N6-methyladenosine-dependent RNA structural switches regulate RNA–protein interactions

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RNA-binding proteins control many aspects of cellular biology through binding single-stranded RNA binding motifs (RBMs)¹⁻³. However, RBMs can be buried within their local RNA structures⁴⁻⁷, thus inhibiting RNA–protein interactions. N6-methyladenosine (m6A), the most abundant and dynamic internal modification in eukaryotic messenger RNA⁸⁻¹⁹, can be selectively recognized by the YTHDF2 binding protein responsible for pre-mRNA processing²⁰⁻²⁴. Combining nuclear ribonucleoprotein C (HNRNPC), an abundant nuclear RNA-non-coding RNA (lncRNA) to facilitate binding of heterogeneous nuclear ribonucleoprotein C (HNRNPC), an abundant nuclear RNA-binding protein responsible for pre-mRNA processing²⁰⁻²⁴. Combining photoactivatable-ribonucleoside-enhanced crosslinking and immunoprecipitation (PAR-CLIP) and anti-m6A immunoprecipitation (MeRIP) approaches enabled us to identify 39,060 m6A-switches among HNRNPC-binding sites; and global m6A reduction decreased HNRNPC binding at 2,798 high-confidence m6A-switches. We determined that these m6A-switch-regulated HNRNPC-binding activities affect the abundance as well as alternative splicing of target mRNAs, demonstrating the regulatory role of m6A-switches on gene expression and RNA maturation. Our results illustrate how RNA-binding proteins gain regulated access to their RBMs through m6A-dependent RNA structural remodelling, and provide a new direction for investigating RNA-modification-coded cellular biology.

Post-transcriptional m6A RNA modification is indispensable for cell viability and development, yet its functional mechanisms are still poorly understood⁸⁻¹⁷. We recently identified one m6A site in a hairpin-stem...
on the human lncRNA metastasis-associated lung adenocarcinoma transcript (MALAT1) \(^{23}\) (Extended Data Fig. 1a). A native gel-shift assay indicated that this m\(^A\) residue increases the interaction of the RNA hairpin with proteins in the HeLa nuclear extract (Fig. 1a). RNA pull-down assays identified HNRNPC as the protein component of the nuclear extract that binds more strongly with the m\(^A\)-modified hairpin than the unmodified hairpin (Fig. 1b and Extended Data Fig. 1b, c). The m\(^A\)-enhanced interaction with the hairpins was validated qualitatively by ultraviolet crosslinking and quantitatively (\(\sim8\)-fold increase) by filter binding using recombinant HNRNPC1 protein (Fig. 1c and Extended Data Fig. 1d).

The HNRNPC protein belongs to the large family of ubiquitously expressed heterogeneous nuclear ribonucleoproteins that bind nascent RNA transcripts to affect pre-mRNA stability, splicing, export and translation\(^{20\text{–}24}\). HNRNPC preferably binds single-stranded U-tracts (five or more contiguous uridines)\(^{20\text{–}24,26\text{–}27}\). In the MALAT1 hairpin, HNRNPC binds a U\(_5\)-tract that is half buried in the hairpin-stem opposing the 2,577–A/m\(^A\) site (Extended Data Fig. 1a, e).

Since m\(^A\) residues within RNA stems can destabilize the thermostability of model RNA duplexes\(^{24}\), we hypothesized that the 2,577–m\(^A\) residue increases the interaction of the RNA hairpin-stem to make its opposing AC motif (Fig. 1d). The m\(^A\) residue that pairs with 2,577–A/m\(^A\) (Extended Data Fig. 1f, g), increased the accessibility of the U-tract and enhanced HNRNPC binding (Extended Data Fig. 1f, g).

This hypothesis. First, according to the RNA structural probing assays, the m\(^A\)-modified hairpin showed significantly increased nuclease S1 digestion (single-strand specific) at the G\(_5\) (A = m\(^A\)) motif, as well as markedly decreased RNase V1 digestion (double-strand stacking specific) at the U-tract opposing the G\(_5\) motif (Fig. 1d). The m\(^A\) residue marked destabilized the stacking properties of the region centred around the U residue that pairs with 2,577–A/m\(^A\) (Extended Data Fig. 1f, g), which was also supported by the increased reactivity between CMCT and the U-tract bases in the presence of m\(^A\) (Extended Data Fig. 1h).

Second, the 2,577–A-to-U mutation increased the HNRNPC pull-down amount from the nuclear extract, whereas U-to-C mutations in the U-tract significantly reduced the HNRNPC pull-down amount regardless of m\(^A\) modification (Fig. 1e). Third, the 2,577–A-to-U mutation increased the accessibility of the U-tract and enhanced HNRNPC binding by \(\sim4\)-fold (Extended Data Fig. 2a–c). Binding results with four other mutated A/m\(^A\) oligonucleotides also supported the U-tract, with increased accessibility alone being sufficient to enhance HNRNPC binding (Extended Data Fig. 2d).

Fourth, RNA terminal truncation followed by HNRNPC binding identified two pairs of truncated hairpins with highly accessible U-tracts, which improved HNRNPC binding significantly but independent of the m\(^A\) modification (Extended Data Fig. 2e–j). All these results confirmed that m\(^A\) modification can alter its local RNA structure and enhance the accessibility of its base-paired

**Figure 2** PAR-CLIP–MeRIP identifies m\(^A\)-switches transcriptome wide. a, CLIP-2dTLC showing the m\(^A\) enrichment in HNRNPC-bound RNA regions. Data are mean \(\pm\) s.d.; \(n = 3\), biological replicates. IP, immunoprecipitation. b, HNRNPC-bound RNA regions had higher anti-m\(^A\) pull-down yield than polyA\(^\pm\) RNA. Data are mean \(\pm\) s.d.; \(n = 3\), biological replicates. c, Illustration of the PAR-CLIP–MeRIP protocol. UV, ultraviolet. d, PAR-CLIP–MeRIP identified an m\(^A\) residue around the MALAT1 2,577 site. e, Binding motifs identified with PAR-CLIP–MeRIP peaks. f, Density plot showing the positive enrichment at RRACH sites. g, h, Validation of two m\(^A\)-switches by S1/V1 structural probing and filter binding. Data are mean \(\pm\) s.d.; \(n = 4\), technical replicates. Annotation is the same as in Fig. 1c, d.
residues or nearby regions to modulate protein binding (Fig. 1f). We term this mechanism that regulates RNA–protein interactions through m^6^A-dependent RNA structural remodelling ‘the m^6^A-switch’.

