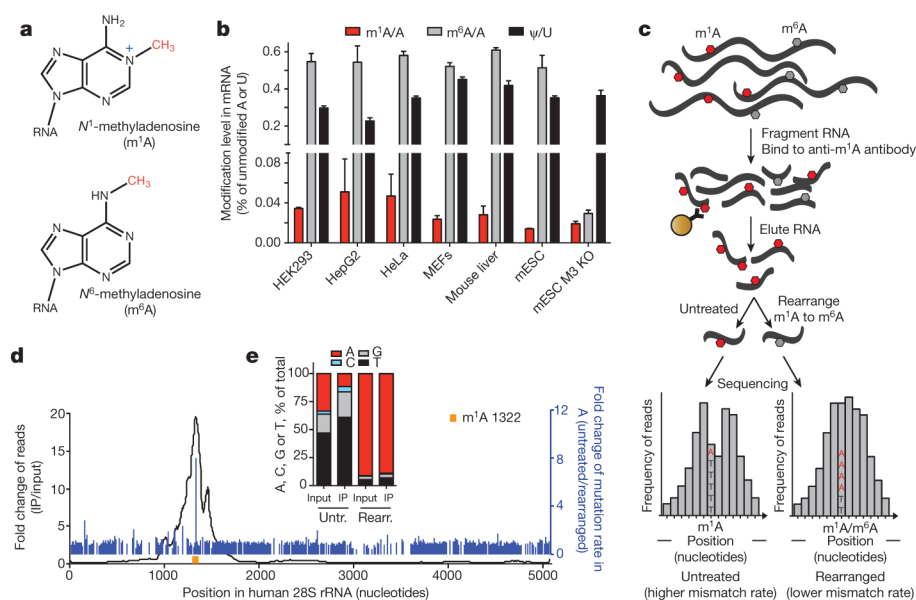


Figure 1. Development of m^1A -seq to map a newly identified constituent of mammalian mRNA
a, Chemical structures of m^1A and m^6A is in blue. **b**, LC-MS/MS quantitation of m^1A , m^6A and Ψ in human and mouse mRNA isolated from the indicated cell types. The level of each modified nucleoside is indicated as a percentage of the unmodified one. Mean values \pm s.e.m. are shown, $n = 3$. MEFs, mouse embryonic fibroblasts; mESC, mouse embryonic stem cells; M3 KO, *Mettl3* knockout. **c**, Schematic outline of m^1A -seq. RNA is fragmented and subjected to immunoprecipitation using anti- m^1A antibody. Eluted RNA fragments are converted to cDNA and sequenced, or treated to induce partial m^1A -to- m^6A rearrangement before cDNA synthesis. m^1A causes both reverse transcription stops and read-throughs accompanied by mismatches, to produce typical peaks with a central trough and an adenosine with increased mismatch rate (left). Partial rearrangement of m^1A to m^6A attenuates the effect (right). m^1A is in red, m^6A is in grey; mismatch rate is illustrated as the ratio between A (red) and T (black). **d**, Detection of the known m^1A 1322 site (yellow) in human 28S rRNA validates m^1A -seq (black curve). High fold-change in mismatch rate (before and after m^1A -to- m^6A rearrangement in immunoprecipitation, blue lines) independently identifies m^1A 1322 with single-nucleotide resolution. **e**, Mismatch rates in m^1A 1322 increase after immunoprecipitation and decrease upon rearrangement.

