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ABSTRACT

Here, we describe an approach to enrich newly transcribed RNAs from primary mouse neurons using 4-thiouridine (s4U) metabolic labeling and solid phase chemistry. This one-step enrichment procedure captures s4U-RNA by using highly efficient methane thiosulfonate (MTS) chemistry in an immobilized format. Like solution-based methods, this solid-phase enrichment can distinguish mature RNAs (mRNA) with differential stability, and can be used to reveal transient RNAs such as enhancer RNAs (eRNAs) and primary microRNAs (pri-miRNAs) from short metabolic labeling. Most importantly, the efficiency of this solid-phase chemistry made possible the first large scale measurements of RNA polymerase II (RNAPII) elongation rates in mouse cortical neurons. Thus, our approach provides the means to study regulation of RNA metabolism in specific tissue contexts as a means to better understand gene expression in vivo.

INTRODUCTION

Tissue-specific regulation of steady-state RNA levels is achieved through the precise control of RNA synthesis, processing, and degradation. These dynamics are critical for control of global transcript levels. The rate of RNA synthesis is regulated through changes in RNA polymerase II (RNAPII) initiation and elongation in response to signaling (1). RNAPII elongation rates have been reported to vary 4-fold for a range of transcripts in mammalian cell culture (2–5), and these rates have been proposed to be determined by the regulation of co-transcriptional splicing, termination, and RNA stability (6). Factors including chromatin context (such as the presence of H3K79me2 in the gene body) are also shown to affect elongation rates (7,8). In addition, differences in the cell-type specific chromatin state, trans-acting factors, and cellular metabolism suggest that elongation rates may differ between tissues (9–11), but these rates have not been routinely studied in primary cells or tissues. This is due to the fact that current experimental approaches have been limited by the scale required for biochemical enrichment (ca. 50 μg of input RNA is generally required for these protocols (12,13)). We therefore sought to develop more efficient biochemical capture of metabolically labeled RNA in order to understand RNA metabolism in primary cells using mouse cortical neurons.

Metabolic labeling allows the identification of newly synthesized RNAs by exposing cells to a non-canonical nucleoside that is incorporated during RNA synthesis. To identify newly synthesized RNA transcripts, total RNA is isolated and new, labeled RNAs are enriched. The three most widely used RNA metabolic labels are 5-bromouridine [BrU], 5-ethyluridine [5-EU] and 4-thiouridine [s4U]. Depending on the nucleoside, labeled RNAs can be captured with a specific antibody (BrU) (14,15), or biotinylated using either click chemistry (5-EU) (16), or thiol-specific reactivity (s4U) (12,17,18), respectively (reviewed in 19). Enriched RNAs can be analyzed by RNA sequencing (RNA-seq), quantitative PCR (qPCR), or microarray. Protocols based on s4U are particularly useful because s4U allows both covalent and reversible capture and is rapidly incorporated into the cellular NTP pool in living cells without the need for cell lysis and nuclear isolation. In addition, s4U shows minimal perturbation to cellular physiology at low concentrations even after long treatment (17,18,20). Therefore, s4U metabolic labeling can be employed to study the stability of both rare, transient RNAs (21,22) as well as stable transcripts (20) in the same experiment (23).

Transcriptional elongation can be studied using 4sUDRB-seq, which combines transcriptional synchronization with s4U labeling (3). Specifically, cells are treated with 5,6-dichloro-1-β-D-ribofuranosylbenzimidazole (DRB), which blocks the release of paused RNAPII into productive elongation. After DRB is removed from the media, the wave of new transcriptional elongation can be followed by s4U metabolic labeling of the new RNAs (3). This approach has been used to calculate rates of transcription genome-wide in HeLa cells (3), but existing methods to capture new RNA require prohibitively large amounts input RNA material to study new transcription in mouse.
cortical neurons. We therefore sought to use 4sUDRB-seq with an improved method to enrich s4U-RNA in order to determine RNAPII elongation rates in mouse cortical neurons.