We performed two experiments to determine the global effect of m^6^A-switches on HNRNPC binding. First, in vivo crosslinking followed by immunoprecipitation and two-dimensional thin-layer chromatography (CLIP-2D TLC) showed that the m^6^A:A ratio of the HNRNPC-bound RNA regions had a ~6-fold higher m^6^A level than the HNRNPC-bound intact RNA, and a ~3-fold higher m^6^A level than the flow-through RNA (Fig. 2a and Extended Data Fig. 3a). Second, the HNRNPC-bound RNA regions had much higher anti-m^6^A pull-down yield (4.3%) than the polya^1^ RNA samples (0.5%) using the previously established m^6^A antibody^1^,^1^ (Fig. 2b). These results indicate a widespread presence of m^6^A residues in the vicinity of HNRNPC-binding sites.

To map the m^6^A sites around HNRNPC-binding sites, we performed PAR-CLIP to isolate all HNRNPC-bound RNA regions (input control sample) followed by MeRIP to enrich m^6^A-containing HNRNPC-bound RNA regions (IP sample). Both the input control and IP samples from two biological replicates were sent for RNA sequencing (RNA-seq) (Fig. 2c and Extended Data Fig. 3b, c). This approach, termed PAR-CLIP–MeRIP, identified transcriptome-wide the m^6^A-proximal HNRNPC-binding site, such as the enriched peak around the MALAT1 2,577 site (Fig. 2d). Remarkably, HNRNPC PAR-CLIP–MeRIP peaks harboured two consensus motifs, the HNRNPC RBM (U-tracts) and the m^6^A consensus motif GRACH (a subset of RRACH^1^,^1^) (Fig. 2e). Both motifs were located mostly within 50 residues, suggesting transcriptome-wide RRACH–U-tract coupling events within the HNRNPC-binding sites (Extended Data Fig. 4a, b). About 62% of all RRACH–U-tract coupling events within HNRNPC-binding sites are enriched at the RRACH motif (Fig. 2f). Our PAR-CLIP–MeRIP approach identified a total of 39,060 HNRNPC m^6^A-switches that corresponded to m^6^A-modified RRACH–U-tract coupling events at a false discovery rate (FDR) ≤ 5% (Extended Data Fig. 4c). These switches account for ~7% of 592,477 HNRNPC-binding sites identified by PAR-CLIP. The majority (87%) of m^6^A-switches occur within introns (Extended Data Fig. 4d, e), consistent with the literature that HNRNPC is nuclear localized and primarily binds nascent transcripts^20,^21. We validated two intronic m^6^A-switches in hairpin structures in which m^6^A residues increase the U-tract accessibility and enhance HNRNPC binding by ~3–4 fold (Fig. 2g, h and Extended Data Fig. 5).

To assess the effect of global m^6^A reduction on RNA–HNRNPC interactions, we performed HNRNPC PAR-CLIP experiments in METTL3 and METTL14 knockdown cells (Extended Data Fig. 6a). We identified 16,582 coupling events with decreased U-tract–HNRNPC interactions upon METTL3 and METTL14 knockdown (METTL3/L14 knockdown) (Fig. 3a and Extended Data Fig. 6b, c). In total, 2,798 m^6^A-switches identified by PAR-CLIP–MeRIP experiments showed decreased HNRNPC binding upon METTL3/L14 knockdown (Fig. 3b) and this number is probably an underestimate due to the fact that METTL3/L14 knockdown reduces the global m^6^A level by only ~30–40% (refs 11, 12). These

Figure 3 | Global m^6^A reduction decreases HNRNPC binding at m^6^A-switches.

- **a**: Density plot showing negative enrichment at the U-tracts. KD, knockdown. **b**: Identification of high-confidence (HCS) m^6^A-switches. **c**: Regional distribution of high-confidence m^6^A-switches. CDS, coding sequence. **d**: Density plot showing m^6^A-switch distribution relative to exon/intron boundaries. **e**: m^6^A-switches in coding RNA were enriched in the 3' UTR and near the stop codon. **f**: Cumulative distribution of HCS m^6^A-switches (black) and control (orange) regarding the S1/V1 cleavage preference (data from ref. 4) at U-tracts and RRACH motif. U-tract can be 3' (top) or 5' (bottom) of the RRACH motif. **P < 0.05, ***P < 10^-3.** Kolmogorov–Smirnov test. **g**: Phylogenetic conservation of high-confidence m^6^A-switches among primates and vertebrates. ***P < 10^-10, Mann–Whitney–Wilcoxon test.
High-confidence m^6^A switches are enriched in the introns of coding and non-coding RNAs (Fig. 3c and Extended Data Fig. 6d). Exonic m^6^A switches are enriched at the middle of exons whereas intronic m^6^A switches are slightly enriched near the 5' end (Fig. 3d). m^6^A switches within coding RNAs tend to locate at very long exons (Extended Data Fig. 6e) and are enriched near the stop codon and in the 3' untranslated region (UTR) (Fig. 3e), consistent with the known topology of the human m^6^A methylome in mRNAs^{13,14}. Transcriptome-wide RNA structural mapping on high-confidence m^6^A-switches yielded consistent structural patterns with our three demonstrated m^6^A-switch hairpins (Fig. 3f). The RR residues in the RRACH motif and the 3' U-tract residues show increased structural dynamics in the presence of m^6^A. Besides, m^6^A switches prefer short RRACH–U-tract inter-motif distances, are not involved in the previously reported inter-U-tract motif patterns and are conserved across species (Fig. 3g and Extended Data Fig. 6f–i).

