Widespread occurrence of 5-methylcytosine in human coding and non-coding RNA

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ABSTRACT

The modified base 5-methylcytosine (m⁵C) is well studied in DNA, but investigations of its prevalence in cellular RNA have been largely confined to tRNA and rRNA. In animals, the two m⁵C methyltransferases NSUN2 and TRDMT1 are known to modify specific tRNAs and have roles in the control of cell growth and differentiation. To map modified cytosine sites across a human transcriptome, we coupled bisulfite conversion of cellular RNA with next-generation sequencing. We confirmed 21 of the 28 previously known m⁵C sites in human tRNAs and identified 234 novel tRNA candidate sites, mostly in anticipated structural positions. Surprisingly, we discovered 10,275 sites in mRNAs and other non-coding RNAs. We observed that distribution of modified cytosines between RNA types was not random; within mRNAs they were enriched in the untranslated regions and near Argonaute binding regions. We also identified five new sites modified by NSUN2, broadening its known substrate range to another tRNA, the RPPH1 subunit of RNase P and two mRNAs. Our data demonstrates the widespread presence of modified cytosines throughout coding and non-coding sequences in a transcriptome, suggesting a broader role of this modification in the post-transcriptional control of cellular RNA function.

INTRODUCTION

The presence of 5-methylcytosine (m⁵C) in DNA and its role as an epigenetic marker of genome activity is well established (1–3). This has been facilitated in large part by the ease of its detection using bisulfite sequencing, which involves chemical conversion of cytosine (but not m⁵C) to uracil (4–6). While DNA is relatively devoid of other modifications, 109 modifications have been identified in different classes of RNA across all three domains of life (7). tRNA is a particularly heavily modified RNA class, and m⁵C sites have been identified in numerous archaeal and eukaryotic tRNAs, commonly around the variable region and the anticodon loop. The modification has been shown to stabilize tRNA secondary structure, affect aminoacylation and codon recognition, and confer metabolic stability (8–13). m⁵C sites are also found in rRNA where they play roles in translational fidelity and tRNA recognition (14). Interestingly, work that led to the discovery of the mRNA cap structure also detected a low level of internal m⁵C in mammalian mRNA (15) and viral RNAs infecting mammalian cells (16–18), although specific m⁵C sites were not mapped and the methylation was not corroborated by all studies at the time (19–23). More recently, it was reported that the methyl-CpG binding protein 2 (MECP2) associates with RNA and can regulate mRNA splicing (24,25) and that reprogramming of cells to pluripotency can be achieved using m⁵C and pseudouridine-modified mRNAs encoding the four Yamanaka factors (26). These observations have rekindled interest in the occurrence and function of m⁵C in mRNA and other non-coding RNA.
Two m⁵C methyltransferases (MTases) have been shown to catalyze the m⁵C modification of eukaryotic RNA. First, NCL1/TRM4 (Nuclear protein 1 / tRNA-specific MTase 4) is responsible for all known m⁵C sites in yeast tRNA (27), however its human ortholog NOP2/Sun domain protein 2 (NSUN2; also known as Misu (28,29)) may have a much narrower target range, selectively methylating the wobble position of tRNA^{Leu(CAA)} prior to intron splicing (30). Secondly, TRDMT1 (tRNA aspartic acid MTase 1, also known as DNMT2), has been shown to methylate position 38 on tRNA^{Asp(GUC)} in eukaryotes (12,31,32), and tRNA^{Val(AAC)} and tRNA^{Gly(GCC)} in Drosophila (11). TRDMT1 was previously thought to act as a DNA MTase; however, it is now primarily regarded as an RNA MTase (33). The range of RNA targets for these two enzymes in animals are largely unexplored (34). Importantly, NSUN2 is cell cycle-regulated (35), directly targeted by MYC (myelocytomatosis viral oncogene homolog) and implicated in cancer cell proliferation (28). NSUN2 knockout mice are small and have revealed a role of the enzyme in balancing stem cell self-renewal and differentiation (36). Loss of TRDMT1 enzymatic activity in zebrafish is responsible for all known m⁵C sites in yeast tRNA (27), whereas its human ortholog NOP2/Sun domain protein 2 (NSUN2) acts on cell RNA to ensure tRNA representation in the sample. We show that known and novel sites appear largely dependent on the m⁵C-specific MTase NSUN2. These observations suggest that cytosine methylation in RNA is important to the control of cell growth and differentiation, thus motivating global screens for the occurrence of m⁵C in RNA.

