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Comprehensive comparative analysis of 5' end RNA sequencing methods

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Abstract

RNA-Seq is an effective method to study the transcriptome, but specialized methods are required to identify 5' ends of transcripts. Several published strategies exist for this specific purpose, but their relative merits have not been systematically analyzed. Here, we directly compare the performance of six such methods – testing five with cellular RNA as well as a novel spike-in RNA assay that helps address interpretation challenges that arise from uncertainties in annotation or RNA processing. Using a single human RNA sample, we constructed and sequenced 18 libraries with these methods and one standard, control RNA-Seq library. We find that the CAGE method performed best for mRNA and that most of its unannotated peaks are supported by evidence from other genomic methods. We then applied CAGE to eight brain-related samples and revealed sample-specific transcription start site (TSS) usage as well as a transcriptome-wide shift in TSS usage between fetal and adult brain.

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J.L., X.A., and A.R. conceived the research. X.A. prepared the 5' end RNA-Seq libraries. J.J. prepared the standard RNA-Seq library. X.S. prepared the *in vitro* neurons under the supervision of J.P. M.L. prepared the brain organoid RNA. A.H., S.S., A.L.M., Z.J., and M.B. developed and performed computational analysis. J.L., X.A., A.H., S.S., and A.R. wrote the paper. All authors assisted in editing the paper.

INTRODUCTION

Precise promoter annotation is central to addressing many questions in biology, including condition and tissue specific gene regulation, differential 5' untranslated region usage, and the impact of genetic variation in non-coding regions on gene expression. In particular, as Genome Wide Association Studies and sequencing studies identify thousands of loci associated with human diseases in non-coding regions, the challenge is to relate genetic variants to their mechanism of action^{1, 2}. A critical step for understanding the functional impact of such genetic polymorphisms is correctly identifying transcription start sites (TSSs). For example, a single nucleotide polymorphism in a regulatory region was shown to create a new TSS that interferes with normal activation of downstream alpha-like globin genes, thereby causing thalassemia³. Additionally, identifying multiple TSSs for a gene and understanding their usage in the relevant tissues can help design follow up experiments. Further, in many cases differential TSS usage is important for gene function and in human disease⁴, such as TP73, where transcription from one promoter leads to a protein acting as a tumor suppressor and the other a protein acting as an oncogene⁵ and NRXN1 in which mutations found in neurodevelopmental disorder patients have differing symptoms that may reflect the disruption of the alpha and beta promoters⁶.

While transcriptome analysis by RNA-Seq is a powerful approach for gene expression measurements, novel transcript discovery, and splice-isoform determination⁷, it is still often difficult to reliably identify more than one TSS per gene in a given transcript isoform. Empirical determination of the correct TSS in a given sample is particularly important in complex transcriptomes, such as human, where 54% of genes are currently annotated as having multiple TSSs⁸. Several methods have been proposed for the identification of the 5' end of transcripts, including CAGE⁹, RAMPAGE^{10, 11}, STRT¹², NanoCAGE^{13, 14}, Oligo Capping (also known as TSS-Seq)^{15, 16}, and GRO-cap¹⁷ (also known as 5' GRO-Seq¹⁸) (Fig. 1), but their relative merits have not yet been systematically compared¹⁹. Even for a widely accepted method such as CAGE, there are many reads aligning to 3' rather than 5' ends of transcripts²⁰, so that further investigation could be beneficial.

Here, we compare six 5' RNA-Seq methods using a comprehensive set of metrics. Starting from total RNA from one human cell line, we constructed a set of libraries for five of the methods, as well as a control library with standard RNA-Seq, and deeply sequenced them. We identify the CAGE method as performing best for mRNA and show that most of its unannotated TSS peaks also have corroborative evidence to support their being bona fide TSSs. For enhancer RNAs (eRNAs), we find GRO-cap identifies many more transcripts than the other methods. We then used CAGE to generate TSS data for eight brain-related samples, identifying many examples of differential promoter usage, and showing evidence for a novel, genome-wide trend of differential TSS usage, where downstream TSSs are preferentially used in adult brain and upstream TSSs are used in fetal brain and *in vitro* differentiated neurons. Our evaluation strategy, results, and brain TSS catalog can serve as resources for the community.

RESULTS

A comparison of 5' RNA-Seg methods

We tested five methods for preparing RNA-Seq libraries that identify the 5' end of transcripts (Fig. 1). We attempted to optimize each method to facilitate efficient library construction and sequencing of indexed libraries on an Illumina sequencing platform (Online Methods). To make a comprehensive comparison, we tested each method by using RNA from the human cell line, K-562, to construct and sequence 18 libraries (Supplementary Table 1). The methods vary in their input RNA requirements, with STRT, which was developed as an ultra-low input method, requiring the least RNA and Oligo Capping requiring the most (Online Methods). We varied the RNA input by method specifications, but aimed to use the lowest recommended amount, recognizing that RNA quantity can be limiting in practice. To assess whether the lower RNA input amount for STRT compared to the other methods affected its performance, we constructed and sequenced an additional set of eight STRT libraries with input amounts ranging from 10 ng to 10 ug (Supplementary Table 1). In addition, we compared published data for K-562 prepared with GRO-cap¹⁷. We constructed one control library using standard RNA-Seq with ribosomal RNA (rRNA) depletion by the RNase H method^{19, 21} to understand the value of using one of these more specialized RNA-Seq methods.