We recently published efficient chemistry for the enrichment of s4U-RNA using the activated disulfide methane thiosulfonate (MTS), which improves yield and decreases biases in metabolic labeling experiments (18,24). We also identified two additional potential improvements that expand the scope of s4U-RNA enrichment methods: decrease the number of steps in the protocol by directly conjugating the MTS activated disulfide to a solid support, and use higher stringency rinses in the context of a fully cova-

cent of the RNA dynamics and metabolism.

MATERIALS AND METHODS

Cell lines and s4U metabolic labeling

K562 cells were cultured in RPM1 media supplemented with 10% (v/v) fetal bovine serum, and 1% (v/v) penicillin/streptomycin. HEK293T cells were cultured in high glucose DMEM media supplemented with 10% (v/v) fetal bovine serum, and 1% (v/v) 2 mM L-glutamine. Cultured cells were treated with s4U as described below and harvested at 80% confluence by the addition of 1 ml TRIzol. For TT-seq, K562 cells were treated with 1 mM s4U for 5 min. For s4U Chase-seq experiments, K562 cells were treated with 1 mM s4U for 2 h. s4U-containing media was exchanged with RPM1 media containing 20 mM uridine, and cells were harvested after 0.5 or 18 h of incubation. HEK293T cells were treated with 100 μM s4U for 2 h. Cells in TRIzol were flash frozen and stored overnight at −80°C.

Mouse cortical neurons and 4sUDRB

Mouse cortical neurons were isolated at E18 as previously described (25). About 2 × 10⁶ cells were plated per six-well dish. Two days post isolation, cells were treated with DRB for 3 h or with DMSO for the minus DRB control. 4sUDRB experiments were performed as previously described (26) with the following changes: (i) Cells were treated with 1 mM s4U for 30 min before being collected; (ii) cells were collected 0, 10, or 20 min after DRB removal. Cells were harvested with TRIzol, flash frozen and stored at −80°C.

Purification of total RNA

Cell lysates were chloroform extracted once and precipitated with one volume of isopropanol (supplemented with 100 μM DTT and 5–10 μg glycoblue), incubated for 10 min at room temperature and centrifuged at 20 000 × g for 20 min at 4°C. The pellet was washed with an equal volume of 75% ethanol. Purified RNA was dissolved in RNase-free water to a concentration of 200 ng/μl. Contaminating DNA was digested with Turbo DNase (1 U per 10 μg RNA, 37°C, 30 min.) Samples were purified by phenol:chloroform extraction followed by a second isopropanol precipitation and resuspension in 40 μl of RNase-free water. For sheared RNA samples, 2× fragmentation buffer (40 μl, 150 mM Tris pH 8.3, 225 mM KCl, 9 mM MgCl₂) was added to the RNA and incubated at 94°C for 4 min. Samples were immediately placed on ice and quenched with EDTA (20 μl 250 mM EDTA, final concentration 50 mM). Samples were cooled on ice (2 min) and were purified by the following modification of the RNeasy MinElute Cleanup Kit (Qiagen). 350 μl of buffer RLT and 250 μl of 100% EtOH were added to the RNA sample and placed over an RNeasy column. Columns were centrifuged for 15 s at 12 000 × g, 4°C, and flow through was discarded. Columns were washed with 500 μl buffer RPE supplemented with 35 μl of 1% β-me (final concentration 10 mM β-me) and centrifuged as above. Columns were washed with freshly prepared 80% EtOH and centrifuged for 2 min at 12 000 × g, 4°C, and flow through was discarded. Columns were dried by centrifugation at maximum speed for 5 min at 4°C. Samples were eluted into a fresh microfuge tube with 14 μl RNase-free water and centrifuged at maximum speed for 1 min at 4°C.