Herein we have devised a method for transcriptome-wide detection of modified cytosine residues at single nucleotide resolution by combining RNA bisulfite conversion with next-generation sequencing. We report that RNA cytosine modification pervades the human transcriptome; we discovered over ten thousand novel candidate sites in mRNAs and various non-coding RNA types that are distributed in non-random patterns. Furthermore, we show that known and novel sites appear largely dependent on the m⁵C-specific MTase NSUN2. These data represent the first high-resolution view of cytosine modification across a transcriptome and provide a basis for exploration of its biological significance for mRNA and non-coding RNA function.

MATERIALS AND METHODS

Unless otherwise stated below, all kits and reagents were used according to the manufacturer’s instructions.

Cell culture and RNAi-mediated methyltransferase knockdown

HeLa cells were cultured in DMEM supplemented with 10% FBS (both Gibco-Invitrogen), and incubated with 5% CO₂ at 37°C. Total RNA was extracted from cells using TRIzol (Invitrogen). For RNAi-mediated knockdown of DNMT1, TRDMT1 and NSUN2, cells were transfected with siGenome SMART pool siRNAs or a non-targeting control pool (Dharmacon) using Lipofectamine RNAiMAX transfection reagent (Invitrogen). Cells were passaged and transfected again 72 h post-initial transfection. RNA was harvested from cells 6 days post-initial transfection. For verification of knockdown, 1 µg of total RNA was used for cDNA synthesis with the SuperScript III Reverse Transcriptase kit (Invitrogen). Real-time PCR was performed using SYBR Green 1 Master (Roche) on a LightCycler 480 (Roche) instrument and results were analysed using Relative Quantification Software (Roche). Oligonucleotides used for real-time PCR are listed in Supplementary Table S1.

RNA isolation, and bisulfite conversion of RNA

Resuspended RNA was treated with DNase (Ambion), and then phenol/chloroform extracted and re-precipitated with ethanol. The RNA was then enriched for mRNA by two iterations of oligo-dT-selection on magnetic beads using an mRNA isolation kit (New England Biolabs). Briefly, 100 µg of the total HeLa RNA was incubated with 200 pmol of biotin-(dT)18 conjugated to streptavidin magnetic beads (New England Biolabs). The sample was washed, eluted and then resubjected to the same process, followed by rRNA depletion using the Ribonuclease Kit (Invitrogen). About 4 µg of the mRNA-enriched sample was subjected to bisulfite treatment, after addition of 400 pg of in vitro transcribed Renilla luciferase (R-Luc) RNA (39) as a negative control and 4 ng of total HeLa cell RNA to ensure tRNA representation in the sample.

Bisulfite conversion of RNA was performed as previously described (40), with the following modifications. About 4 µg of RNA was mixed in 100 µl of 40% sodium bisulfite (Sigma), 600 µM hydroquinone (Sigma) solution (pH 5.1) and incubated at 75°C for 4 h. The reaction mixture was then desalted by two passages through Micro Bio-spin 6 chromatography columns (Bio-Rad). RNA was desulfonated by adding an equal volume of 1 M Tris (pH 9.0) to the reaction mixture and incubated for 1 h at 75°C, followed by ethanol precipitation.

Conventional bisulfite sequencing

A 220 ng aliquot of bisulfite-converted RNA was converted to cDNA using random hexamers and the Superscript III Reverse Transcriptase kit (Invitrogen). PCR conditions were optimized for each respective primer set (Supplementary Table S2) and the target products were isolated from primers and spurious products by agarose gel electrophoresis followed by band excision and purification using a Gel Extraction Kit (Qiagen). Amplicons were ligated into the pGEM-T Easy Vector system (Promega) and individual clones sequenced to determine bisulfite conversion efficiency of selected RNAs.

Novel m⁵C candidate sites were verified using new batches of HeLa cell total RNA. Where indicated, negative control transcripts corresponding to the local sequence around the candidate site were generated by in vitro transcription and spiked into the HeLa cell RNA to guard against non-conversion artefacts due to RNA sequence or structure. Briefly, multiple gene
constructs were generated from unconverted HeLa cell cDNA to contain a stretch of sequence surrounding selected candidate m^5C sites and flanked by unique priming sequences. Amplicons were cloned using the pGEM-T Easy Vector system (Promega) and used as templates for in vitro transcription using the MEGAscript high yield transcription kit (Ambion). Transcribed RNA was purified using MEGAclear columns (Ambion) and 400 pg of each in vitro transcript was spiked into HeLa cell total RNA (4 μg). The pool of RNA was then DNase treated, bisulfite-converted and conventionally sequenced as described above. Oligonucleotides used to generate in vitro transcription constructs and sequencing clones are listed in Supplementary Table S2.