To reveal the function of m^6^A switches on RNA biology, we performed polyA^- RNA-seq from HNRNPC, METTL3 and METTL14 knockdown and control cells (Extended Data Fig. 7a). METTL3/L14 knockdown, which has been shown to decrease HNRNPC binding transcriptome-wide, co-regulated the expression of 5,251 genes with m^6^A hypermethylation (Extended Data Fig. 7b). The m^6^A-switch-regulated expression of genes within these GO categories was validated by quantitative polymerase chain reaction (qPCR) (Fig. 4a and Extended Data Fig. 7c). The m^6^A-switch-regulated expression of genes within these GO categories was validated by quantitative polymerase chain reaction (qPCR) (Fig. 4a and Extended Data Fig. 7d–g). We also found that HNRNPC, METTL3 and METTL14 knockdown decreased the cell proliferation rate to similar extents (Extended Data Fig. 7b).

Besides the mRNA abundance level changes, we also observed splicing pattern changes within high-confidence m^6^A-switch-containing transcripts by testing the differential exon usage in RNA-seq data (DESeq)^{30}. HNRNPC knockdown co-up/downregulated 131/127 exons with METTL3 knockdown and 130/115 exons with METTL14 knockdown. These co-regulated exons occur more frequently in the vicinity of m^6^A switches than non-co-regulated exons (Fig. 4b, c), indicating that m^6^A switches tend to regulate splicing events at nearby exons. We investigated the splicing pattern at two exons with neighbouring m^6^A switches: the PAR-CLIP–MeRIP and METTL3/L14 knockdown data confirmed the HNRNPC-binding signature at the m^6^A switch site neighbouring these exons; and HNRNPC and METTL3/L14 knockdown co-inhibited exon inclusion in both cases (Fig. 4d–f and Extended Data Fig. 8b–f). Besides, we identified 155 genes with multiple m^6^A switches exhibiting more

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Figure 4 | m^6^A switches regulate mRNA abundance and alternative splicing. a, HNRNPC, METTL3/L14 knockdown (KD) co-regulated the abundance of m^6^A-switch-containing transcripts by RNA-seq and qPCR. Ctrl, control. b, Illustration of the relative exon distance to m^6^A-switches. c, Co-regulated exons by HNRNPC knockdown and METTL3 knockdown (left) and METTL14 knockdown (right) were more enriched around m^6^A-switch sites than non-co-regulated exons, Kolmogorov–Smirnov test. HCS, high confidence m^6^A switches. d–f, Validation of the m^6^A-switch-regulated splicing at one exon neighbouring the CDS2 m^6^A switch as shown in PAR-CLIP–MeRIP data (d), METTL3/L14 knockdown data (e), and PCR with reverse transcription (RT–PCR) results (f). The red triangle and square mark the m^6^A site and U-tract, respectively. Data are mean ± s.d.; n = 4, biological replicates.
than two splice variants, and 221 m^6^A-switch-containing genes with differentially expressed splice variants in HNRNPC and METTL3/L14 knockdown samples. Further analysis suggested that m^6^A-switches have an effect on intron excision (Extended Data Fig. 8g). Consistent with previous reports about splicing regulation by both HNRNPC and m^6^A, our results indicate that m^6^A functions as an RNA structure remodeler to affect mRNA maturation through interference with post-transcriptional regulator binding activities.

We demonstrated that post-transcriptional m^6^A modifications could modulate the structure of coding and non-coding RNAs to regulate RNA–HNRNPC interactions, thus influencing gene expression and maturation in the nucleus. It is possible that m^6^A could also recruit additional accessory factors, such as the YTH domain proteins, which can directly recognize m^6^A—as previously reported—to destabilize the RNA structure and facilitate HNRNPC binding. Besides HNRNPC, m^6^A-switches may regulate the function of many other RNA-binding proteins through modulating the RNA-structure-dependent accessibility of their RBMs. Our work indicates widespread m^6^A-induced mRNA and lncRNA structural remodelling that affects RNA–protein interactions for biological regulation.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions N.L., G.Z. and M.P. designed and performed experiments, and analysed data. Q.D. synthesized all RNA oligonucleotides. N.L., M.P. and T.P. conceived the project. N.L. and T.P. wrote the paper with input from C.H. and M.P.

Author Information RNA-seq data have been deposited in the Gene Expression Omnibus under accession number GSE56010. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to M.P. (marc.parisien@mcgill.ca) or T.P. (taopan@uchicago.edu).
METHODS

Mammalian cell culture, siRNA knockdown and western blot. Human cervical cancer cell line HeLa (CCL-2) and embryonic kidney cell line HEK293T (CRL-11268) were obtained from the American Type Culture Collection (ATCC) and were cultured under standard conditions. Control short interfering RNA (siRNA) (1027281, Qiagen), METTL3 siRNA (S014317096, Qiagen), METTL14 siRNA (S014317096, Qiagen) or HNRNPC siRNA (10620318, Invitrogen) were transfected into HEK293T cells at a concentration of 40 nM using lipofectamine RNAiMAX (Invitrogen) according to the manufacturer’s instructions. Cells were collected after 48 h from the transfection, shock-frozen in liquid nitrogen, and stored at −80 °C for further studies. Western blot analysis using METTL3- (HPA038002, Sigma), METTL14- (HPA038002, Sigma), HNRNPC- (sc-32308, Santa Cruz) and GAPDH- (A00192-20, Genescript) specific antibodies was performed under standard procedures. Blotting membranes were stained for Protein Kinase (RN2232, GE Healthcare) and visualized by a digital imaging system (G:BOX, SYNGENE). All synthetic oligonucleotides were synthesized by Q.D.