Synthesis of MTS-resin

For 10 samples, 100 μl NHS magnetic sepharose (GE Healthcare) was added to 1 ml RNase-free water, mixed well, and captured on a magnetic rack, discarding supernatant. One milliliter ice cold 1 mM HCl was added to beads, incubated 2 min with agitation, and beads were captured on a magnetic rack, discarding supernatant. Beads were washed twice with 1 ml 1 × PBS, then 100 μl MTSEA (Biotium, 10 mg/ml in PBS) plus 0.75 μl DIEA was added. Beads were incubated at room temperature for 15 min with rotation. Beads were washed twice with 1 ml 1 × PBS and blocked with 1 ml blocking buffer (1 × PBS, 5 μl acetic anhydride) for 15 min with rotation. Beads were washed twice with 1 ml 1 × PBS and twice with 1 ml quench buffer (1 M Tris pH 7.4, 200 mM ethanolamine). Beads were quenched with 1 ml quench buffer for 15 min with rotation. Beads were washed twice with 1 ml quench buffer and twice with 1 ml binding buffer (100 mM NaCl, 10 mM HEPES pH 7.4, 1 mM EDTA, 0.05% Tween). Beads can be stored at 4°C for several hours in binding buffer or aliquoted (10 μl/sample) and binding buffer removed to use immediately.

Purification of s4U-RNA with MTS-resin

1–5 μg total RNA in 10.5 μl RNase-free water was mixed with 1.5 μl 10 × binding buffer (1 M NaCl, 10 mM HEPES pH 7.4, 10 mM EDTA, 0.5% Tween), and 3 μl DMF, added to MTS-resin (10 μl aliquot), and incubated at room temperature in the dark for 2 h with rotation. Beads were captured on a magnetic rack and supernatant removed. Beads were incubated for 5 min with the following wash buffers:
once with 100 µl 1× binding buffer, twice with 100 µl high salt wash buffer (1 M NaCl, 100 mM Tris pH 7.4, 10 mM EDTA, 0.05% Tween), twice with 100 µl denaturing buffer (8 M guanidinium chloride) and three times with 100 µl buffer TE (10 mM Tris pH 7.4, 1 mM EDTA) at 55°C. RNA was eluted with 10 µl elution buffer (100 mM NaCl, 10 mM Tris pH 7.4, 1 mM EDTA, 10 mM DTT and 100 µg fragmented total RNA from S. pombe (a generous gift from the Berro lab, Yale University) and incubated at room temperature in the dark for 15 min with rotation. Eluent was ethanol precipitated with 5–10 µg glycoblu, and RNA concentration was assayed by Bioanalyzer RNA 6000 Pico Kit (Agilent) according to the manufacturer’s instructions.

MTS-resin binding capacity

A solution of 100 nmol s4U nucleoside in 18 µl RNase-free water was mixed with 3 µl 10× binding buffer and 6 µl DMF (total volume 30 µl) was enriched using 1, 2, 5, 10 or 20 µl MTS-resin as above. Enriched nucleoside was quantified by UV–vis spectroscopy based on A334 absorbance, and resin binding capacity was calculated using a standard curve of different concentrations of s4U.

MTS-resin saturation

1 µg of total RNA from K562 cells (prepared as above) was combined with 1 ng total RNA from S. pombe and enriched on 1, 2, 5, 10, 20 or 40 µl MTS-resin as above. Enriched RNA and 10% input were each reverse transcribed using the Superscript VILO cDNA synthesis kit (Invitrogen) and analyzed by qPCR using iQ Universal SYBR Green Supermix (Bio-Rad). Enrichment of s4U-RNA was analyzed using H. sapiens qPCR primers for CDKN1B total RNA (Forward: 5’-TTTGAAGCTGACATGAAGAGCTGTTGCAT-3’ and Reverse: 5’-AGCTGTTCTTGAAGGGACATTGAGAAGGGATGTTGGACCTGCTT-3’ (27)). Background from S. pombe RNA was analyzed using S. pombe qPCR primers for 28S rRNA (Forward: 5’-TGAGAGGGATGTGGACCTTGTCATTGAG-3’ and Reverse: 5’-ATTGCGTCACACCCACTTCTTGCGC-3’) (28).