Library preparation and SOLiDTM sequencing of bisulfite-converted RNA

To ensure decapping and phosphorylation of the 5' end of the RNA, 500 ng of the bisulfite-converted sample was treated with five units of Tobacco Acid Pyrophosphatase (Epicentre Biotechnologies) at 37°C for 45 min followed by phenol/chloroform extraction. The sample was then subjected to T4 Polynucleotide Kinase (New England Biolabs) treatment and subsequent phenol/chloroform extraction. A next-generation sequencing library was prepared using the SOLiDTM Whole Transcriptome Analysis kit (Applied Biosystems). Since the sample was already sufficiently fragmented by the bisulfite-conversion reaction, the RNA fragmentation step prior to adaptor hybridization and ligation was not performed. cDNA with an approximate insert length of 50–120 nt was selected by polyacrylamide gel electrophoresis. Beads were prepared, deposited and sequenced on a SOLiDTM Version 3 instrument (Applied Biosystems). Sequenced read data was deposited in the National Center for Biotechnology Information sequence read archive (accession number SRA027832.1).

Sequenced read mapping and analysis

Sequenced reads were mapped against reference sequences consisting of all known human transcripts from the Ensembl v61 database (mRNA, rRNA, tRNA, mitochondrial RNA and all non-coding RNA) (41), predicted and known tRNA sequences obtained from the GrtRNAdb (42) and tRNAdb (43), miRNA hairpin sequences from miRBase v16 (44), rRNA sequences obtained from the NCBI RefSeq database, and R-Luc spike-in negative control sequences (Supplementary Table S3). A 'CCA' sequence was appended to tRNAs lacking one since this non-templated addition is common to tRNA sequences. To reduce mapping ambiguity, identical sequences were collapsed and represented only once in the reference.

Sequenced reads were mapped using the SOCS-B program (45), an alignment tool designed to map bisulfite-converted SOLiDTM colour-space reads to nucleic acid reference sequences. This program disregards mismatches between 'C' in the reference and 'T' introduced in reads as a result of bisulfite conversion. SOCS-B mapping parameters were chosen such that low-quality reads (average quality across a read <18) and reads with ambiguous colour-call represented by '.' in the colour-space sequence were discarded. During mapping, up to four colour-space mismatches were allowed between the reference and read sequences to accommodate sequencing errors and natural variation. Given the large proportion of 'predicted' tRNA sequences in the reference, reads multi-mapping to both known and predicted tRNA sequences were preferentially assigned to the known tRNA sequence. For all other reads, SOCS-B was set to assign multi-mapping reads to a single randomly selected mapped locus. Reads mapping to the anti-sense strand of the reference were discarded.

Identification of modified cytosine sites in RNA

Non-conversion of a cytosine in read sequences was taken to indicate the presence of m^5C. To delimit m^5C site prediction to high confidence candidates, threshold parameters were set to a read coverage ≥10 and conversion rate ≥80% per cytosine position in the reference (see ‘Results’ section). Read coverage was defined as the number of reads that overlap a given cytosine. The following equation describes the conversion rate calculations.

\[
\frac{\text{Number of reads representing non-conversion (C)}}{\text{Number of reads representing conversion (T)}} \times \frac{\text{Conversion rate (i x 100)}}{\text{at a given cytosine position = j}}
\]

Read coverage and cytosine conversion rate were calculated directly from mapped data for tRNA, rRNA and R-Luc spike-in negative control sequences. For all other RNA types, cytosine-mapping data was transferred to genomic coordinates before applying threshold criteria (Supplementary Figure S1). Ensembl Perl API v61 was used for coordinate transfers from transcript sequence to the human genome assembly version GRCh37 (hg19).

Analysis of modified cytosine location bias within the transcriptome

Enrichment of non-converted sites across genomic regions was computed by chi-squared and binomial tests, relative to the proportion of all potential sites (cytosines with read coverage ≥10) in each measured category. Enrichment of methylation sites for regulatory elements was estimated by overlap with the 3'UTR structural RNA prediction set of (46) (binomial test relative to the proportion of structured to unstructured nucleotides in 3'UTR).

Enrichment of non-converted sites near binding regions for Argonaute (I-IV) and Pumilio 2 proteins was estimated by overlapping site coordinates with the PAR-CLIP dataset of (47) mapped as described in (48). Putative RNA binding protein (RBP) binding sites here defined as regions with more than one PAR-CLIP read. P-value of enrichment was computed by permutation test against a shuffled set of positions within each genomic region. A plot of RBP binding site density in the vicinity...
of the m\(^5\)C sites was computed by averaging a 2000 bp window centred on all methylation sites.

