Single-base mapping of m⁶A by an antibody-independent method

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N⁶-methyladenosine (m⁶A) is one of the most abundant messenger RNA modifications in eukaryotes involved in various pivotal processes of RNA metabolism. The most popular high-throughput m⁶A identification method depends on the anti-m⁶A antibody but suffers from poor reproducibility and limited resolution. Exact location information is of great value for understanding the dynamics, machinery, and functions of m⁶A. Here, we developed a precise and high-throughput antibody-independent m⁶A identification method based on the m⁶A-sensitive RNA endoribonuclease recognizing ACA motif (m⁶A-sensitive RNA-Endoribonuclease–Facilitated sequencing or m⁶A-REF-seq). Whole-transcriptomic, single-base m⁶A maps generated by m⁶A-REF-seq quantitatively displayed an explicit distribution pattern with enrichment near stop codons. We used independent methods to validate methylation status and abundance of individual m⁶A sites, confirming the high reliability and accuracy of m⁶A-REF-seq. We applied this method on five tissues from human, mouse, and rat, showing that m⁶A sites are conserved with single-nucleotide specificity and tend to cluster among species.

INTRODUCTION

Among more than 100 RNA chemical modifications, N⁶-methyladenosine (m⁶A) is one of the most abundant forms in eukaryotic mRNA, accounting for 0.1 to 0.4% of all adenosine (1). The m⁶A modification influences various stages of mRNA metabolism and the biogenesis of long noncoding RNA and microRNA, with effects across diverse biological processes including neuronal development, cell fate transition, immune response, DNA damage response, and tumorigenesis (2–10). Most m⁶A sites were found in conserved motif DRACH (D=G/A/U, R=G/A, H=A/U/C) (1), with an estimated methylation ratio ranging from 6 to 80% (11). Previous whole-transcriptome m⁶A maps have suggested that m⁶A modifications are enriched around stop codons, implying the functional importance of distribution pattern for m⁶A (12–14).

The chemical properties of m⁶A are similar to adenosine, making it difficult to discriminate by chemical reactions (15). The recently developed high-sensitivity liquid chromatography–tandem mass spectrometry and blotting methods relying on antibodies were widely used to quantify the overall m⁶A level. To detect individual sites, methyl-sensitive ligase has been applied to confirm the methylation status of specific adenosines (16, 17), while the method called SCARLET (site-specific cleavage and radioactive-labeling followed by ligation-assisted extraction and thin-layer chromatography) can quantify the methylation level of individual m⁶A site (11). Other methods were developed to identify m⁶A in single-base resolution during reverse transcription, taking advantage of m⁶A-sensitive reverse transcriptase (18, 19), chemoenzymatic substitution of the N⁶-methyl group (20), or selective dTTP (deoxythymidine triphosphate) analog such as 4SdTTP (21). Along with the rapid progress of single-molecule sequencing technologies, Oxford Nanopore Technologies sequencing platform is able to detect modifications on model RNA (22), although the systematic error prevents the practical application in biological samples. Nevertheless, comprehensive interrogating of m⁶A at the transcriptome level is pivotal to reveal the biological importance of this mRNA modification. Methylation RNA immunoprecipitation and sequencing (MeRIP-seq or m⁶A-seq) has been widely used to profile m⁶A, identifying approximate region with m⁶A in ~100 nucleotide (nt) length, while the exact location of individual m⁶A site remains undetermined (12, 13). Many refined methods have been developed to improve resolution, such as PA-m⁶A-seq, miCLIP, and m⁶A-CLIP (23–26). However, all of these methods are dependent on m⁶A-specific antibodies, suffering from poor reproducibility and complicated process. In addition, affinity variation and batch effects of antibodies make it difficult to quantify the methylation level (27). Therefore, a convenient and single-base resolution method is still needed for whole-transcriptome m⁶A identification and quantification, advancing the comprehension of m⁶A for its dynamics and cellular functions in posttranscriptomic regulation.

Several DNA endonucleases belonging to the restriction-modification system have demonstrated sensitivity to DNA methylation. For instance, Dpn I specifically cleaves methylated sites, while Dpn II is blocked by the same modified sequence. This feature has been adopted in genome-wide studies to detect the DNA 6mA modification in single-base resolution (6mA-RE-seq/DA-6mA-seq) (28). For RNA endoribonucleases, a large group of sequence-specific enzymes belonging to the bacterial type II toxin-antitoxin system has been found with determined cleavage motifs. Recently, an Escherichia coli toxin and RNA endoribonuclease, MazF, was reported to be sensitive to m⁶A modification within ACA motif, specifically cleaving the unmethylated ACA motif, leaving methylated (m⁶A)CA motifs intact (29). By screening the endoribonuclease pool, we identified ChpBK as another m⁶A-sensitive endoribonuclease, which can discriminate m⁶A-modified motifs from unmodified sequences. Taking advantage of the m⁶A-sensitive endoribonucleases, we developed m⁶A-sensitive RNA-endoribonuclease–facilitated sequencing method or m⁶A–REF-seq, which can identify transcriptomic m⁶A sites at specific motifs.
and quantify the methylation level in single-base resolution. To validate the m\textsuperscript{6}A sites identified by this method, we used a ligation-based method testing individual sites and confirmed the high-reliability results. Further analysis revealed distribution pattern of m\textsuperscript{6}A in single-nucleotide resolution. Last, we applied m\textsuperscript{6}A-REF-seq to five tissues from human, mouse, and rat, revealing the conservation of m\textsuperscript{6}A at both single-base and regional levels across diverse tissues and species.