Gel shift, RNA pull-down and filter-binding assays. HeLa nuclear extracts were isolated using the NE-PER Nuclear and Cytoplasmic Extraction Reagents (78833, Thermo Scientific) according to the manufacturer’s instructions. The purified radioactively labelled RNA oligonucleotides were refolded by heating at 90 °C for 1 min, then at 30 °C for 5 min. Three microtubes HeLa nuclear extract and 6 μl refolded RNA were incubated at room temperature for 30 min and then at 4 °C for 2 h. Each sample was mixed with 1 μl 50% glycerol, separated on an 8% native 1X TBE gel, and visualized by phosphorimaging using the Personal Molecular Imager (Bio-Rad).

The in vitro pull-down assay was performed as described31. The eluted protein samples were separated on 4–12% polyacrylamide Bis-Tris gels (NP0321BOX, Thermo Scientific) and stained with SYPRO Ruby (S12000, Invitrogen) according to the manufacturer’s instructions. Protein in gel slices or the entire pulled-down protein samples were digested with trypsin and identified using Liquid chromatography-tandem mass spectrometry by the Donald Danforth Plant Science Center (Washington University). The RNA oligonucleotides used in Fig. 1f were: 2,577–UC-A, 5′-AACUUAAUUGUUAUUGGCUUCAUGAGA-Biotin; 2,577–CC-A, 5′-AUCUUAAUUGUUAUUGGCUUCAUGAGA-Biotin; 2,577–CC-m6A, 5′-AACUUAAUUGUUAUUGGCUUCAUGAGA-Biotin; 2,577–mC-m6A, 5′-AACUUAAUUGUUAUUGGCUUCAUGAGA-Biotin.

The full-length HNRNPC protein was purified and used in the in vitro ultraviolet cross-linking assay as performed previously as described35. Filtered and eluted RNA samples were performed as described previously41.

CLIP-2dTLC. HEK293T cells at 70–80% confluency were ultraviolet irradiated with 400 mJ cm−2 at 254 nm, and harvested by centrifuging at 4,000 r.p.m. for 3 min at 4 °C (with centrifugation rotor 75003524, Fisher Scientific). The pellet of cross-linked cells was resuspended in 1 ml lysis buffer (1X PBS, 0.1% SDS, 1% Nonidet P-40, 0.5% sodium deoxycholate, protease inhibitor cocktail and RNase inhibitor) and incubated on ice for 4 h. Cell lysate was isolated by centrifuging at 3,000 r.p.m. for 5 min and pre-blocked with 50 μl protein A beads in 300 μl lysis buffer. Another 50 μl protein A beads (Invitrogen) were incubated with 8 μg corresponding antibodies for 4 h at room temperature, and then mixed with the pre-blocked cell lysate at 4 °C overnight. The beads were washed three times with 1 ml wash buffer (20 mM Tris-HCl pH 7.4, 10 mM EDTA, 1% SDS) containing 2 mg ml−1 T4 PNK, 600 Ci mmol−1 γ-[32P] ATP, 5X PKN buffer) and incubation at 37 °C for 30 min. Unreacted [γ-32P] ATP was removed using Illustra MicroSpin G-25 columns. The eluted RNA was digested with 1 μl of 1 μl−1 nuclease P1 at 37 °C for 1 h. Samples were spotted on cellulose TLC plate and 2dTLC was run as described24 using isobutyric acid: 0.5 M NaH2O (5.3, v/v) as the first dimension and isopropanol:HCl:water (70:15:15, v/v/v) as the second dimension.

RNA structural probing and RNA terminal truncation. The synthetic RNA oligonucleotides were 5′-end-labeled with [γ-32P]ATP by T4 PNK (70031, Affymetrix), gel purified, and re-folded. Structural probing assay with RNAse T1, RNAse S1 and RNAse V1 was performed as previously described30. Note that 3′-end-labeled HNRNPH1 oligonucleotides were used for the RNA structural probing assay shown in Fig. 2g.

CMCT RNA structural probing assay was performed as reported31. RNA refolding: 3 pmol RNA was annealed in 50 mM potassium borate (pH 8) by heating at 90 °C for 1.5 min then incubation at room temperature for 3 min.

RNA terminal truncation assay was carried out as previously reported31. RNA samples were first alkaline-hydrolysed as in the RNA structural probing assay, and then incubated with HNRNPC protein in the same conditions as in the filter binding assay. The RNA–protein complexes were then loaded onto filter papers and washed twice with chilled binding buffer. Air-dry filters and RNA samples were then extracted from the filters and loaded onto denaturing gel as in the RNA structural probing assay.

PAR-CLIP and PAR-CLIP–MeRIP. PAR-CLIP procedures were performed as previously reported32 with the following modification. HEK293T cells in 15-cm plates treated following normal plates for PAR-CLIP procedures were ligated and digested with a combination of RNase I (Ambion, AM2295, 15 μl/l 50 μl diluted with H2O) and Turbo DNases (2 μl) for 3 min at 37 °C, shaking at 1,100 r.p.m. The lysate was then immediately cleared by spinning at 14,000 r.p.m. for 30 min, and placed on ice for further use. HNRNPC-binding sites were identified by PARalyzer v.1.1 (ref. 33) with default settings.