RESULTS
A transcriptome-wide survey of m\(^5\)C candidate sites in human RNA

To develop a method for single nucleotide mapping of m\(^5\)C sites in a cellular transcriptome we adapted an RNA bisulfite conversion protocol, originally devised for primer extension-based detection of m\(^5\)C (40), for use with a sequencing-based readout (see ‘Materials and Methods’ section for details). As in DNA bisulfite sequencing, sites of m\(^5\)C in RNA will be read as cytosine in cDNA sequence, while unmodified cytosines will appear as thymidine. Although some other types of modified cytosine can also be resistant to bisulfite treatment (see ‘Discussion’ section), for simplicity we refer in the following to sites of non-conversion as ‘m\(^5\)C candidate sites’. We chose to analyse RNA preparations from HeLa cells (a human cervical cancer cell line) and used both positive and negative control RNAs to monitor success of the procedure. Our positive control was the endogenous tRNA\(^{\text{Asp(GUC)}}\), which harbours three previously identified m\(^5\)C sites at structural positions 38, 47 and 48 (7,31,32). Our negative control was \textit{in vitro} transcribed Renilla luciferase (R-Luc) mRNA lacking m\(^5\)C, which was spiked into the cellular RNA sample prior to bisulfite treatment. An aliquot of this RNA was used for conventional bisulfite sequencing reactions to examine the expected m\(^5\)C patterns in our controls. The R-Luc negative control RNA exhibited virtually complete cytosine conversion (Figure 1A; 99.8% conversion overall), while all three known m\(^5\)C sites in tRNA\(^{\text{Asp(GUC)}}\) selectively displayed low levels of conversion (Figure 1B; position 38 and 48: 0%, position 47: 12.5% conversion). These results showed that our RNA conversion protocol was efficient and accurate detection of m\(^5\)C sites is achieved.

We then performed SOLiD\(^\text{TM}\) next-generation sequencing of the converted RNA sample and obtained \~96 million sequence reads (50 nt in length). Approximately 41 million reads could be mapped to a custom human transcriptome reference (described in the ‘Materials and Methods’ section) using the SOCS-B program (45) and allowing up to four colour-space mismatches without counting those resulting from a C to T sequence change (indicative of bisulfite conversion). We next analysed the coverage and extent of conversion at cytosine residues in our controls, setting successively more stringent cut-offs, from four to zero-mismatch (4–0 mm) mapping data. Our controls had sufficient coverage across cytosines and displayed the expected C to T conversion patterns (Figure 1C and D show results for 2 mm data; see also Table 1 and Supplementary Table S4). R-Luc RNA showed almost complete C to T conversion overall (e.g. 99.5% at 2 mm; Figure 1C), and no individual position showed less than 90% conversion at a coverage threshold of 10 irrespective of the number of mismatches allowed. The three known m\(^5\)C sites in tRNA\(^{\text{Asp(GUC)}}\) showed low levels of conversion; this was consistent at all stringencies of mapping (e.g. 17.9% at 2 mm; average across all three

![Figure 1. Single-nucleotide resolution mapping of m\(^5\)C candidate sites in RNA. HeLa cell RNA preparations were spiked with a trace amount of \textit{in vitro} transcribed R-Luc RNA and bisulfite-converted as detailed in the ‘Materials and Methods’ section. (A) Negative control R-Luc and (B) endogenous tRNA\(^{\text{Asp(GUC)}}\) as a positive control were first analyzed by conventional sequencing to establish the efficacy of bisulfite conversion (top panels; columns signify cytosine positions along the RNA sequence, rows represent individually sequenced alleles, open boxes indicate cytosine to uracil conversion read as thymidine in cDNA and filled boxes indicate a retained cytosine). Numbers below refer to cytosine positions in the primary RNA sequence. Nucleotide positions highlighted in red designate previously identified m\(^5\)C sites in tRNA\(^{\text{Asp(GUC)}}\). Dual axis charts (bottom panels) display next-generation sequencing data mapped at 2 mm for the same control RNAs. Blue bars represent bisulfite-induced cytosine conversion, while red lines represent read coverage across individual residues. Top and bottom panels are aligned to each other by interrogated cytosine residues.](https://academic.oup.com/nar/article-abstract/40/11/5023/2409239)
sites; Figure 1D). These data indicate that our combination of bisulfite conversion with next-generation sequencing can accurately detect m\(^5\)C sites in RNA, and importantly, has a low false-positive rate.

**Discovery of novel m\(^5\)C candidate sites**

We next examined all reads mapping to cytoplasmic and mitochondrial tRNA sequences. There are 28 known m\(^5\)C sites in human tRNAs (43) and 27 of these were detectable in our dataset with at least one read spanning the site (Supplementary Dataset S1, Supplementary Table S5). Confirming our approach, we called the large majority of these sites as modified when applying a C to T conversion threshold of \(80\%\) and read coverage of at least \(10\) (shaded area). (B) Dependence of the proportion of novel tRNA candidate sites in anticipated tRNA structural positions (red circles in tRNA cloverleaf cartoon) on chosen conversion cut-off. Read coverage threshold was \(\geq 10\). Colour code for dots and lines refers to mapping at different colour-space mismatch limits.