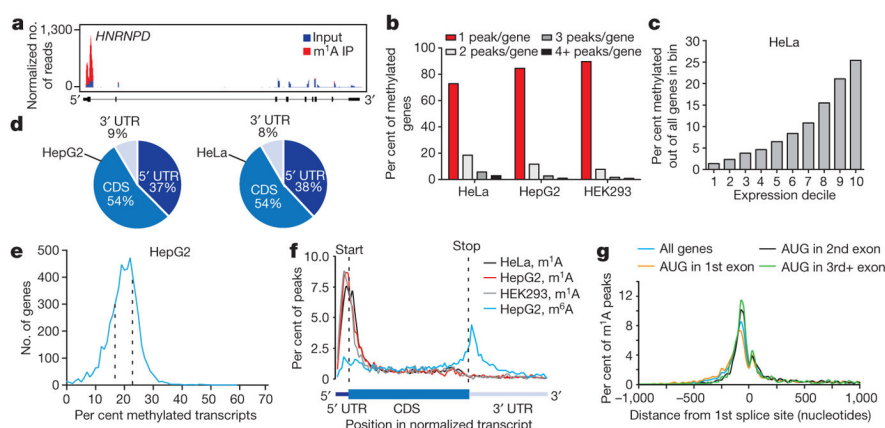


Figure 2. m¹A is associated with translation initiation sites (TISs) in the human transcriptome
a, Representative plot of a methylated transcript. Normalized sequence coverage of immunoprecipitation (m¹A IP, red) and input (blue) are indicated above the gene architecture in UCSC format. Thin black boxes represent the 5' and 3' untranslated regions (UTRs), thick black boxes represent the coding sequences (CDS) and thin lines represent introns. **b**, The percentage of methylated genes that carry 1, 2, 3 or 4+ peaks per gene in the indicated human cell types. Most methylated genes carry only one m¹A peak. **c**, The percentage of methylated genes in HeLa cells increases with expression level. **d**, Pie charts presenting the fraction of m¹A peaks in each of three non-overlapping transcript segments (5' UTR, CDS and 3' UTR) in the indicated human cell types. **e**, Stoichiometry (fraction) of methylated transcripts in genes that carry one m¹A peak in HepG2 mRNA. Plotted is the number of genes sorted by percentage of methylated transcripts. Dashed lines define the stoichiometry interquartile range (17–23%). **f**, Metagene profiles of m¹A peak distribution along a normalized transcript composed of three rescaled non-overlapping segments illustrated below, in the indicated human cell types. m¹A peaks cluster around the AUG start codon. For comparison, the distribution of m⁶A peaks in HepG2 is superimposed. **g**, Metagene profiles of m¹A peak distribution in a non-normalized window centred on the first splice site. Peaks are sorted by the exon containing the AUG codon in that gene.

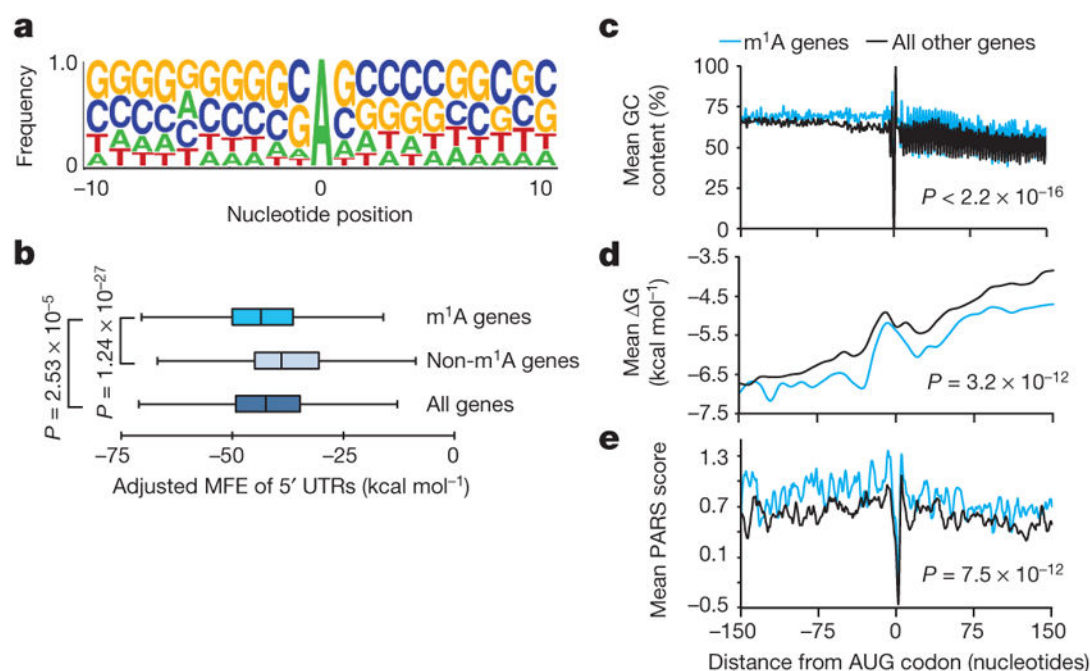


Figure 3. m¹A occurs in GC-rich sequence contexts and in genes with structured 5' UTRs