RESULTS
Identification and quantification of m\textsuperscript{6}A by endoribonuclease

The application of methylation-sensitive DNA endonuclease in genome-wide 6mA identification inspired us to inspect the possibility of finding similar tools for m\textsuperscript{6}A determination (28). To discover endoribonuclease with m\textsuperscript{6}A sensitivity, we expressed and screened the candidate enzymes by testing the cleavage capacity to synthetic RNA oligonucleotides (RNA oligos) with or without m\textsuperscript{6}A. Two enzymes, MazF and ChpBK, both belonging to the bacterial toxin-antitoxin system, were able to distinguish m\textsuperscript{6}A from unmethylated A. MazF recognized and cleaved the motif sequence ACA from the 5′ side of first A, leaving the methylated (m\textsuperscript{6}A)CA motif intact (Fig. 1A). The cleavage activity of ChpBK was also blocked by m\textsuperscript{6}A at the recognition motif UAC (Fig. 1B). Synthetic RNA oligos with different m\textsuperscript{6}A/A ratios were used to simulate the partially methylated sites on mRNA. The cleavage assay demonstrated that the fraction of digested oligo versus intact part was proportional to the methylation ratio, implying that MazF has the potential to quantify the methylation level (Fig. 1, C and D, and fig. S1).

The high sensitivity and specificity of selected endoribonuclease that enzymatically distinguished m\textsuperscript{6}A urged us to develop a new high-throughput method to identify single-base m\textsuperscript{6}A at the transcriptome level. Since the MazF recognition motif ACA is more prevalent than that of ChpBK among known m\textsuperscript{6}A sites, we chose MazF in the following studies. In general, mRNA from human embryonic kidney (HEK) 293T cells was digested into fragments by MazF. After end repair and purification, the mRNA fragments were ligated to next-generation sequencing (NGS) adaptors and amplified by polymerase chain reaction (PCR). To prevent potential false positives caused by unknown factors, we also included a parallel sample batch demethylated by fat mass and obesity-associated protein (FTO) in advance as the negative control (fig. S2). We call this method m\textsuperscript{6}A-REF-seq, which stands for m\textsuperscript{6}A-sensitive RNA-Endoribonuclease-Facilitated sequencing (Fig. 1E).

Transcriptome-wide identification of m\textsuperscript{6}A sites in single-base resolution

We developed a pipeline to analyze the high-throughput sequencing data generated by m\textsuperscript{6}A-REF-seq (Fig. 2A). As we anticipated, most reads contained the ACA at the 5′ terminus, in line with the effect of MazF treatment (fig. S3). The intact ACA motif sequence observed internally within an RNA fragment was supposed to be...
methylated. We verified this using the known m^6A site with ACA motif on the 18S ribosomal RNA (rRNA) as an example. The unmethylated rRNA fragments were cleaved on the ACA motif, and the methylated fragments were retained (Fig. 2B). Next, we scanned each ACA motif within the transcriptome and interrogated the methylation sites by calculating the reads harboring intact internal ACA motifs. To further eliminate the potential false positive caused by RNA structure, we predicted the probability of each RNA fragment forming secondary structure according to minimum free energy and removed the candidate sites tending to reside in double-stranded regions. We performed three biological replicates and retained the recurrent candidate sites detected in at least two replicates (Fig. 2C and table S1). For each site, the ratio of sequence reads with internal ACA motif versus reads split at the end represents the relative methylation ratio. We compared the methylation ratio of each candidate site in MazF treatment to the sample treated with FTO in advance, which was considered as demethylated RNA. Most of m^6A sites had marked reductions in methylation level with FTO treatment (Fig. 2D). Only candidate sites with a substantial decrease of methylation ratio in FTO treatment were considered as accurate m^6A sites. In total, we identified 4260 high-confidence m^6A sites for further studies.