In total, we detected 255 modified cytosine candidate sites in tRNA sequences using the above threshold criteria (Table 1 and Supplementary Dataset S2). Of these sites, 21 were previously known as m\(^5\)C in human tRNAs, for 51 cases other human isodecoder tRNA variants were known to harbour m\(^5\)C at those sites, 68 sites were reported as methylated in other animal tRNA orthologs, while 115 sites were entirely novel before this study.

**Figure 2.** Defining parameters for m\(^5\)C candidate site selection using tRNA data. (A) Plot of conversion against read coverage at 28 known m\(^5\)C sites in human tRNAs (43). The majority of the previously identified tRNA m\(^5\)C sites had a conversion of \(80\%\) or less and read coverage of at least \(10\) (shaded area). (B) Dependence of the proportion of novel tRNA candidate sites in anticipated tRNA structural positions (red circles in tRNA cloverleaf cartoon) on chosen conversion cut-off. Read coverage threshold was \(\geq 10\). Colour code for dots and lines refers to mapping at different colour-space mismatch limits.

| RNA type           | # of Cytosine | # of Cytosine with ≥10 read coverage | # of m\(^5\)C (≤80% conversion rate) | % m\(^5\)C |
|--------------------|---------------|-------------------------------------|--------------------------------------|-----------|
| R-Luc              | 255           | 232                                 | 0                                    | 0.0       |
| tRNA               | 14 584        | 2943                                | 255                                  | 8.7       |
| mRNA               | 16785 229     | 2247 702                            | 8495                                 | 0.4       |
| Other non-coding RNA | 5783 482     | 1447 999                            | 1780                                 | 1.2       |

Table 1. Number of cytosine residues identified as m\(^5\)C in different RNA categories in 2 mm data

Within rRNA sequences, we clearly detected two previously known m\(^3\)C sites at positions 3782 and 4447 in human 28S rRNA (7,32). We independently verified these sites as well as a site within the decoding centre of 12S mitochondrial rRNA (position 841; Supplementary Figure S2) that was previously identified as m\(^3\)C in hamster mitochondrial 12S rRNA (49,50). However, we also saw several prominent clusters of non-conversion (e.g. clusters of 10 or more sites in a 50 nt window, Supplementary Dataset S2), particularly in the cytoplasmic and mitochondrial large subunit rRNAs, which are among the longest, highly structured cellular RNAs.
These are likely due to incomplete denaturation of some highly structured regions of RNAs, thus we did not consider further our mapping data for rRNAs. Only ~7% of the m^C sites seen in other cellular RNAs occurred in such clusters, indicating a low proportion of false positive m^C predictions due to RNA structure in our data.

We then applied our selection criteria to identify novel m^C candidate sites throughout the transcriptome. To simplify the interpretation of results, we transferred the cytosine mapping information for the remaining RNA types to genomic co-ordinates and calculated cytosine conversion and coverage for each genomic locus. Approximately 2.4 million cytosine positions in the genome corresponding to known transcripts were covered by at least 10 reads, representing 10.6% of the total cytosines in known transcripts. Of these, 10275 sites (0.43% of all assessed cytosines) were identified with ≤80% conversion rate; 1863 sites remained when applying the highly stringent requirement of ≤50% conversion rate. Genomic coordinates, extent of conversion and additional information on all identified sites are listed in Supplementary Dataset S2. About 1780 of these sites were present in a range of non-coding RNA types, including lincRNA, pseudogenes and processed pseudogenes, while 8495 sites were located in mRNA sequences (Table 1).

Validation of m^C candidate sites by conventional bisulfite sequencing

To validate our global site mapping data, we verified three candidate sites from mRNAs, eight from tRNAs, and two from other non-coding RNAs by conventional bisulfite sequencing (Figure 3 and Supplementary Figures S2 and S3). Among the selected sites there were six that had been previously identified as m^C in other animal homologs (tRNA^Gly(UCC), 3 sites; tRNA^Lys(CU)U, 1 site; mitochondrial tRNA^Glu(UUC), 1 site; 12S mitochondrial rRNA, 1 site). The three sites in mitochondrial tRNA^Ser(GCU) were entirely novel, as those were chosen in protein-coding mRNAs (cyclin-dependent kinase 2 interacting protein, CINP; nicotinate phosphoribosyltransferase domain containing 1, NAPRT1; NADH dehydrogenase 1 beta subcomplex 7 mRNA, NDUF7B; 1 site each) and a non-coding RNA (RPPH1, Ribonuclease P RNA component H1). To assess potential non-conversion due to local RNA structure, we designed and prepared in vitro transcribed negative control transcripts mimicking the sequence context of five of the sites to be verified. These negative controls were spiked into independently prepared HeLa cell RNA prior to bisulfite treatment. None of these negative controls showed any evidence of non-conversion at the test sites, whereas all sites except position 47 in tRNA^Gly(UCC) were convincingly confirmed (Figure 3 and Supplementary Figure S3). This demonstrated that most of the novel m^C candidate sites predicted by our next-generation sequencing-based mapping are independently verifiable.