PAR-CLIP–MeRIP experiment applied m6A-antibody immunoprecipitation13,24 to the HNRNPC PAR-CLIP RNA samples. The HNRNPC PAR-CLIP RNA sample was incubated with m6A-specific antibody (202003, SYSY), RNase inhibitor (80 units, Sigma-Aldrich), human placental RNase inhibitor (NEB) in 200 μl 1× IP buffer (50 mM Tris-HCl pH 7.4, 750 mM NaCl and 0.5% (v/v) Igepal CA-630) at 4 °C for 2 h under gentle shaking conditions. For each PAR-CLIP–MeRIP experiment, 20 μl protein A beads (Invitrogen) were washed twice with 1 ml 1× IP buffer, blocked with 2 h incubation with 100 μl 1× IP buffer supplemented with bovine serum albumin (BSA) (0.5 mg ml−1), RNasin and human placental RNase inhibitor, and then washed twice with 100 μl 1× IP buffer. The pre-blocked protein A beads were then combined with the prepared immuno-reaction mixture and incubated at 4 °C for 2 h, followed by three washes with 100 μl 1× IP buffer. After that, the RNA was eluted by 1 h incubation with 20 μl elution buffer (1× IP buffer and 6.7 mM m6A, Sigma-Aldrich) under gentle shaking conditions, and purified by ethanolic precipitation. The purified RNA sample (IP) as well as the input PAR-CLIP RNA sample (input control) were used for library construction by Truseq small RNA sample preparation kit (illumina).

Libraries were prepared using TruSeq Small RNA Sample Preparation Kit (RS-2000-0122, Illumina) according to the manufacturer’s instructions, and then sequenced by Illumina HiSeq2000 with single-end 50-bp read length. The control and IP samples from PAR-CLIP–MeRIP experiments (same case for the control and knockdown samples from METTL knockdown experiments) were sequenced together in one flowcell on two lanes, and the reads from two lanes of each sample were combined for remaining analysis. The raw sequencing data were trimmed using the Trimmomatic computer program v.0.30 (ref. 35) to remove adaptor sequences, and mapped to the human genome version hg19 by Bowtie 1.0.0 (ref. 36) without any gaps and allowed for at most two mismatches.

Detection of PAR-CLIP–MeRIP peaks and differential PAR-CLIP peaks. The raw read counts of the biological replicates confirmed the reproducibility between replicates (Extended Data Fig. 9), and replicates were combined for subsequent analysis. For the genomic regions in the reads, we calculated the average read counts within an 11-nucleotide window centred at that site, as the normalized read counts for that site. This normalization smoothed the raw mapping curves, and facilitated identification of peaks within each mapping cluster. To correct for changes in sequencing depth or expression levels between samples, we then normalized the read counts at each genomic site to the total number of read counts on the respective gene. The above defined double-normalization procedures enabled precise identification of changes in the mapping reads at specific genomic locations by directly comparing the normalized read counts between samples. No read counts in the intergenic region were compared between samples, because the transcription boundaries are not defined and the intergenic read counts cannot be normalized to correct for transcription expression.

Detection of PAR-CLIP–MeRIP peaks involves comparing the read counts of the IP sample with that of the control (Ctrl) sample as follows: (1) we identified all peaks within HNRNPC-binding sites in the IP sample; (2) we performed transcriptome-wide scanning to compare read counts of each identified peak in (1) with read counts at the same genomic locations in the Ctrl sample to calculate the fold change score, score = log2 (H1/H0). The score threshold was set to be 1, corresponding to a twofold increase compared with control. The detection of decreased HNRNPC-binding sites involved comparing HNRNPC knockdown (KD) sample with that in the control as follows: (1) we identified all peaks within HNRNPC-binding sites in the METTL knockdown sample; (2) we performed transcriptome-wide scanning to compare read counts of each identified peak in (1) with read counts at the same genomic locations in control to calculate the fold change score, score = log2 (H0/H1).

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Identification of enriched motifs and HNRNPC m6A-switches. To identify enriched motifs, we first sorted the 12,998 HNRNPC PAR-CLIP–MeRIP peaks (with IP/input enrichment \( \geq 2 \)) by the T-o-C mutation frequency. We then chose the top 4,500 peaks with the highest T-o-C mutation frequency for motif analysis using FIRE \(^*\) with default RNA analysis parameters. The top two enriched motifs are the GRACH and the U-tract motif. We also used the top 1,024 and 2,048 peaks for motif analysis, yielding the same motif results as the top 4,500 peaks.

To identify transcriptome-wide HNRNPC m\(^6\)A-switches, we first searched for all coupling events within 50 nucleotides between U5 and RRACH motifs, with the binding protein, the RNA-seq data from METTL3/L14 and reverse transcribed using SuperScript III First-Strand Synthesis System (Life Technologies, catalogue no. 18080-051). In order to validate the splicing changes identified from our RNA-seq data, we performed RT–PCR measurements using Thermo Scientific Taq DNA Polymerase under the following conditions: 95 °C for 3 min, 30 cycles of 95 °C for 30 s, 55 °C for 30 s, 72 °C for 1 min, and then finally 72 °C for 10 min. For the target alternate exon, we designed and used primers annealing to both neighbouring constitutive exons. The PCR products were separated on 1.2% agarose gel and ethidium bromide stained. In order to validate the gene expression level changes identified from our RNA-seq data, we performed qRT–PCR measurements using SYBR Green PCR Master Mix (Life Technologies, catalogue no. A6020) under the following conditions: 50 °C for 3 min followed by 95 °C for 10 min, 40 cycles of 95 °C for 15 s, 60 °C for 1 min, and then 40 °C for 1 min and 95 °C for 15 s and finally 60 °C for 30 s.