About 65% of these sites were located in the coding sequence (CDS) region, and others were in untranslated regions (UTRs) (Fig. 2E). The transcriptome-wide distribution showed strong enrichment surrounding the stop codon (Fig. 2F), in line with the typical pattern found by antibody-dependent methods such as MeRIP-seq or m^6A-seq. To compare the m^6A location with MeRIP-seq in detail, we downloaded the profiling data deposited in Gene Expression Omnibus (GEO) database (GSE29714) and calculated the cumulated reads distribution of input and IP (immunoprecipitation) samples within the range of −400 to 400 nt flanking to every m^6A site identified.
by m6A-REF-seq. The reads from IP samples were markedly concentrated in this region, with a peak summit at almost the exact position of individual m6A sites; whereas reads from input sample did not show any enrichment, implying that the m6A sites identified by m6A-REF-seq coincided well with the m6A peaks reported by MeRIP-seq (Fig. 2G). By expanding the context sequences flanking m6A sites, we found that GGACA was the most enriched consensus motif. More generally, the consensus motifs DRACA and RRACA were overrepresented, accounting for 56 and 45% of all identified m6A sites, respectively (Fig. 2H). These results were in concordance with previous studies, in which DRACH/RRACH was the most preferred m6A motif in native mRNA (12, 13, 24).

High reliability of individual m6A sites identified by m6A-REF-seq
To show the reliability of m6A-REF-seq as a new method, we sought another independent principle to validate individual m6A sites. The T3 ligase was reported to be sensitive to m6A sites during the ligation reaction, which had been applied practically to determine m6A sites in mRNA (17). We adopted this method and designed the probe L (left) and probe R (right) based on the sequence flanking the exact site (table S2) and then used the T3 ligase to concatenate the two probes to an integrated template, which could be amplified by PCR. According to previous result, the ligation efficiency at m6A site was significantly suppressed comparing to that at unmethylated A sites (17), and PCR amplification magnified the difference of ligation efficiency (fig. S4). Thus, the amount of PCR products could be assessed on gel to represent the ligation efficiency, implying the methylation status of the interrogated site (Fig. 3A). We also applied the same protocol on FTO-demethylated mRNA as negative control. In total, we picked nine m6A sites for validation, including eight m6A sites and one unmethylated site. Six of eight m6A sites were validated by the ligase-based method, while the unmethylated site was also confirmed (Fig. 3B and fig. S5). To the best of our knowledge, this is the first batch of individual m6A sites confirmed by two independent methods. One validated m6A site (chr1: 29070177; Fig. 3B) is located at 343 nt upstream from the stop codon of m6A reader protein YTHDF2, suggesting a self-regulating feedback loop mediated by RNA modification.

Two other antibody-dependent methods, m6A-CLIP and miCLIP, had reported comprehensive m6A maps at transcriptome level. We checked whether the six individual m6A sites could be identified by these two methods. Peaks from MeRIP-seq data were also included as reference. Of the six sites, one of them was not reported in either previous method. The negative control site that supposed to be unmethylated was reported by miCLIP as an m6A site but not by other methods (Fig. 3B). We suspect that differences in samples, as well as the dynamic nature of m6A methylation, contribute to these inconsistencies and potential false positives. Nevertheless, these results underline the need of independent methods to confirm authentic m6A sites for downstream analysis, especially considering m6A as a highly dynamic RNA modification.

Quantification of m6A abundance by m6A-REF-seq
m6A-REF-seq not only determines the methylation status but also can be used to quantify the methylation level of each m6A site by calculating the ratio of sequence reads with internal ACA versus reads split at the motif. To verify this, we designed a digestion-quantification assay that used reverse transcription quantitative PCR (RT-qPCR) to evaluate the methylation level of identified m6A sites. First, mRNA was treated by MazF. The unmethylated sites located in ACA motif were cleaved, leaving intact methylated sites. After reverse transcription, complementary DNA (cDNA) spanning the intended m6A site was amplified by PCR to estimate the relative abundance. The same amount of mRNA without MazF digestion was amplified simultaneously as a control to compare with the digested sample (Fig. 3C). We picked six m6A sites including three sites with high methylation level and three sites with low methylation level determined by m6A-REF-seq (table S3). The ΔΔCt values calculated from RT-qPCR results were comparable to the average methylation ratio determined by m6A-REF-seq (Fig. 3D and fig. S6), demonstrating that m6A-REF-seq can principally quantify the methylation fraction of m6A sites.

Characterizing m6A distribution pattern in single-nucleotide resolution
Previous studies demonstrated that one of the most remarkable features of m6A was the distribution enrichment surrounding stop codon (12, 13). However, limited by the resolution of antibody-based methods, it was hard to know the precise position relative to the exact stop codon site, which divides coding regions and 3’UTRs. The single-nucleotide m6A identification method largely increased the resolution of m6A map. Our results showed that this stop codon–concentrated pattern could be reproduced by using single-nucleotide m6A sites, without any preference to coding regions or 3’UTRs (Fig. 4A). Meanwhile, it was worth noting that consensus motifs DRACA and DRACH were also enriched near the stop codon, suggesting that sequence motif might be one factor to shape this pattern (Fig. 4A). m6A had been proposed to be involved in various key regulatory processes of RNA metabolisms, including splicing (31). We calculated the relative distances of m6A sites to intron-exon boundaries in each exon and plotted in Fig. 4B. The result showed m6A distributed evenly along exon regions, with slight depletion toward the 3’end. By mapping single m6A sites to exons, we compared the lengths of exons containing m6A modifications versus all exons. The lengths of m6A-containing exons were significantly greater than all exons (P < 2.2 × 10^-16, Wilcoxon test; Fig. 4C), in line with previous report that m6A modification usually occurred within long exons (13).