Identification of NSUN2 and TRDMT1 dependent m^C sites

To gain insight into the enzymes mediating RNA methylation, we performed RNAi-mediated knockdown of the RNA methyltransferases TRDMT1 and NSUN2 by

Figure 3. Validation of novel m^C candidate sites. Conventional bisulfite sequencing data is shown for three novel sites, (A) residue 48 in tRNA^Lys(CU)U, (B) residue 174 in the RNase P RNA component H1 (RPPH1), and (C) residue 748 in cyclin-dependent kinase 2 interacting protein mRNA (CINP). Top panels display results for endogenous transcripts. Data for spiked-in in vitro transcribed negative controls harboring the same sequence flanked by unique priming sites are also shown (middle panels) as are corresponding next-generation sequencing results (lower panels). Numbering of cytosine positions is as described in Figure 1, positions highlighted in red designate m^C sites identified by next-generation sequencing. See Supplementary Figure 3 for additional validation data.
transient transfection of HeLa cells. Knockdown of the DNA methyltransferase DNMT1 was used as a negative control, with knockdown efficiency determined by quantitative RT–PCR (Supplementary Figure S4). Conventional bisulphite sequencing of target regions in tRNA<sub>Asp(GUC)</sub> and tRNA<sub>Aeu(CAA)</sub> revealed marked disappearance of known target m<sup>5</sup>C sites for TRDMT1 (residue 38 in tRNA<sub>Asp(GUC)</sub>) and NSUN2 (residue 34 in tRNA<sub>Aeu(CAA)</sub>) in the respective knockdown cells (Figure 4A and B). Furthermore, methylation of residues 47 and 48 in tRNA<sub>Asp(GUC)</sub> was also shown to disappear with NSUN2 knockdown. Novel sites in CINP and NAPRT1 mRNAs, and in the non-coding RPPH1, were all clearly NSUN2-dependent (Figure 4C–E). Two previously identified m<sup>5</sup>C sites in 28S rRNA, as well as two novel candidate sites in 12S mitochondrial rRNA and tRNA<sub>Lys(CUU)</sub> did not respond to either enzyme knockdown (Supplementary Figure S2). Modulations at these sites may be placed by other MTases, or they may correspond to other cytosine modifications that could protect against bisulphite conversion (see ‘Discussion’ section); most likely these sites reside in RNA species that are not turned over rapidly enough to be amenable to assay in short-term knockdown experiments. Of the 11 sites tested, we found that six were dependent on NSUN2 and one was dependent on TRDMT1, also corroborating the identity of the modification at these sites as m<sup>5</sup>C. This further implies a major role for NSUN2 in modifying the human transcriptome, increasing the number of its known target sites from one to six, and extending its substrate range to an additional tRNA as well as a non-coding RNA and two mRNAs.

Analysis of m<sup>5</sup>C location bias within the transcriptome

To discover the wider role of m<sup>5</sup>C in the human transcriptome, we searched our data for any bias in the location of m<sup>5</sup>C candidate sites. To this end, we considered a subset of 9177 sites residing in canonical transcripts (longest known cDNA coding sequence; 89% of all predicted sites). The distribution of m<sup>5</sup>C candidate sites varied across different RNA types (P-value <2.2 x 10<sup>-16</sup>, chi-squared test). Sites were significantly enriched in a variety of non-coding transcript types, including several pseudogene categories (P-value <2.2 x 10<sup>-16</sup>, binomial test), whereas they were significantly depleted in mRNA (P-value <2.2 x 10<sup>-16</sup>, binomial test) (Supplementary Table S6). Given that many non-coding RNAs are expected to be of low abundance, we deemed our dataset to be of insufficient coverage to further analyse the patterns of m<sup>5</sup>C distribution in these RNA types. By contrast, despite being relatively devoid of m<sup>5</sup>C, candidate sites in mRNAs constituted the great majority (~83%) identified in our screen. We therefore, searched for any bias in site distribution within mRNA sequences. This revealed a significant enrichment within the untranslated regions (both 5' and 3' UTRs), and a relative depletion within coding regions (P-value <2.2 x 10<sup>-16</sup>, binomial test) (Figure 5A). Gene ontology enrichment analysis using the elin method (51) of genes harbouring m<sup>5</sup>C candidate sites in different transcript regions (3'UTR, 5' UTR and CDS) did not show any statistically significant enrichment at an FDR of 0.05 (data not shown).