a, Sequence frequency logo for a set of 192 adenosines in peak areas that have a higher mismatch rate in immunoprecipitation relative to input (FC = 6) in HepG2 demonstrates the GC-rich context of m¹A. **b**, Length-adjusted minimum free energy (aMFE) for 5' UTRs of methylated, unmethylated and all genes. Box limits represent 25th percentile, median and 75th percentile, whiskers represent 2.5 and 97.5 percentiles. P-values are indicated, *t*-test. Analysis was based on human common peaks (see Methods). **c–e**, A sliding window of mean GC content (**c**), ΔG (**d**) and PARS scores (**e**) in a ± 150 -nucleotide window centred on the AUG start codon of methylated genes compared to all the rest, in HepG2 cells. P-values are indicated; Kolmogorov–Smirnov test, *t*-test and Kolmogorov–Smirnov test, respectively.

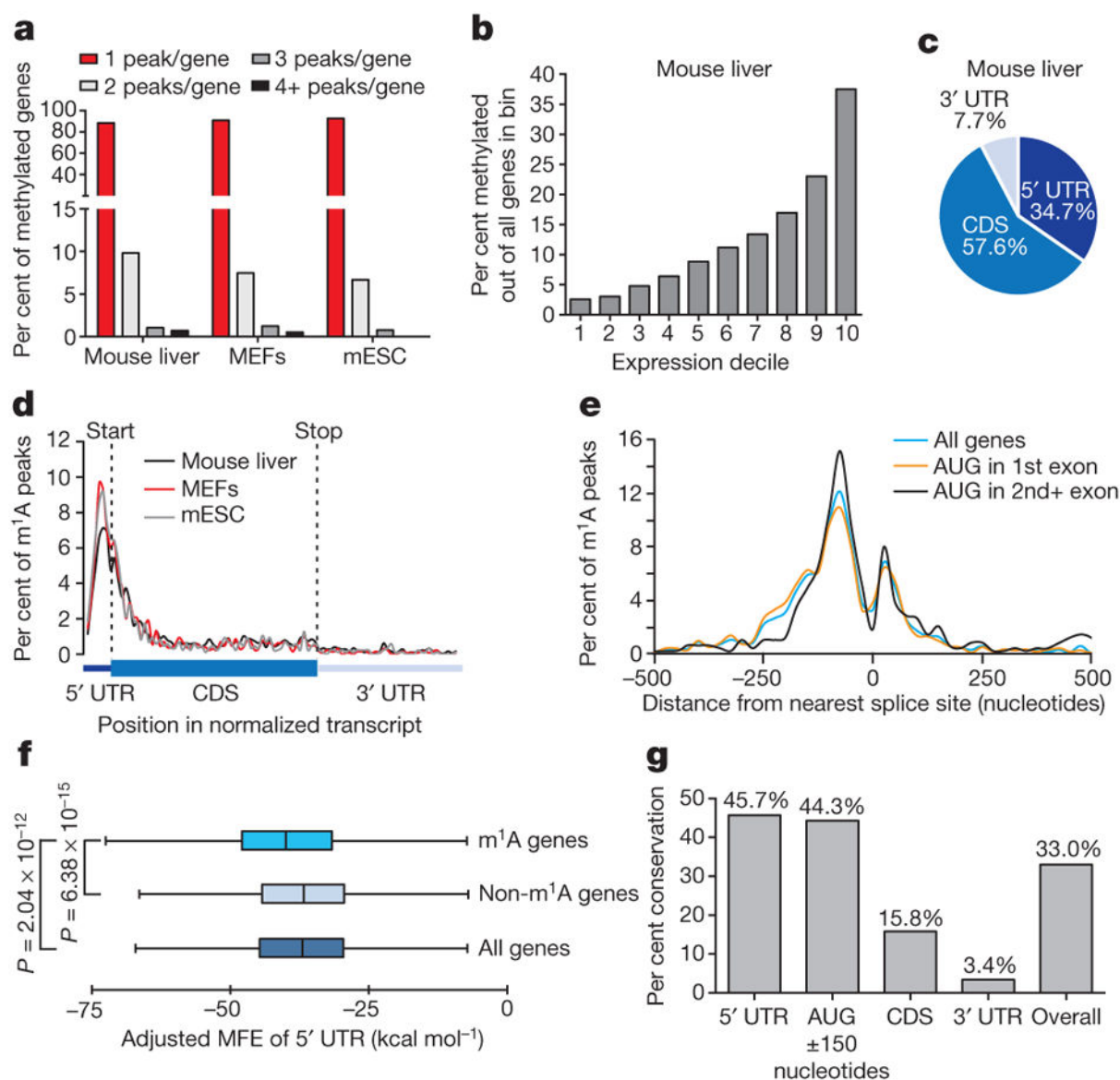


Figure 4. m¹A methylome conservation between human and mouse

a, The percentage of methylated genes that carry 1, 2, 3 or 4+ peaks per gene in the indicated mouse cell types. Most genes carry only one m¹A peak. **b**, The percentage of