Some indirect evidence showed that m6A tended to locate in short regions, suggesting a specialized working model of m6A as clusters. However, antibody-based methods could barely divide multiple sites from one m6A peak to demonstrate clustering. Using the single-nucleotide m6A positions identified by m6A-REF-seq, we calculated the distances between each two adjacent m6A sites and compared to the control dataset, which contained the same number of sites randomly sampled from transcriptome 1000 times. Our results showed that two adjacent m6A sites were statistically prone to aggregate within 200-nt regions (P = 6.6 × 10^-5 and P = 0.0065, Fisher’s exact test), supporting the hypothesis that m6A functions as clusters of modification (Fig. 4D).

Conservation of m6A sites in mammals
We next applied m6A-REF-seq to identify m6A modifications in different tissues of human, mouse, and rat and identified more than 3500 individual m6A sites in each sample (tables S4 and S5). The transcriptomic distribution displayed similar patterns among tissues in three species (Fig. 5A and figs. S7 to S9). To examine the conservation of m6A at the individual gene level, we first collected 13,117 one-to-one orthologs in three species. By mapping the m6A sites to these ortholog genes in each type of tissues, we identified 2439
human genes, 3596 mouse genes, and 1343 rat genes in the brain, among which 571 of them were shared in three species. The numbers of shared genes were 536 and 262 in the kidney and the liver, respectively (figs. S8 and S9). The ortholog genes modified by m6A in all three species were significantly more than expected even after adjusting for gene expression (Fig. 5B, fig. S10, and table S6), indicating a well-conserved distribution of m6A at the gene level.

Similarly, previous studies have reported that the count of shared m6A peaks between human and mouse were significantly higher than expected, proposing the conservation and important functions of m6A modification (13).

Since m6A-REF-seq identified high-confidence sites, we were able to investigate the conservation of m6A at single-nucleotide level. Previous work analyzed the evolutionary conservation of yeast and
human m\textsuperscript{6}A sites in protein-coding regions and concluded that m\textsuperscript{6}A were overall not conserved in yeasts and only slightly conserved in mammals (32). However, the m\textsuperscript{6}A sites used in this work were predicted based on the consensus motif DRACH, which could contain underestimated false positives. Therefore, we mapped the single-nucleotide m\textsuperscript{6}A sites identified by m\textsuperscript{6}A-REF-seq to orthologous genes in three species (table S6). By using similar method (32), we tested whether m\textsuperscript{6}A site were more conserved than unmethylated A. We randomly sampled the human unmethylated A sites as the same number of m\textsuperscript{6}A sites in each orthologous gene and tested whether the corresponding sites in mouse and rat remained to be base A (fig. S11). A total of 1854 to 3722 m\textsuperscript{6}A sites were compared to the same number of unmethylated A sites from 1286 to 2306 genes across three tissues. The results indicated that m\textsuperscript{6}A sites were more conserved than unmethylated A sites in brain, kidney, and liver tissues for all three codon positions ($P = 0.002$, $0.021$, and $0.075$, respectively; figs. S11 to S13), confirming the m\textsuperscript{6}A conservation at single-nucleotide level.

Furthermore, we tested whether the nonpairwise m\textsuperscript{6}A sites tended to cluster in narrow corresponding regions among species. The shortest distances between pairwise m\textsuperscript{6}A sites in ortholog genes were averaged (Fig. 5C). Random sampling of the same number of ACA motifs were performed for 1000 times as control. The average distances between pairwise m\textsuperscript{6}A sites were shorter than the control (Fig. 5D, and figs. S8 and S9), indicating that, although some m\textsuperscript{6}A sites were not conserved at single-nucleotide level, they tended to cluster within conserved short regions across multiple species.

More extensively, we computed the PhastCons scores representing the conservation of each nucleotide among multiple species (33). Identified m\textsuperscript{6}A sites in the brain were attributed and classified into sites conserved in three species or sites residing in ortholog genes. As expected, conserved m\textsuperscript{6}A sites at single-base level and sites conserved at gene level had highest conservation scores (medians 0.991 and 0.970, respectively). The scores of overall m\textsuperscript{6}A sites were also distinctively higher than that of all A sites in ACA motifs with the medians 0.791 and 0.338, respectively ($P < 9 \times 10^{-10}$, Wilcoxon test; Fig. 5E). Similar results hold for other tissues in human, mouse, and rat (figs. S8, S9, and S14).