A complementary approach to look for functional links was to test for any spatial association between m<sup>5</sup>C candidate sites and cis-acting regulatory motifs within RNA. Thus, we overlaid our modification mapping data with a set of recently identified human regulatory RNA elements within 3' UTRs (46). This showed a moderate but statistically significant enrichment of those elements in the vicinity of m<sup>5</sup>C sites (1.42-fold; P-value = 0.04, binomial test; data not shown). Furthermore, we overlaid our mapping data with publicly available transcriptome-wide maps of binding sites for major regulatory RNA-binding proteins in human cells generated by PAR-CLIP-Seq (47). This showed a substantial association of m<sup>5</sup>C candidate sites in mRNA with binding regions for the Argonaute I-IV proteins (Figure 5B), the central components of the miRNA/RISC complex (~1.76-fold over 3'UTR; P-value <1 x 10<sup>-3</sup>, permutation test) but, interestingly, not for binding sites of another post-transcriptional regulator, the Pumilio 2 protein (1.15 fold; P-value = 0.13). We also inspected the local sequence context for candidate sites and saw an increased frequency of C and G as flanking bases and a depletion of U either side of the modified cytosine, but no strict requirement for a targeting context was evident (data not shown). While more parsimonious analyses of subclasses of m<sup>5</sup>C candidate sites may yield further insight, it appears that m<sup>5</sup>C context requirements differ from those for m<sup>6</sup>A where a clear sequence context was reported [e.g. Gm<sup>6</sup>A in yeast (52)]. The enrichment patterns detailed above persisted when the analyses were done with more stringently selected candidate sites (i.e. ≤50% conversion rate), and altogether they are highly suggestive of an involvement of cytosine modification in post-transcriptional gene regulation.

DISCUSSION

Herein we described the successful combination of bisulphite treatment with next-generation sequencing of a cellular RNA library. This allowed us to survey the modification status of over two million cytosine sites in the transcriptome of the human HeLa cell line, which led to the mapping of over ten thousand novel candidate m<sup>5</sup>C sites in diverse types of cellular RNA (Table 1; Supplementary Dataset S2). The accuracy of our approach and site selection criteria was supported by multiple tests, including verification of a subset of known and novel sites by conventional bisulphite sequencing and m<sup>5</sup>C MTase knockdown.

Some cytosine modifications other than m<sup>5</sup>C may also be resistant to bisulphite treatment, including 3-methylcytidine, N4-methylcytidine (m<sup>4</sup>C), N4,2'-O-dimethylcytidine (m<sup>4</sup>Cm), and N4-acetylated variants. Schaefer et al. (32) were able to detect m<sup>4</sup>Cm at position 1402 of bacterial 16S rRNA by locus-specific bisulphite sequencing. The equivalent region in mitochondrial 12S rRNA from the hamster is methylated in the motif ‘Gm<sup>5</sup>CCm<sup>3</sup>CG’ (49,50). We surveyed the orthologous
Figure 4. Analysis of methyltransferase target sites. HeLa cells were transfected with siRNAs targeting **DNMT1**, **TRDMT1** and **NSUN2** or a non-targeting control siRNA (NTC) as indicated on the left. Conventional bisulfite sequencing data was obtained as described in Figure 1 and is shown for (A) tRNA<sup>Asp</sup>(GUC) (residue 38 is a known TRDMT1 target) and (B) tRNA<sup>Leu</sup>(CAA) (residue 34 is a known NSUN2 target) (C) **CINP** and (D) nicotinate phosphoribosyltransferase domain containing 1 (**NAPRT1**) mRNAs, and (E) **RPPH1** non-coding RNA. Nucleotide positions highlighted in red below designate m⁵C sites identified by next-generation sequencing. Green boxes indicate sites selectively responding to MTase depletion. Hatched boxes indicate intronic sequence. See Supplementary Figure S4 for data on RNAi knock-down efficiency and Supplementary Figure S2 for analysis of additional target sites.
region of the human counterpart by next generation and conventional sequencing and could only detect the m^4C site (Supplementary Figure S2). This may suggest that human mitochondrial 12S rRNA does not carry the m^4C modification or that m^4C and m^4Cm display differential sensitivity to bisulfite conversion. Perhaps, more likely, it reflects differences in treatment protocols as used by these colleagues and us. On balance, our evidence is consistent with m^5C as the modification underlying the majority of novel candidates sites reported here, although definitive proof for any given site would require application of more elaborate direct detection methods. Transcriptome-wide bisulfite sequencing of material from cells depleted for specific MTases would be a further tractable option to confirm modification identity on a global scale. Our current method also has limits with regard to surveying extremely structured regions such as those found in rRNA, which are among the longest, highly structured RNAs, so we excluded suspect regions of rRNA from further analysis. In addition, we included unmodified spike-in sequences in our verification efforts to control for effects of local structure (Figure 3), and verified sites through other means, such as MTase knockdown (Figure 4). While this work was in progress, Schaefer and colleagues used their locus-specific bisulfite sequencing approach to detect known m^5C sites in several tRNAs and rRNAs in different contexts (12,32,38), further underscoring the validity and scope of our transcriptome-wide method. Future efforts to improve the method will focus on optimization of RNA denaturation and conversion, while sufficiently preserving RNA integrity.