methylated genes in mouse liver cells increases with expression level. **c**, Pie chart presenting the fraction of m¹A peaks in each of three non-overlapping transcript segments (5' UTR, CDS and 3' UTR) in mouse liver. **d**, Metagenes profiles of m¹A peak distribution along a normalized transcript composed of three rescaled non-overlapping segments illustrated below, in the indicated mouse cell types. m¹A peaks cluster around the AUG start codon. **e**, Metagenes profiles of mouse m¹A peak distribution in a non-normalized window centred on the nearest splice site. Peaks are sorted by the exon containing the AUG codon in that gene. **f**, Length-adjusted minimum free energy (aMFE) for 5' UTRs of methylated, unmethylated and all genes in mouse liver. Box limits and whiskers are as indicated in Fig. 3b. *P* values are

indicated, *t*-test. **g**, Human–mouse m¹A conservation expressed as per cent of orthologous positions with shared m¹A peaks according to their location in the transcript.

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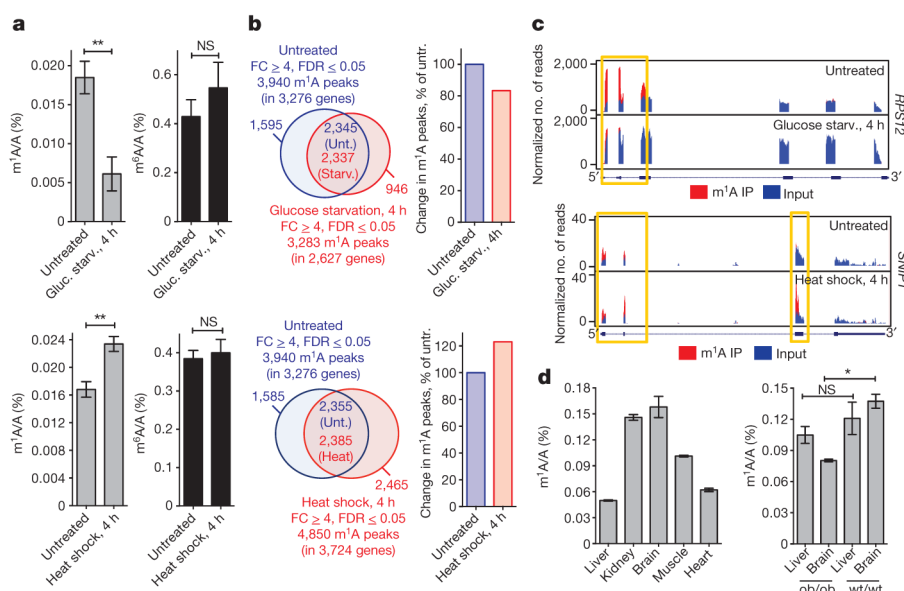