The primer sequences are as follows (listed as gene name: forward primer; reverse primer).

| Gene       | Forward Primer | Reverse Primer |
|------------|----------------|----------------|
| METTL3     | 5' TTGGGGTGTTTCGATTGTTT 3' | 5' CCACTCAGGCTTGAAGCT 3' |
| METTL14    | 5' CCACTCAGGCTTGAAGCT 3'   | 5' CCACTCAGGCTTGAAGCT 3' |

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Extended Data Figure 1 | m^6A increases the accessibility of the U-tract to enhance HNRNPC binding. a, Secondary structure of the MALAT1 hairpin with m^6A methylation at the 2,577 site shown in red^25. Nucleotide position numbers correspond to their locations along the human MALAT1 transcript (NCBI accession NR_002819). b, RNA pull down showing that HNRNPC preferably binds methylated RNA. c, The list of proteins with identified peptides by mass spectrometry in b. d, Recombinant HNRNPC1 binds more strongly with that MALAT1 2,577–m^6A hairpin compared with the unmethylated hairpin, as determined by an in vitro ultraviolet crosslinking assay^23. e, HNRNPC shows binding around the 2,577–A site along MALAT1 in vivo, as determined by previously published HNRNPC iCLIP data^20. The underlying genomic sequence is shown at the bottom with a red square marking the 2,577–m^6A site. The slight shift of the iCLIP signal to upstream of the U-tract-binding site is probably due to the steric hindrance of the peptide fragment remaining on RNA, which can cause reverse transcription to terminate more than one nucleotide upstream of the crosslink site^20. f, Quantification of the RNase V1 cleavage signal for the U-tract region from the RNA structural mapping assay in Fig. 1e. To correct for sample loading difference, each band signal was normalized to the band signal of the immediate 3' residue to the U-tract. Data are mean ± s.d.; n = 3, technical replicates. g, Quantitative analysis of the RNase T1 cleavage signal from the RNA structural mapping assay in Fig. 1e. An increased RNase T1 cleavage signal (single-strand specific and cleavage after guanosines) was observed due to the surrounding m^6A residue. To correct for sample loading difference, the ratio for each band signal among all bands in each lane was calculated. Relative T1 cleavage = (m^6A native/m^6A denatured)/(A native/A denatured). n = 2, technical replicates. h, Quantitative CMCT mapping showing increased signals for the U-tract bases around the U base-pairing with m^6A. Quantitation of band signals within the U-tract region is shown on the right. Data are mean ± s.d.; n = 4, technical replicates.
Extended Data Figure 2 | Increased accessibility of U-tracts enhances HNRNPC binding. a, Structure probing of the 2,577–A-to-U mutated MALAT1 hairpin (2,577–U). The annotation is the same as in Fig. 1d. b, Quantification of the RNase V1 cleavage signal for the U-tract region from RNA structural mapping assays as in a. To correct for sample loading difference, each band signal was normalized to the band signal of the 39-most U of the U-tract. n = 2, technical replicates. c, Filter-binding curves displaying the binding affinities between recombinant HNRNPC1 and 2,577–U/A oligonucleotides. Data are mean ± s.d.; n = 3, technical replicates. d, Filter-binding results showing the binding affinities between recombinant HNRNPC1 and four mutated MALAT1 oligonucleotides. (1) Mutate G–C to C–C, 2,577–A: predicted to weaken the hairpin stem and increase HNRNPC binding. Results: binding improved from 722 nM Kd to 142 nM (fivefold). (2) Mutate G–C to C–C, 2,577–m6A: in this context of weaker stem, m6A is predicted to confer a smaller effect compared to wild-type hairpin. Result: improved binding only twofold instead of eightfold. (3) Restore C–C to C–G, 2,577–A: predicted to restore the hairpin stem and decrease HNRNPC binding compared to C–C mutant. Result: binding decreased by 6.4-fold. (4) Restore C–C to C–G, 2,577–m6A: in this context of restored stem, m6A is again predicted to confer a smaller effect compared to wild-type hairpin. Result: improved binding by 2.5-fold. Data are mean ± s.d.; n = 3 each, technical replicates. e, RNA alkaline hydrolysis terminal truncation assay showing recombinant HNRNPC1 binding to terminal truncated MALAT1 hairpin oligonucleotides (2,577 site m6A methylated or unmethylated). In this assay, 39-radiolabelled MALAT1 2,577 hairpin oligonucleotides were terminal truncated by alkaline hydrolysis into RNA fragments that were then incubated with HNRNPC1 protein followed by filter binding wash steps. The remaining RNA on the filter paper was isolated and analysed by denaturing gel electrophoresis, as indicated in the lane ‘C1-bound or C1-B’. ‘Input’ refers to alkaline-hydrolysis-truncated RNA oligonucleotides used for incubation with hnRNP C1; ‘G-L or G-ladder’ was generated from RNase T1 digestion; ‘Ctrl’ refers to the intact MALAT1 hairpin without alkaline hydrolysis truncation. One pair of methylated/unmethylated terminal truncated oligonucleotides (CUT1, marked by green arrows) was selected for subsequent biochemical analysis, due to their strong interaction with HNRNPC. f, RNA terminal truncation assay as in e except 59 32P-labelled oligonucleotides were used. One pair of methylated/unmethylated terminal truncated oligonucleotides (CUT2, marked by green arrows) was selected for subsequent biochemical analysis. g, Structure probing of the CUT1 oligonucleotides using RNase V1 and nuclease S1 digestion. Annotation is the same as in g, i. Structure probing of the CUT2 oligonucleotides using RNase V1 and nuclease S1 digestion. Annotation is the same as in g, i. Truncated oligonucleotides with exposed U-tracts increased HNRNPC binding regardless of m6A. Data are mean ± s.d.; n = 3, technical replicates.
Extended Data Figure 3 | m^6A is enriched in the vicinity of HNRNPC-binding sites. a, Schematic diagram of the CLIP-2dTLC protocol. IP, immunoprecipitation; nt, nucleotide; UV, ultraviolet. The RNase T1 used in our 2dTLC assay cleaves single-stranded RNA after guanosines, so the m^6A/A ratio determined here represents the m^6A fraction of all adenosines following guanosines. b, Analysis of crosslinked RNA–HNRNPC complexes (CLIP RNP) using denaturing gel electrophoresis (lanes 1 and 2). Positions of the protein size standards are shown on the left. HNRNPC IP RNA region (RNA samples within RNA–HNRNPC crosslinked complexes) were extracted from the gel slices marked by the red rectangle. c, Denaturing gel analysing the size distribution for the HNRNPC PAR-CLIP RNA samples (lane 2). The RNA size standards were loaded in lanes 1 and 3.
Extended Data Figure 4 | PAR-CLIP–MeRIP identifies transcriptome-wide m^6A-switches in the vicinity of HNRNPC-binding sites. a, Density plots illustrating the distribution of distance between the PAR-CLIP–MeRIP/input peaks and the nearest GRACH motif (top) or the nearest U-tracts (bottom). b, Definition and identification of HNRNPC m^6A-switches based on the PAR-CLIP–MeRIP analysis. Approximately 89% of PAR-CLIP–MeRIP peaks harbouring both the U-tract and RRACH motifs have an RRACH–U-tract inter-motif distance within 50 nucleotides, significantly higher than the 64% of such coupling within the genomes. HNRNPC m^6A-switches are identified as m^6A-methylated RRACH–U-tract coupling events. c, Volcano plot depicting all coupling events (open circles) as defined in b, according to their P values^36 (y axis) and fold-change values at RRACH sites (x axis). To identify HNRNPC m^6A-switches, we generated the π value, π = E (−log_{10} P), as one comprehensive parameter to pick meaningful genomic loci^37. HNRNPC m^6A-switches identified from PAR-CLIP–MeRIP experiments should fulfill the following requirements: (1) read counts at both the control and IP sample ≥ 5; (2) π value ≥ 0.627, corresponding to FDR ≤ 5%. d, Pie chart depicting the region distribution of HNRNPC m^6A-switches identified by PAR-CLIP–MeRIP. e, Pie chart depicting HNRNPC PAR-CLIP peaks. These are enriched in introns, consistent with previous reports that HNRNPC binds mainly nascent transcripts^19,23,25.
Extended Data Figure 5 | Validation of two identified m^A-switches.