We first assessed performance of each method by standard RNA-Seq metrics (Supplementary Table 1). CAGE and NanoCAGE-XL produced fewer reads per library because of limited quantities of library DNA for CAGE and poor sequencing yield for NanoCAGE-XL. All methods showed acceptable levels of reads aligning to rRNA (<20% of reads) and excellent strand specificity (>99% correct strand reads) with the exception of GRO-cap (90% correct strand reads). GRO-cap also had a much higher fraction of reads aligning to introns (20%) and intergenic regions (47%) than the other 5' end RNA-Seq methods.

For an initial assessment of specificity for 5' ends, we examined coverage of reads from 5' to 3' (Fig. 2a,b). When considering only reads that aligned to exonic regions, we observed GRO-Cap performed best, followed by CAGE and RAMPAGE and NanoCAGE-XL performed worst (Fig. 2a). Analyzing reads aligning to the entire gene including introns, we observed that RAMPAGE and CAGE performed best followed by GRO-cap (Fig. 2b) – likely reflecting the high fraction of GRO-cap reads aligning to introns (Supplementary Table 1). This global analysis is congruent with our observations for representative highly-expressed genes (Supplementary Fig. 1). Overall, even the best performing method, CAGE, has a sizeable fraction of reads (24% of reads for the average gene) mapping to regions not close to transcript 5' ends (farther away than 10% of the length of the transcript) by this measure (Fig. 2a).

Assessment of 5'-end specificity with synthetic spike-in RNAs

We next explored why many reads were not aligned at the annotated 5' end of transcripts (Fig. 2a). Such non-5' end reads could reflect either technical limitations, incomplete annotation, or biological explanations like RNA re-capping²². To focus on technical

performance, we developed a spike-in RNA assay using ERCC transcripts 23 (Online Methods).

The methods showed similar relative performance based on 5' end coverage of these artificial transcripts (Fig. 2c and Supplementary Table 2) as with cellular RNA (Fig. 2a), with CAGE and RAMPAGE performing best. STRT performance did not improve with increased input amounts (Supplementary Table 2). The 5' end specificity differs among the spike-in transcripts – indicating that there is some variability in method performance for different transcripts.

We also assessed how accurately these methods quantitated the relative fraction of each ERCC spike-in transcript. This is important for quantifying differences in TSS usage across samples. For each transcript, we compared the relative input amount to the fraction of reads aligned (Supplementary Table 3). We assessed the uniformity of read coverage using the mean quantitation error (Supplementary Table 4). The CAGE libraries had the lowest error (1.1%) and performed better than RAMPAGE (2.0%).

TSS peak calling to help assess 5' specificity

To better compare the 5' specificity of methods for cellular transcripts, we used computational peak calling to identify TSS locations. We reasoned that this would distinguish noisy background reads spread across the length of a gene from peaks of reads aligning to specific locations in that gene. Furthermore, many studies seek to identify TSS peaks and associate them with other genomic and biological information. To enable equivalent comparisons across methods, we randomly sampled 20 million aligned reads for each method, with two exceptions. First, NanoCAGE-XL did not have sufficient reads (Supplementary Table 1), so that we used all its aligned reads. Second, we did not have enough reads from any single CAGE library, and pooled replicate libraries for subsequent analysis. We called peaks from aligned reads using Paraclu²⁴, with parameters optimized for best sensitivity and precision for each library (Online Methods). We also incorporated two additional filtering steps to ensure that our peak calling results accurately represented lab method performance (Supplementary Note 1, Supplementary Fig. 2-4).

CAGE performs best in TSS peak-based comparison

We evaluated each of the six methods for their ability to identify TSSs relative to known annotation. We used the UCSC transcriptome annotation⁸, which does not rely on data gathered with 5' end RNA-Seq methods, and as such should not be biased towards any of the methods *a priori*. For precision, CAGE, STRT, and NanoCAGE-XL performed best (Fig. 3a), while for sensitivity GRO-cap was the best method followed by CAGE and RAMPAGE (Fig. 3a). Combining both considerations, CAGE performed best followed by RAMPAGE, GRO-cap, STRT, NanoCAGE-XL, and Oligo Capping (Fig. 3a,b). STRT performance did not improve with increased input amounts (Supplementary Fig. 5). The same rankings were obtained when using the Gencode annotation²⁵ (Supplementary Fig. 6). In addition, we analyzed published datasets in which two of these methods were tested on the same cell line or tissue in different laboratories. In three cases, we were able to compare CAGE and another method – CAGE²⁶⁻²⁸ performed best compared with RAMPAGE¹⁰, STRT²⁹, and

Oligo capping¹⁵ (Supplementary Fig. 7). We also compared the performance to 5' ends identified from standard RNA-Seq (Online Methods). Standard RNA-Seq had more peaks – leading to both more true and false positives – and a ROC curve that suggests a similar false positive to true positive ratio compared to NanoCAGE-XL, which identified the fewest peaks, but worse than all the other 5' end methods, except Oligo capping (Fig. 3b).