**DISCUSSION**

Antibody-dependent pull-down methods are widely used to profile transcriptomic m\textsuperscript{6}A, the most prevalent modification in mRNA. An antibody-independent, easy-to-use method is strongly desired to cross-validate the known m\textsuperscript{6}A sites and study the dynamic functions in single-nucleotide resolution. We developed m\textsuperscript{6}A-REF-seq, an antibody-independent, high-throughput, and single-base m\textsuperscript{6}A detection method based on m\textsuperscript{6}A-sensitive RNA endoribonuclease. High sensitivity and specificity of endoribonuclease that we found to discriminate m\textsuperscript{6}A modification, as well as stringent data process
by removing potential RNA secondary structure and the introducing of FTO demethylation control, ensured reliability of m⁶A sites identified by this method. Transcriptome-wide m⁶A maps in human HEK293T cell and mammalian tissues displayed similar distribution pattern to previous antibody-based pull-down methods. To validate the m⁶A sites identified by m⁶A-REF-seq, we introduced another independent method, which used T3 ligase to differentiate individual m⁶A or A at template RNA. The high consistency between different methods demonstrated the efficiency of m⁶A-REF-seq with high confidence. We also applied qPCR to show that m⁶A-REF-seq could quantify the methylation level of individual m⁶A site.

The RNA endoribonuclease MazF has high sensitivity and specificity for RNA substrates. It works very well on extremely limited substrate, as little as nanograms or even picograms. Rapid and simplified experiment design, without the need for antibody-enrichment step, largely reduces the requirement of starting RNA amount and sample preparation time. Therefore, this method could be applied to the studies for rare biological materials such as samples from pathological tissues or early embryos. In addition, m⁶A-REF-seq offers the potential to capture the subtle changes of m⁶A during metabolic processes, advancing the dynamic studies of this post-transcriptional RNA modification in different life stages.

By screening RNA endoribonucleases, we found two m⁶A-sensitive enzymes from the bacterial type II toxin-antitoxin system, which recognized specific motifs “ACA” and “UAC,” respectively. To cover all the RRACH motifs in the transcriptome, we need new enzyme(s) capable of recognizing more universal sequence motifs while retaining the m⁶A sensitivity. On the basis of our preliminary screening, we speculated that the m⁶A sensitivity might be a general feature of endoribonucleases belonging to type II TA system. Further exploring in the endoribonucleases pool would be helpful to expand the toolbox for RNA modification studies. Another strategy is to transform the existing enzyme to adapt for new recognition sequences while retaining the methylation sensitivity. We studied the structure of MazF protein and mutated the 56th lysine to alanine. The MazF-K56A mutant showed irregular recognition sequences not meeting our anticipation, although the methyl sensitivity was retained (fig. S15). On the other hand, directed evolution would be a promising way to expand the recognition sequence scope of MazF or other related proteins, as long as dedicated screening system was created.

We applied m⁶A-REF-seq to five different tissues from human, mouse, and rat. The single-base m⁶A maps displayed a distribution pattern similar to previous findings that m⁶A tended to enrich surrounding stop codons. This unique pattern seemed to be conserved among different species. However, because of limited resolution, previous methods could not tell to what extent the modification is conserved. In this study, we mapped the m⁶A sites to orthologous genes in these three species and compared the conservation score at single-nucleotide level. We found that significantly more m⁶A-modified genes were shared among species than expected, and a large group...
of m^6^A sites could be aligned to the syntenic regions, showing that the m^6^A modification was conserved at both the gene and individual base level. The well-conserved feature restated the biological importance of m^6^A, implying other uncovered roles of this RNA modification.

Together, m^6^A-REF-seq provides a new perspective for single-base m^6^A identification at the transcriptome level. The new principle eliminates the need for m^6^A-specific antibodies, making it unique and amenable for future applications in small and precious samples. The features of this efficient and convenient method will largely expand the scope of m^6^A studies.

**MATERIALS AND METHODS**

**RNA materials**
The RNA oligos synthesized from Takara Bio Inc. and Sangon Biotech were as follows: RNA-m^6^A 5′-UUGGGUUUUUUGG(m^6^A)CUGUAUUGAGU-3′, RNA-ACA 5′-UUGGGUUUUUUUG-GACAUGUAUAUGU-3′, RBl-Um^6^A 5′-GUUGUGUGUAGAU(m^6^A)-CAUUGUGUGUGG-3′, and RBl-UAC 5′ GUUGUGUAAUACAUGUGGUGG-3′.

HEK293T cell line was cultured with Dulbecco’s modified Eagle’s medium (DMEM; Corning) supplemented with 10% (v/v) fetal bovine serum (FBS; Gibco) at 37°C in the presence of 5% CO_2_ and tested to be mycoplasma negative. Total RNA from HEK293T cells was extracted using TRIZol reagent (Thermo Fisher Scientific), and the mRNA was purified using the Dynabeads mRNA Purification Kit (Thermo Fisher Scientific, catalog no. 61006). The mRNA of human tissues (brain, liver, and kidney), mouse tissues (brain, liver, heart, tests, and kidney), and rat tissues (brain, liver, and kidney) were purchased from Takara Bio Inc.