Comparison between MTS biotin and MTS-resin enrichment

1 µg of total RNA from HEK293T cells was combined with 1 ng total RNA from S. pombe and enriched with MTS biotin exactly as in (29) or with MTS-resin as above. Enriched samples were assayed by RT-qPCR as above and fold enrichment was calculated as

\[ \frac{2^{\Delta \Delta C_{\text{Input}}}}{2^{\Delta \Delta C_{\text{MTS}}}} \]

An RNA ladder of 100–1000 nt was transcribed in vitro using the RNA Century Plus Marker Template and Maxiscript T7 transcription kit (Invitrogen) using Cy5-CTP at a ratio of 1:1 Cy5-CTP:CTP for downstream visualization, with the option of adding s4UTP (TriLink Biotechnologies) at a ratio of 1:1 s4UTP:UTP to the reaction. After the reaction, template DNA was digested with Turbo DNase (Thermo Fisher). Enzymes were removed by phenol-chloroform extraction and the RNA was purified using the RNeasy Mini Kit (QIAGEN) with the following changes: β-mercaptoethanol was added to wash buffer RPE to a final concentration of 1%. 400 ng of RNA ladder was enriched with MTS-resin following the protocol described above, or with HPDP-biotin and MTS-biotin following protocols in (18). Enriched samples were separated on a 5% urea-PAGE gel, visualized by Typhoon fluorescence imager (GE), and bands were quantified using ImageJ software.

Library preparation and sequencing

10 ng input as well as enriched RNA were prepared using the SMARTer Stranded Total RNA-seq Pico Input Mammalian Kit (Clontech) according to the manufacturer’s instructions. Samples were multiplexed with Illumina TruSeq i7 barcodes, and sequencing was performed at the Yale Center for Genomic Analysis on Illumina HiSeq 2500 instruments with paired-end 2 × 75 nt sequencing runs.

Bioinformatic analysis

Sequencing reads were aligned using STAR (version 2.4.2a) (30) to a joint index of the H. sapiens and S. pombe genomes (GRCh38 and sp2) and transcriptomes (NCBI and Ensembl Fungi v22; respective) or to the M. musculus genome and transcriptome (GRCm38, NCBI) (31,32). Alignments and analysis were performed on the Yale High Performance Computing clusters. Following alignment, HTSeq-count (version 0.6.1p1) (33) was used to quantify annotated H. sapiens and S. pombe transcripts for total RNA (+t gene) and mRNA (+t exon). Tracks normalized using the S. pombe reads were uploaded to the UCSC genome browser. Pearson correlations between samples were visualized in R using the corrplot package. GO-enrichment analysis was performed using the PANTHER classification system (34) using the statistical overrepresentation test and GO molecular function—complete gene list. TT-seq data from Schwalb et al. (GEO series accession GSE75792) were randomly downsampled for the same read depth as MTS-TT-seq data and processed as above. Counts were filtered to remove mitochondrial and ribosomal RNAs. PRO-seq data were downloaded from GEO (series accession GSM1480327), and ChIP data for RNAPII, H3K36me3 and H3K4me1 were downloaded from the ENCODE ChIP data matrix. Enhancer RNAs were quantified using BEDtools coverage (50% minimum read overlap) of the Chromatin State Segmentation by HMM from ENCODE/Broad (35) and filtered for strong enhancers. MicroRNA transcription start sites were assigned using the mirSTP pipeline (36) and relaxed annotations are reported.