Our study detected 255 candidate sites for cytosine modification in tRNAs, most of them not previously mapped in humans. We found that most high-quality candidate sites were found at positions within tRNA secondary structure known to be occupied by m^4C in animals (43) (Figure 2), broadly confirming existing expectations of the role of m^4C in modulating tRNA function (8–13). Nevertheless, an emerging facet of tRNA biology is their processing into smaller regulatory RNAs species (53), and TRDMT1-mediated placement of m^5C has been shown to protect several tRNAs against stress-mediated cleavage in Drosophila (12). The present work thus provides a wealth of mapped m^5C sites that can now be assessed for a role in this phenomenon.

Human NSUN2 was thought to have a narrow substrate range, possibly confined to a single site within tRNA_{Leu(CAA)} (30). In contrast, our evidence demonstrates that its substrate range is likely to be much broader, as it is responsible for modification of further sites in tRNA_{Asp^GUC}, the RNA subunit RPPH1 of RNase P (54), and even mRNAs (Figure 4). The unexpected relationship between a tRNA-modifying and a tRNA-processing enzyme is intriguing and experiments will now have to address whether RPPH1 modification affects the enzymatic properties of RNase P. In support of a broader role of NSUN2 in mRNA modification it was found to be among the most highly enriched proteins in an in vivo capture screen for HeLa cell proteins binding to polyadenylated RNA (A Castello & MW Hentze, personal communication). NSUN2 is frequently overexpressed in a number of different tumour types and varies in its expression and intracellular localization (i.e. nucleolar versus cytoplasmic) during the cell cycle, while its enzymatic activity is regulated by phosphorylation (28,35,55). Importantly, it furthermore functions in balancing stem cell self-renewal and differentiation (36). Given the broad role for NSUN2 in modifying different RNA types we uncovered here, it is thus plausible that aspects of the transcriptome-wide RNA methylation patterns we have observed are dynamically responding to, and/or supportive of, distinct states of cellular growth, differentiation and transformation. Comparative studies of differential RNA methylation, particularly in the cancer and stem cell contexts, and in cells depleted of the candidate RNA MTases are now warranted.

A key outcome of this study was the clear demonstration of a widespread presence of m^5C within mRNAs. Pioneering work in the 1970’s had suggested the presence of m^5C, as well as N6-methyladenosine (m^6A) in mRNA, but their low abundance combined with a lack of suitable methods to map individual sites hampered investigation of their relevance. The
identification of the methyltransferase responsible for m^6A in mRNA subsequently allowed progress in its investigation (e.g. 52,56), but work on m^C in mRNA had all but seized since then. We detected candidate sites for cytosine modification in several thousand mRNAs, indicating that the presence of m^C is not limited to a highly specialized subset of mRNAs. Our analyses also indicate that m^C distribution within the transcriptome is not random, as we found the modification enriched within non-coding RNAs, whereas functional mRNAs were relatively devoid of it. Analyses of datasets with more complete mRNA coverage will be required to more precisely map the preferred locations of m^C within mRNAs, as they will likely hold clues to its molecular function(s). Our finding of an enrichment of m^C within the UTRs of mRNA and in the vicinity of Argonaute protein binding regions, already presents an intriguing pretext to further investigate a role of m^C in post-transcriptional gene regulation. Possibilities include, but are not limited to, a role in protecting mRNAs against innate antiviral defence mechanisms (26) or promoting their efficient translation, as has been suggested for m^6A (52).

The potential of RNA editing and modification to further diversify functions of cellular transcriptomes has recently come to the fore (57). We provided here a first global map of the patterns of m^C modification in the human transcriptome, which link it to mechanisms of post-transcriptional gene regulation and control of cell growth and differentiation. Our approach to the transcriptome-wide mapping of m^C can now be utilized to survey distribution patterns of this modification in different organisms and tissues, as well as to assess its potential for dynamic change under different cellular conditions.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1–6, Supplementary Figures 1–4, Supplementary Datasets 1–2.

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