**RNA endoribonuclease validation for MazF**
RNA oligos (10 pmol) were incubated with 2.5 U of MazF (mRNA interferase MazF, Takara Bio Inc., code no. 2415A) in the 20-μl reaction mixture of MazF buffer [40 mM sodium phosphate (pH 7.5) and 0.01% Tween 20] at 37°C for 30 min. The RNA oligos were mixed with the m^6^A percentage of 0, 20, 40, 60, 80, and 100%, and the oligo mixtures were incubated with 2.5 U of MazF in the 20-μl reaction mixtures. The samples were loaded into 15% urea polyacrylamide gel and electrophoresed in 0.5× tris-boric acid–EDTA (TBE) buffer.

The FTO demethylation reaction was conducted in the reaction mixture that contained 10 pmol of m^6^A RNA oligo, 0 to 10 μg of FTO demethylase, 283 μM (NH_4)_2Fe(SO_4)_2·6H_2O, 300 μM α-ketoglutarate (α-KG), 2 mM L-ascorbic acid, 20 U of RNase inhibitor (Takara Bio Inc., code no. 2313A), and 50 mM tris-HCl buffer (pH 7.5). The reaction was stopped by 40 mM EDTA after 3 hours of incubation at room temperature. The MazF cleavage reaction was performed to test the efficiency of FTO demethylation.

**Dot-blot assay of FTO demethylation**

m^6^A FTO demethylase with different concentrations (0, 0.25, and 2.5 μg) was used in the dot-blot assay to test the demethylation of RNA. The total RNA was incubated with FTO, 283 μM (NH_4)_2Fe(SO_4)_2·6H_2O, 300 μM α-KG, and 2 mM L-ascorbic acid for 3 hours at room temperature. After purification by an RNA Clean & Concentrator-5 kit (Zymo Research), the concentration of total RNA was determined by the Qubit RNA HS Assay Kit (Thermo Fisher Scientific) and diluted to 10-μl volume with 500 ng. Spotting and thermal cross-linking were carried out on a metal base plate preheated to 70°C. After thermal cross-linking for 15 min, ultraviolet cross-linking was conducted twice with 2000 kJ. Then, 3% bovine serum albumin (BSA) was used to block at room temperature for 1 hour. The anti-m^6^A antibody (Synaptic Systems, catalog no. 202003) was diluted to 1:2000 and incubated with the nylon membrane for 2 hours. The nylon membrane was washed for three times with 1% tris-buffered saline with Tween 20 (TBST) and then incubated with the 1:5000 diluted anti-rabbit antibody for 1 hour. The membrane was washed for five times with 1% TBST. After drying off the water, the chemiluminescence detection signal was performed using the enhanced chemiluminescence (ECL) system. The membrane was washed twice with 1% TBST. After drying off the water, it was dyed with methylene blue solution for 8 min and then washed with 1% TBST. The signal was detected by a white light system.

**Expression and purification of recombinant ChpBK protein**
To express recombinant ChpBK protein [National Center for Biotechnology Information (NCBI) reference sequence: NP_418646.1], E. coli BL21(DE3) strain carrying PET-28a-chpBK plasmid was grown in lysogeny broth (LB) medium (Sangon Biotech, A507002) supplemented with Kan (10 μg/ml) at 37°C and 200 rpm until the optical density at 600 nm (OD_{600}) reach to 0.5. Isopropyl-β-D-thiogalactopyranoside (IPTG) was added to a final concentration of 0.4 mM, and cell growth resumed for 2 hours under the same conditions. Harvested cell pellets were lysed by sonication in 30 ml of buffer A [10 mM bis-tris (pH 7.0) and 30 mM imidazole] and centrifuged for 20 min at 16,000g and 4°C. The soluble fraction was incubated on ice for 30 min in the presence of DNase and RNase (10 g/ml; Thermo Fisher Scientific). The His-ChpBK fusion protein was purified from clarified lysates using Ni-NTA (nickel-nitrilotriacetic acid) beads (Invitrogen). Protein purity was determined by SDS–polyacrylamide gel electrophoresis (SDS-PAGE) analysis, and samples were quantified using the Bio-Rad protein assay and stored at ~80°C.

**Expression and purification of MazF-K56A in E. coli**
The 56th lysine of MazF protein was mutated to alanine. The protein sequence was shown below, where the underlined bold “A” indicates the mutated amino acid: MazF-K56A (MVSRYVPDMGDLIW-GYPFEVVLSQGERDGVKIAWRAGAT-KKGTVAPEELQIKAINVLAG).