RNAPII elongation rates were calculated using the scripts published by Fuchs et al. (3), with the following modifications: (i) Due to the lower sequencing depth in our samples, the bin size was changed from 100 to 500 bp. (ii) Because the time of s4U metabolic labeling was 10 and 20 min, rather than 4 and 8 min published by Fuchs et al., we expanded our x-intercept filtering to –2.5 min to 10 min, rather than –1 to 4 min. (iii) We calculated rates for the most abundant isoform of each transcript. To expand our analysis to more genes, we also calculated RNAPII elongation rates from 0 to 10 min, assuming a linear fit (e.g. a constant rate of transcription). Genes were filtered for expression (RPK > 1), active RNAPII elongation over 20 min
following DRB washout (average transcriptional boundary at 20 min is greater than the average transcriptional boundary at 10 min), agreement of two replicates (transcriptional boundary of replicate 1 within 70% of replicate 2). In addition, we only considered genes were the transcriptional boundary of one transcript did not overlap another gene.

H3K79me2 and H3K36me3 ChIP data from mouse cortical neurons (GEO series accession GSE95831 and GSE103214 (37), respectively) were re-processed as follows: reads were aligned to the mm10 genome using Bowtie2 (38) and \( \log_2 \) fold enrichment over input was calculated using MACS2 (39). Mean fold enrichment was calculated over the first 10 kb from the TSS.

RESULTS

Resin synthesis and characterization

Developing solid-phase chemistry to capture \( s^4U \)-RNA requires the use of a highly activated disulfide, as the loading of the resin limits the extent to which mass-action can be used to drive the chemical reaction. Previously, we demonstrated that methane thiosulfonate (MTS) activated disulfides react more efficiently with \( s^4U \) than the commonly used HPDP-biotin, and that this improved chemistry enables more sensitive applications of \( s^4U \) metabolic labeling (18). We hypothesized that coupling the MTS moiety directly to magnetic sepharose beads should decrease the loss that occurs during the multiple steps of a biotin/streptavidin purification and increase fold enrichment of \( s^4U \)-RNA by enabling higher stringency rinses. After \( s^4U \)-RNAs are eluted by reducing the disulfide bond, these RNAs could be analyzed by microarray, qPCR, or RNA-seq. To test this approach, we coupled methane thiosulfonate (MTS) to a solid support (Figure 1) by reacting methane thiosulfonate ethyamine (MTSEA) with commercially available N-hydroxysuccinimide (NHS) activated magnetic beads using a modification of standard protocols (Supplementary Figure S1A, Materials and Methods). We found these beads are capable of capturing \( s^4U \) nucleosides as well as metabolically labeled \( s^4U \)-RNA (Figure 1B and C).

To determine the binding capacity of MTS-resin, we incubated beads with a vast molar excess of \( s^4U \) nucleoside to saturate the available sites on the resin. After rinsing and elution, we quantified the amount of the captured nucleoside. Binding was linear with increasing quantity of resin and the resin could capture 0.4 nmol \( s^4U \) per \( \mu l \) MTS-resin slurry (Figure 1C). While this loading capacity provides an upper bound of the amount of \( s^4U \)-RNA that can be captured, other factors such as site accessibility could provide practical limits to the resin loading in a biological sample. Therefore, we used metabolically labeled RNA from K562 cells (2 h, 1 mM \( s^4U \)) to test the relationship between amounts of MTS-resin slurry used and the amount of \( s^4U \)-RNA retrieved (Figure 1D). We observed an increase in \( s^4U \)-RNA enrichment with increasing amounts of MTS-resin (plateauing at 10 \( \mu l \) resin per \( \mu g \) of input RNA). To measure nonspecific background, we also added exogenous unlabeled RNA from \( S. \) pombe (1 ng unlabeled standard per 1 \( \mu g \) input labeled RNA) and found that this background did not significantly increase even at the highest amounts of MTS-resin that we tested. While background did not increase, we did observe some bias for enrichment of longer RNAs (with more \( s^4U \)) when using the MTS-resin that is not observed with MTS-biotin (Supplementary Figure S1B–D). This type of length bias has been observed previously (40) and can be mitigated by pre-shearing RNA samples before enrichment (22).
Next, we compared the sensitivity of this small-scale s⁴U-RNA enrichment (1 μg input RNA) on MTS-resin to our previously published protocols using MTS-biotin (18,29) and tested the effect of stringent rinses on both purifications (Figure 1E). We measured the retrieval of a metabolically labeled mRNA (CDKN1B) in comparison with an unlabeled RNA from S. pombe. While stringent rinses improve the fold enrichment in MTS biotin experiments (13-fold enrichment with standard rinses, 340-fold with stringent rinses), the enrichment was significantly increased with MTS-resin (1500-fold enrichment with standard rinses, 5200-fold with stringent rinses). In addition, while the stringent rinses decrease s⁴U-RNA yield in MTS biotin purification (33% input with standard rinses, 19% input with stringent rinses), we did not detect a significant difference in yield with MTS-resin enrichment stringent rinses (53% input with standard rinses, 55% input with stringent rinses). These results demonstrate that high levels of enrichment can be achieved when using an entirely covalent purification, which was made possible by the MTS-resin.