a, b, PAR-CLIP–MeRIP data detected positive IP/input enrichment at the RRACH sites (red arrowheads) on the DNAJC25-GNG10 gene (a) and HNRNPH1 gene (b) in HEK293T cells. c, d, Quantification of RNase V1 cleavage signals around the U-tract region of m^A-switches on the DNAJC25-GNG10 (c) and HNRNPH1 (d) transcript, related to Fig. 2g, h. Data are mean ± s.d.; n = 3, technical replicates each. e, Quantitative CMCT mapping of DNAJC25-GNG10 m^A-switch shows increased band signals around the uridine base that pairs with m^A. The red vertical line marks the U-tract region. Quantitation of band signal for the U-tract region is shown on the right. Data are mean ± s.d.; n = 3, technical replicates. The HNRNPH1 m^A-switch hairpin is not suitable for CMCT probing, because its reverse transcription binding primer region is too short. f, g, In vivo DMS mapping of the DNAJC25-GNG10 hairpin (f) and HNRNPH1 (g); data are from ref. 7. A and C residues are marked with orange dots and the m^A residue is marked with a red dot. The hairpin loops are indicated by red bars. h, Transcriptome-wide S1/V1 mapping around the HNRNPH1 m^A-switch site. Blue bars represent V1 signal; magenta bars represent S1 signal. The hairpin loop is indicated by a red bar; data are from ref. 4. Not enough reads could be collected to make a plot for the DNAJC25-GNG10 m^A-switch region.
Extended Data Figure 6 | Molecular features of high-confidence m^6A-switches. 

**a**, Western blot (WB) showing stable HNRNPC protein abundance upon METTL3/L14 knockdown. 

**b**, Volcano plot of the METTL3/L14 knockdown (KD) data depicting RRACH–U-tract coupling events (open red circles) as defined in Extended Data Fig. 4b, according to their P values\(^3\) (P; y axis) and fold-change values at the U-tracts (E; x axis). 

**c**, Overlap of RRACH–U-tract coupling events with decreased HNRNPC binding by METTL3 and METTL14 knockdown. 

**d**, The intron fraction of HCS m^6A-switches in coding RNA and non-coding RNA. 

**e**, Density plot displaying the distribution of exonic m^6A-switches/HNRNPC PAR-CLIP peaks according to exon length. 

**f**, Inter-motif (RRACH–U-tract) distance distributions suggest that m^6A-switches have a preference for shorter distances between the RRACH and U-tract (>5×U) motifs. The distribution curves are from PAR-CLIP–MeRIP data (green), METTL3/L14 knockdown (red) and high-confidence (HCS) m^6A-switches (black). 

**g**, Analysis of the inter-motif (U-tract–U-tract) distance patterns, previously identified by iCLIP\(^20\), in PAR-CLIP–MeRIP, METTL3/L14 knockdown and high-confidence m^6A-switch data. The peaks at \(\sim 165\) and \(\sim 300\) nucleotides are clearly present. For the 2,798 high-confidence switches, we analysed those in which the other U-tract motif is also in a PAR-CLIP-identified sequence; the long-range peaks seem to have shifted to longer distances (\(\sim 220\) and \(\sim 370\) nucleotides). 

**h**, METTL3/L14 knockdown does not affect the inter-motif (U-tract–U-tract) distance distributions for U-tracts (\(\sim 5\times U\)) in HEK293T cells. 