The plasmid DNA pCOLD-MazF was transformed into E. coli Rosetta strain, and a single colony was picked by sterilized loop and inoculated into a 50-ml culture (containing ampicillin) in sterile culture flask. E. coli was cultured at 37°C until OD_{600} reached 0.6 to 0.8. IPTG (1 mM) was added to induce protein expression, and cells grew overnight at 18°C. Cells were harvested by centrifugation and suspended in 200 ml of lysis buffer [50 mM tris (pH 8.0), 350 mM NaCl, and 10 mM imidazole]. Cell pellets were lysed by sonication, and the supernatant was collected by centrifugation and applied to nickel affinity purification using HisTrap HP. The column was washed by lysis buffer to remove nonspecific binding protein, and target protein was eluted with linear gradient of elution buffer [50 mM tris (pH 8.0), 350 mM NaCl, and 500 mM imidazole]. We concentrated the protein and changed protein storage buffer to ion-exchange binding buffer [20 mM Hepes and 50 mM NaCl (pH 7.5)] by HiTrap Desalting column. Tobacco etch virus (TEV)
protease was used to remove the His-tag, and target protein was loaded to HiTrap HP Q column. Target protein was eluted by linear gradient of ion-exchange elution buffer [20 mM Heps and 1 M NaCl (pH 7.5)]. The concentrated protein was analyzed by SDS-PAGE.

**RNA endoribonuclease validations for ChpBK and MazF-K56A**

The digestion reaction of ChpBK was conducted in a 20-μl reaction mixture with 10 pmol of RNA oligo (RB1), 1 μg of ChpBK, 20 U of RNase inhibitor, and 50 mM tris-HCl buffer (pH 7.5). The mixture was incubated at 37°C for 30 min. The digestion reaction of ChpBK with different m6A percentage was conducted as described above. The MazF-K56A cleavage reaction was performed in a 20-μl reaction mixture with 10 pmol of RNA oligo, 2 μg of MazF-K56A, and 1× MazF buffer [40 mM sodium phosphate (pH 7.5) and 0.01% Tween 20] at 37°C for 30 min. All samples were loaded on the 15% urea polyacrylamide gel and electrophoresed in 0.5× TBE buffer.

**Library construction for NGS**

mRNA was first heated at 85°C for 5 min and chilled on ice for 2 min to denature RNA secondary structure. The FTO demethylation reaction was conducted before MazF reaction in a 20-μl reaction mixture containing 100 to 200 ng mRNA, 2.5 μg of FTO demethylase, 283 μM (NH4)2Fe(SO4)2·6H2O, 300 μM α-KG, 2 mM l-ascorbic acid, 20 U of RNase inhibitor, and 50 mM tris-HCl buffer (pH 7.5). After incubating at room temperature for 3 hours, the reaction was stopped by adding 40 mM EDTA. To prevent the influence of FTO buffer, the MazF treatment was supplemented with the same reaction mixture except FTO protein. The MazF and FTO-MazF samples were heated at 85°C for 5 min, chilled on ice for 2 min, and then incubated with 10 U of MazF in the MazF reaction buffer at 37°C for 30 min. The fragmented mRNA was purified by an RNA Clean & Concentrator-5 kit (Zymo Research) and eluted into 20 μl of RNase-free water. The eluted RNA fragment was end repaired by the T4 polynucleotide kinase (T4PNK; Vazyme Biotech) in 50 -μl reaction mixture with 1× T4PNK buffer and incubated at 37°C for 30 min. After purification by an RNA Clean & Concentrator-5 kit (Zymo Research), the concentration of fragmented mRNA was measured by the Qubit RNA HS Assay Kit (Thermo Fisher Scientific). The NGS library was constructed by using the VAHTS Small RNA Library Prep Kit (Vazyme Biotech, NR801) with 10 ng of fragmented mRNA and amplified by PCR for 15 cycles. Libraries were loaded on the 8% urea polyacrylamide gel and electrophoresed in 0.5× TBE buffer, and purified libraries were sequenced on the Illumina HiSeq X10 platform. Three replicates have been conducted for mRNA of HEK293T cell line. The same protocol was also conducted for mammalian tissues.

**NGS data analyses**

First, quality control was performed on raw sequencing reads by the open software FastQC. Then, adaptors were filtered by cutadapt (34) with at least 15 nt remaining length of paired-end reads. The clean reads were mapped to the reference genome (hg19) using HISAT2 (35) with default parameters, except reporting only one result. Integrative Genomics Viewer was used for data visualization (36). We scanned the motif ACA (and the reverse complementary motif TGT) on the reference genome and mapping result file, respectively, counting the number of ACA/TGT at the internal or terminal of a sequence read. As short sequence reads with lengths less than 15 nt were filtered, only ACA/TGT with the distance from the end of a read more than 15 nt was treated as “internal” of a read. Each ACA/TGT motif on the reference genome, which has more than 10 reads supporting the motif located internal of reads, was treated as a candidate m6A site. The ratio of sequence reads with internal ACA versus reads split at the end represents the relative methylation ratio of each site. Candidate sites with repeated or continuous ACA motifs were removed.