4-Thiouridine pulse-chase labeling (s⁴U Chase-seq)

After we synthesized MTS-resin and validated that it can capture s⁴U-RNA, we next tested if the MTS-resin could be used to distinguish fast turnover RNA populations from slower turnover populations, a common application of s⁴U metabolic labeling experiments (12,17,40,41). We chose to perform a pulse-chase experiment similar to Yi et al. (20), where the cells are exposed to a s⁴U pulse (1 mM, 2 h) followed by a chase phase where the cells were treated with 20 mM uridine (Figure 2A). Those RNAs that were labeled with s⁴U at 0.5h were compared with the population of s⁴U-RNA that remained after 18 h to detect long-lived RNAs. For applications such as measuring transcription elongation rates in primary cells, s⁴U-RNA capture method must perform well at small scale and also have the sensitivity to capture rare, newly made RNAs from the higher concentrations of pre-existing RNA pool. We tested MTS-resin in the context of a transient transcriptome sequencing (TT-seq) experiment (Figure 3A), which uses very short s⁴U labeling (5 min, 1 mM) followed by RNA fragmentation and enrichment. This approach captures rare RNAs including introns, enhancer RNAs (eRNAs), and pri-miRNAs (22). Using MTS-resin, we performed this experiment using only 2.5 μg of input RNA, substantially below (>20-fold) the scale that is generally used for these experiments (22,48). Enriched RNA samples, as well as input RNA, were analyzed by high-throughput sequencing and reads were mapped to the human genome. Enrichment was consistent across biological replicates (Pearson’s r = 0.99). As we would expect if the MTS-resin successfully enriched the transient RNA population, the MTS-TT-seq samples correlate better with each other than with s⁴U controls or input, suggesting that the resin is capturing new transcripts over nonspecific background (Figure 3B, Supplementary Figure S3B). Consistent with this conclusion, MTS-TT-seq samples correlated well with previously published TT-seq data (Pearson’s r = 0.89, Supplementary Figure S3A and B). Notably, we found that MTS-TT-seq enriches transient RNA species including pre-mRNA (as revealed by intronic RNA enrichment), eRNAs, and primary microRNAs (pri-miRNAs) relative to input (Figure 3C and D), consistent with previous data (22). We conclude that MTS-resin is able to enrich rare RNA pop-
Figure 2. s^4U Chase-seq identifies fast- and slow-turnover RNAs in K562 cells. (A) Schematic of s^4U Chase-seq. K562 cells were metabolically labeled with 1 mM s^4U for 2 h, followed by a 20 mM uridine chase for 0.5 h or 18 h. (B) Example genome browser view of s^4U-RNA enriched after 0.5 h and 18 h uridine chase. FOS and HSP90AA1 were identified as fast and slow turnover, respectively, by differential expression analysis. Reads from biological triplicates were summed for display. (C) Scatterplot of fold change versus normalized expression based on comparative analysis of RNAs that are significantly enriched or depleted in early time points (0.5 h chase) relative to late time points (18 h chase). Fast-turnover RNAs (fold difference > 2; $P < 2 \times 10^{-5}$) are colored red; slow-turnover RNAs (fold difference < 0.5; $P < 2 \times 10^{-5}$) are shown in blue. (D) Scatterplot of log₂ fold change from (C) versus log₂ RNA half-life from Schofield et al. (E) Box and whisker plot of log₂ RNA half-life from Schofield et al. for RNAs binned as fast turnover, slow turnover, or not significant in (C). RNAs were filtered for normalized expression > 2 (logCPM).