**i**, EVOfold analysis for the 2,798 high-confidence m^6A-switches. The chances for high-confidence m^6A-switches to have EVOfold records are significantly higher than random genomic sequences. We first calculated the number of high-confidence sites in the EVO database if occurring in random to be \(1.7\). We found that 18 high-confidence sites are present in the EVO database, resulting in \(11.60\) enrichment. This result is further divided into intronic and exonic regions.
Extended Data Figure 7 | m^6A-switches regulate the abundance of target mRNAs. a, HNRNPC, METTL3/L14 knockdown confirmed by western blots. b, HNRNPC knockdown (KD) and METTL3/L14 knockdown co-regulated the expression of a large number of genes. Gene expression changes between control (Ctrl) and HNRNPC, HNRNPU, METTL3/L14 knockdown HEK293T cells were analysed by Cuffdiff2 (refs 38, 39), and the absolute numbers of differentially expressed genes are shown. HCS-containing genes refers to the 1,815 genes containing high-confidence m^6A-switches. The RNA-seq data from HNRNPU knockdown HEK293T cells (Gene Expression Omnibus accession GEO34995 data set40) were analysed for comparison with a different mRNA-binding protein. HNRNPU did not show preferential interaction with the 2,577–m^6A modified MALAT1 hairpin (Fig. 1b, c). c, GO analysis of the m^6A-switch-containing genes whose expression levels were co-differentially regulated by HNRNPC and METTL3/L14 knockdown, against all m^6A-switch-containing genes as background. d, An example of an m^6A-switch among co-regulated transcripts is the ARHGAP5 transcript (NCBI accession NM_001030055). Its proposed secondary structure with the m^6A methylation site in red is shown with the opposing the U-tract in a stem. e, f, PAR-CLIP–MeRIP detected positive IP/input enrichment at the RRACH site (red arrowhead) of the ARHGAP5 m^6A-switch (e), while METTL3/L14 knockdown decreased HNRNPC binding at the U-tract (red square) of this m^6A-switch (f). g, The expression level of the ARHGAP5 gene was co-upregulated by HNRNPC, METTL3/L14 knockdown, as shown by the RNA-seq data from HEK293T cells. The vertical black line represents the m^6A-switch site. h, HNRNPC, METTL3/L14 knockdown decreased the proliferation rates of HEK293T cells to a similar extent. Data are mean ± s.d.; n = 4, biological replicates.
Extended Data Figure 8 | m\(^{\text{6}A}\)-switches regulate alternative splicing of target mRNAs. 

**a**, Fold changes (knockdown (KD)/control (Ctrl), \log_{2}) in normalized exon expression against RNA-seq reads detect the exons in HNRNPC knockdown, METTL3 knockdown, METTL14 knockdown and control samples. Statistically significant differentially expressed exons (SSDEEs) called by DEXSeq are indicated in red. 

**b**, Proposed secondary structure of the CDS2 hairpin with the m\(^{\text{6}A}\) methylation site shown in red, opposing the U-tract region. Nucleotide position numbers correspond to their locations along the human CDS2 transcript (NCBI accession NM_003818).

**c**, Proposed secondary structure of the YTHDF2 hairpin with the m\(^{\text{6}A}\) methylation site shown in red, opposing the U-tract region. Nucleotide position numbers correspond to their locations along the human YTHDF2 transcript (NM_001173128).

**d**, **e**, PAR-CLIP–MeRIP detected a positive enrichment at the RRACH site (red arrowhead) (d), while METTL3/L14 knockdown decreased HNRNPC binding at the U-tract (red square) of this YTHDF2 m\(^{\text{6}A}\)-switch (e). The inclusion level of one YTHDF2 exon is co-downregulated by HNRNPC knockdown, METTL3 knockdown and METTL14 knockdown, as validated by RT–PCR. Data are mean ± s.d.; \(n = 3\), biological replicates.

**f**, The inclusion level of one YTHDF2 exon is co-downregulated by HNRNPC knockdown, METTL3 knockdown and METTL14 knockdown, as validated by RT–PCR. Data are mean ± s.d.; \(n = 3\), biological replicates.

**g**, We analysed our polyA\(^{+}\) RNA-seq data to look for reads that span intron/exon junctions on CDS m\(^{\text{6}A}\)-switch containing genes. We find that the control sample has significantly higher reads spanning intron/exon junctions than HNRNPC and METTL3/L14 knockdown samples. This result indicates that m\(^{\text{6}A}\) depletion at the CDS m\(^{\text{6}A}\)-switches promotes intron exclusion.
**Extended Data Figure 9 | Summary of the sequencing samples.**

**a**, For PAR-CLIP–MeRIP and PAR-CLIP experiments from HEK293T cells, the number of mapped reads and 'T-to-C' mutation rates are given for each replicate.

| Experiment     | Sample/replicates | Mapped reads | T-to-C mutation rate |
|----------------|-------------------|--------------|----------------------|
| PAR-CLIP–MeRIP | Input-rep1        | 26,755,227   | 0.67                 |
|                | Input-rep2        | 34,022,750   | 0.67                 |
|                | IP-rep1           | 22,515,469   | 0.83                 |
|                | IP-rep2           | 48,570,954   | 0.76                 |

**b**, For RNA-seq experiments from HEK293T cells, the number of total reads, the number of mapped reads as well as the mapping rates is given for each replicate.

| Experiment     | Sample/replicates | Total reads | Mapped reads | Mapped rate |
|----------------|-------------------|-------------|--------------|-------------|
| RNA-seq        | Ctrl-rep1         | 78,453,301  | 68,058,993   | 86.70%      |
|                | Ctrl-rep2         | 78,823,710  | 70,907,544   | 90.00%      |
|                | HNRNPC KD-rep1    | 81,933,883  | 72,840,271   | 88.90%      |
|                | HNRNPC KD-rep2    | 78,133,137  | 68,053,888   | 87.10%      |
|                | METTL3 KD-rep1    | 58,140,943  | 48,123,493   | 82.80%      |
|                | METTL3 KD-rep2    | 58,923,206  | 53,011,571   | 90.00%      |
|                | METTL14 KD-rep1   | 77,932,017  | 68,703,305   | 88.20%      |
|                | METTL14 KD-rep2   | 75,442,883  | 65,812,269   | 87.20%      |

**c**, Scatter plots comparing transcripts for all PAR-CLIP replicate experiments. The square of Spearman’s rank correlation value ($r^2$) for each pair is shown in the top left corner of the respective panel.

**d**, The detected expression level changes show a strong correlation between gene knockdown replicates. Scatter plots comparing the fold changes (log2) in normalized gene expression from replicates of HNRNPC, METTL3 and METTL14 knockdown. The square of Spearman’s rank correlation value ($r^2$) for each pair is shown in the top left corner of the respective panel.