Figure 5. m¹A in mRNA is a dynamic modification that responds to changing physiological and stress conditions, and varies between tissues

a, LC-MS/MS quantification of m¹A (left, grey) and m⁶A (right, black) in mRNA of untreated and glucose-starved (upper panels) or heat shock-treated (lower panels) HepG2 cells, presented as percentage of unmodified A. Mean values ± s.e.m. are shown, *n* = 3, ***P* < 0.01, not significant (NS) *P* > 0.05, unpaired *t*-test. **b**, Venn diagrams of differential and shared m¹A peaks in untreated and glucose-starved (left upper panel) or heat shock-treated (left lower panel) HepG2 cells. The overall change in the number of identified m¹A peaks is presented as per cent of untreated (right upper and lower panels). **c**, Representative plots of differentially methylated transcripts in untreated and glucose starved (upper panel) or heat-shock-treated (lower panel) HepG2 cells. Plot format as in Fig. 2a. Yellow boxes frame differential peaks. **d**, LC-MS/MS quantification of m¹A in the indicated wt/wt and ob/ob mouse tissues, presented as percentage of unmodified A. Mean values ± s.e.m. are shown, *n* = 3, **P* < 0.05, NS, *P* > 0.05, one-way ANOVA.

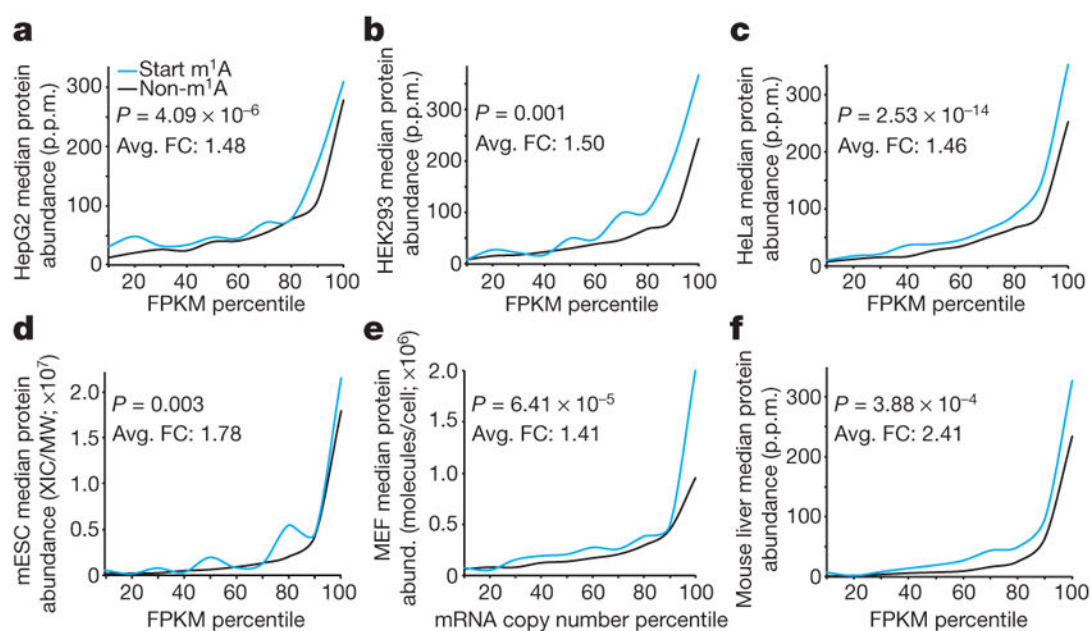


Figure 6. m¹A around the start codon correlates with higher protein levels

a–f, Genes methylated in a 300-nucleotide window centred on the start codon have higher protein levels than all other genes with similar RNA expression levels in HepG2 (**a**), HEK293 (**b**), HeLa (**c**), mESC (**d**), MEFs (**e**) and mouse liver (**f**). ANOVA m¹A *P* values against log₂ protein levels and the average fold-change (Avg. FC) of median protein levels across all gene expression bins (Start m¹A/Non-m¹A) are indicated. FPKM, fragments per kilobase of transcript per million mapped reads.