Afforded by MTS-resin for TT-seq analysis led us to conclude that this approach will allow the study of transcriptional dynamics analyses in a wider variety of cell types and tissues than was previously feasible.

RNAPII elongation rates in mouse cortical neurons

Having established that MTS-resin is compatible with s^4U-RNA capture from small numbers of cells, we next applied the resin in the context of a 4sUDRB-seq experiment to study the transcriptional dynamics in primary neurons. We treated $\sim 2 \times 10^6$ neurons with DRB to synchronize RNAPII at the promoter-proximal pause site. Next, s^4U was added to the media (before DRB washout to allow s^4UTP to build up in the cellular pool) and the DRB was removed to allow the polymerase to elongate and incorporate s^4U into the newly synthesized RNA. Total RNA from cells harvested after 0, 10 or 20 min of DRB washout was purified, and s^4U-RNA was enriched on MTS-resin followed by high-throughput sequencing (Figure 4A and Supplementary Figure S4A).

Read counts mapping to transcripts were consistent between biological replicates (Pearson’s $r = 0.81–0.97$). As we would expect if 4sUDRB-seq is capturing the transcriptional wave of newly synthesized RNA, $t = 10$ min samples correlated better to $t = 20$ min (Pearson’s $r = 0.81–0.90$) than to RNA-seq (Pearson’s $r = 0.58–0.63$), $t = 0$ min (Pearson’s $r = 0.58–0.67$) or no treatment controls (Pearson’s $r =$
After filtering the RNAPII transcriptional boundary for RNA-seq expression and agreement between replicates (see Materials and Methods), the RNAPII elongation rate from 0 to 10 min after DRB washout was calculated to be much faster than the elongation rate from 10 to 20 min for most genes (Supplementary Figure S4E), which is consistent with previous reports that RNAPII reaches its maximum elongation speed approximately 15 kb through the gene (51). Therefore, we first calculated the elongation rate from 0 to 10 min following DRB washout, which is a timescale more consistent with previous experiments in cell culture (3,8).

We estimated the rates of RNAPII elongation from 0 to 10 min after DRB washout for 260 genes in mouse cortical neurons, where the median transcription elongation rate was 2.74 kb/min, and gene-specific elongation rates varied from 0.35 to 5.83 kb/min (Figure 4D, Supplementary Table S3).

We examined how the gene-specific rates of RNAPII elongation in mouse cortical neurons related to the presence of chromatin modifications that have been previously reported to correlate with RNAPII elongation rate in cell culture (3,8,51). We found H3K79me2 to be significantly enriched in the fastest quartile (Wilcoxon test, \( P = 0.0024 \)), whereas H3K36me3 did not show any significant enrichment (\( P = 0.22 \), Figure 4E). We did not find any significant correlation between RNAPII elongation rate and gene length, expression, or exon density (Figure 4E).

In an effort to quantify RNAPII elongation rates using transcriptional boundaries calculated after 10 and 20 min of DRB washout, we more stringently filtered our transcriptional boundary measurements using criteria established by Fuchs et al. (26). We limited our analysis to the most abundant isoforms of transcripts >50 kb in length where transcriptional boundary algorithms (see Materials and Methods) gave convergent values for both biological replicates in the \( t = 10 \) and \( t = 20 \) min samples. In addition, we filtered our elongation rates for an x-intercept between –2.5 min and 10 min, which is proportional to the longer times used in our experiments (10 and 20 min, compared to 4 and 8 min in (3)). Based on these conservative criteria, we were able to calculate the elongation rate for 55 genes in cortical neurons (Figure 4F, G, Supplementary Table S4). The s4U-RNAs that mapped to these regions show a similar transcriptional wave compared to the profile for all genes >50 kb (compare Supplementary Figure S4D and Figure 4C). We found that the median transcription elongation rate in mouse cortical neurons for these transcripts was 2.95 kb/min, which is similar to the median transcription elongation rate that Fuchs et al. calculated in HeLa cells (3.72 kb/min) (3). In addition, the RNAPII elongation rate varied more than 3-fold between genes (1.5 to >6 kb/min), which is consistent with the spread of RNAPII elongation rates from 0 to 10 min (Figure 4D), as well as previous findings in cell culture (2–5).