As the RNA secondary structure could affect the reaction efficiency, we predicted the probability of each RNA fragment forming secondary structure and removed the candidate sites tending to reside in double-stranded regions. The RNAfold program of ViennaRNA package (30) was used to predict intramolecular secondary structure of each read, which supported the candidate sites. The condensed representation of the pair probabilities of each nucleotide was parsed according to the tutorials of ViennaRNA package. The pairing probability value of NNACA motif was calculated, and candidate sites with high pairing probability were discarded. The FTO demethylation treatment was also treated as a negative control. Candidate sites with at least 10% methylation ratio decrease were remained as m6A sites. The metagene plot of m6A sites, which was described the relative location of each site on the 3′ UTR, CDS, and 5′ UTR of mRNA, was calculated and plotted as well. The raw reads of MeRIP-seq datasets were downloaded from Meyer et al. (12) (GSE29714) and mapped to human (hg19) reference genome using HISAT2. The reads within the range of –400 to 400 nt flanking to every m6A site were plotted. As the sequencing data were single end, we extended the reads to 200 nt to simulate the IP peak.

**Single-base validation using ligase-based method**

Refrained single-base T3 ligase–based method (17) was used to validate m6A sites identified by m6A-REF-seq. The FTO demethylation sample was used as negative control (i.e., the FTO treatment in Fig. 3A). DNA probe L and probe R were designed to match theprivate m6A sites with repeated or continuous ACA motifs were removed. The ratio of sequence reads with internal ACA versus reads split at the end represents the relative methylation ratio of each site. Candidate sites with repeated or continuous ACA motifs were removed. The FTO demethylation treatment was also treated as a negative control. Candidate sites with at least 10% methylation ratio decrease were remained as m6A sites. The metagene plot of m6A sites, which was described the relative location of each site on the 3′ UTR, CDS, and 5′ UTR of mRNA, was calculated and plotted as well. The raw reads of MeRIP-seq datasets were downloaded from Meyer et al. (12) (GSE29714) and mapped to human (hg19) reference genome using HISAT2. The reads within the range of –400 to 400 nt flanking to every m6A site were plotted. As the sequencing data were single end, we extended the reads to 200 nt to simulate the IP peak.

**qPCR validation for m6A quantification**

The MazF treatment was performed in 10-μl volume with 500 ng of mRNA and 10 U of MazF at 37°C for 30 min. The digested mRNA and nondigested mRNA were subjected to reverse transcription using 5× TransScript All-in-One First-Strand cDNA Synthesis SuperMix for qPCR (One-Step gDNA Removal, TransGen Biotech, AT341-02). TB Green Premix Ex Taq II (Tli RNaseH Plus, Takara Bio Inc., code no. RR820A) was used on BIO-CFX96 Touch.
Characterizing m^6A distribution pattern in single-base resolution
Consensus motifs DRACA and DRACH were scanned on the reference exons for m^6A, regular A sites in DRACA, and DRACH were computed. To estimate the distance between two adjacent m^6A sites, only genes with more than two m^6A sites were analyzed. We randomly sampled the same number of ACA motifs as the number of observed m^6A and repeated this step for 1000 times as control. The distances between adjacent A sites in control were calculated. Fisher’s exact test was used to test the significance of first two bins.

Conservation of m^6A in the same mammalian tissues
m^6A-REF-seq protocol was used to identify m^6A sites in the different tissues of human (brain, liver, and kidney), mouse (brain, liver, heart, testis, and kidney), and rat (brain, liver, and kidney). The one-to-one orthologous genes in pairwise species of human (GRCh37.p13), mouse (GRCm38.p6), and rat (Rnor_6.0) were downloaded from Ensembl (Ensembl genes 94). A total of 13,117 one-to-one orthologous genes among three species were retained for downstream analyses. Protein sequences of 13,117 orthologs in human, mouse, and rat were aligned by MUSCLE (37) and then converted to corresponding coding sequence alignments using PAL2NAL (38). The shared m^6A-modified genes and share m^6A sites among species were calculated and tested for significances based on the well-aligned orthologous coding-region sequences as mentioned above. The average pairwise distances among closely m^6A sites in three species were calculated and compared with control random sample. The control sample was randomly picked for the same number of ACA motifs as the number of observed m^6A for 1000 times. Analyses of conservation score were conducted by using PhastCons scores of human (hg19), mouse (mm10), and rat (rn6) downloaded from University of California, Santa Cruz (UCSC).

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/7/eaax0250/DC1
Fig. S1. Quantitative demonstration of various fractions of m^6A-containing oligo mixed with unmethylated oligo digested by ChpBK.
Fig. S2. FTO demethylate assay.
Fig. S3. The base composition of mRNA products cleaved by MazF.
Fig. S4. The single-site validation for 18
Fig. S5. The base composition of mRNA products cleaved by MazF.
Fig. S6. Quantitative PCR results of six m^6A sites.
Fig. S7. Metagene plots of m^6A in mouse heart and mouse testis.
Fig. S8. Conservation of m^6A in mammalian kidney.

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Competing interests: C.H. is a scientific founder and a member of the scientific advisory board of Accent Therapeutics, Inc.

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