**DISCUSSION**

Here we show that MTS conjugated to beads can be used capture s4U metabolically labeled RNA. This one-step enrichment reduces handling loss that is inherent in other s4U enrichment protocols, and thereby makes it possible to use
Figure 4. MTS-resin reveals RNAPII elongation rates in mouse cortical neurons. (A) Scheme of 4sUDRB-seq using MTS-resin. NT: no DRB treatment (30 min s4U-seq); t = 0 min: no DRB washout. S4U is added to cells for 30 min, but DRB is not removed from cells; t = 10 min, t = 20 min: s4U-RNA is enriched with MTS-resin, followed by high-throughput sequencing. (B) 4sUDRB-seq enriched RNA from two representative genes, Rsf1 and Shc3. Arrows mark the direction of transcription. (C) Average distribution of reads in all genes longer than 50 kb from s4U-RNA enriched after 0, 10 or 20 min of DRB removal from two biological replicates. (D) Distribution of RNAPII elongation rates from 0 to 10 min following DRB washout, calculated from 260 genes in cortical neurons. (E) Box-whisker plot of log2 transformed H3K79me2 and H3K36me3 mean fold enrichment in the first 10 kb from the TSS, gene length (kb), gene expression (RPK), and exon density (% exon) of the 25% of genes with the highest calculated elongation rate and the 25% of genes with the lowest elongation rate. P-values were calculated by the Wilcoxon test. (F) Linear fit of transcriptional boundaries calculated after 10 and 20 min of DRB washout. The slope (v) represents the elongation rate in kb/min, with confidence intervals as indicated. (G) Distribution of RNAPII elongation rates calculated from 55 genes in cortical neurons using criteria established by Fuchs et al. (3).

In addition, we sought to analyze RNAPII transcription boundaries after both 10 and 20 min of DRB washout using the more stringent criteria established by Fuchs et al. (3) which focuses on rates that are linear during this time course. This limited set of 55 genes showed a similar median elongation rate to the larger set of RNAPII elongation rates during the first 10 min of DRB washout (2.74 versus 2.95 kb/min). Despite the differences in metabolic state of primary neurons compared to rapidly dividing HeLa cells, the median transcription rates are remarkably similar (3 kb/min in neurons versus 3.7 kb/min, Figure 4D). We found that these rates vary >3-fold across the set of genes we analyzed, which is consistent with rates reported in cell culture (2–5). Given the extensive differential processing of RNA in neurons in particular (54), the ability to measure RNAPII elongation rates in mouse cortical neurons provides the foundation for understanding the relationship between the control of RNA elongation and the generation of different neuronal RNA isoforms.

These results demonstrate how MTS-resin can be a useful tool for studying RNA population dynamics in samples where input RNA is limiting. We suggest that MTS-resin is particularly well suited for smaller-scale purification of s4U-RNA (e.g. primary tissues or microdissections). When large quantities of RNA are available, MTS-biotin has some advantages over MTS-resin. When using MTS-biotin, higher concentrations of the activated disulfide can be used to drive s4U-RNA modification, but an additional step is required to remove unreacted MTS-biotin. When using MTS-resin, we have observed variable amounts of length bias...
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DATA AVAILABILITY

Raw and processed RNA-seq datasets have been submitted to the NCBI Gene Expression Omnibus under the accession code GSE110951.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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