For each method, we also assessed TSSs at single base resolution (Supplementary Note 2, Supplementary Fig. 8), reproducibility and gene expression quantification accuracy (Supplementary Note 3, Supplementary Figs. 9,10, Supplementary Tables 2-4).

Evidence supporting unannotated TSS peaks

Given that CAGE performed best, we explored its performance further using evidence from other sources. DNase-seq and H3K4me3 ChIP-Seq data can be used to further understand which false positives (based on prior annotation) may actually be true positives. DNase-Seq³⁰ identifies genomic regions with open chromatin such as promoters, and H3K4me3 is associated with TSS regions of active promoters³¹. While such corroborative data cannot provide a definitive conclusion about which individual peaks are true positives, it adds valuable information about the confidence in each peak.

Indeed, when considering both epigenetic data, Gencode annotations and the consensus peaks identified by the other 5' end RNA-Seq methods, the vast majority of CAGE peaks have additional support (Fig. 4). First, ~55% of false positive and ~42% of intergenic peaks for CAGE have support from Gencode annotation and/or the consensus peaks identified by the other 5' end RNA-Seq methods (Fig. 4a). (We omitted NanoCAGE-XL from this consensus because it detected so few peaks (Fig. 3a,b).) Next, with corroborative evidence from DNase-Seq or H3K4me3 ChIP-Seq²⁸, the great majority of the remaining unannotated peaks have some evidence that they may be actual TSSs (Fig. 4a,b). Corroborative evidence for other methods was also analyzed (Supplementary Note 4, Supplementary Fig. 11)

Although TSSs are associated with promoters, they can also be found at the start of eRNAs and would be classified as "intergenic" in the annotation we used ^{32, 33}. For Paraclu-called peaks found in intergenic regions (Supplementary Fig. 11), we explored their locations relative to enhancer regions in the genome. Using three different approaches based on histone modifications and open chromatin to identify enhancer regions (Online Methods), we observed that GRO-cap had ~5,000 peaks in such regions compared to much fewer such peaks for the other methods (Supplementary Table 5). The percentage of intergenic peaks that were in enhancer regions showed the same trend as for peaks in annotated genes (Fig. 3) with most methods performing well, except for Oligo capping (Supplementary Table 5). Because eRNAs are expressed at lower levels than mRNAs and are expressed as divergent, non-overlapping transcript pairs, we took a second approach based on a previous study ³³ (Online Methods) to identifying peaks in such regions. We obtained similar results in that GRO-cap had the most TSSs in enhancer regions (Supplementary Table 6).

When considering whether CAGE data are sufficient to identify TSSs in a given sample, it is worth knowing whether collecting corroborative data such as DNase-Seq or H3K4me3 ChIP-Seq can improve the quantity or confidence of the identified TSSs. Figure 4c shows

how TSS prediction can be refined using such corroborative data together with CAGE data. More CAGE peaks would be filtered out by requiring DNase-Seq than H3K4me3 ChIP-Seq evidence (Fig. 4c). For standard RNA-Seq without CAGE data, the corroborative evidence is even more important in identifying "true positive" 5' ends, based on annotation, with DNase-Seq evidence again having a bigger impact than H3K4me3 ChIP-Seq (Supplementary Fig. 12).

Differential TSS usage in brain-related samples

Having identified CAGE as best, we applied it to a set of eight brain-related samples to explore differential TSS usage. We selected cell type populations derived from post-mortem brain (neurons, astrocytes, endothelial, and smooth muscle), post-mortem fetal and adult frontal lobe samples, 26 day old in vitro neurons produced with an NGN1 and NGN2 overexpression differentiation protocol³⁴, and 60 day old *in vitro* brain organoids³⁵. For each sample, we sequenced CAGE libraries, sampled 13 million aligned reads, called peaks using ParaClu, and applied CapFilter (Online Methods). We intentionally focused on differences in TSS usage rather than expression levels (Online Methods) to explore the specific information added by CAGE compared to standard RNA-Seq. Also, we wanted to identify the most significant differences in TSS usage and this can be difficult to discern for genes with low expression in a given sample. To identify TSS peaks differentially used between pairs of samples, for each gene we compared the fraction of reads in a TSS peak between samples. This method identified 2,312 TSS peaks in 1,015 genes that were significantly different (FDR < 0.05, Fisher's exact test) between at least one pair of these eight samples (Supplementary Table 7). Unsupervised hierarchical clustering of the differentially used TSSs showed relationships among these samples, such as the in vitro models of neuron development being most closely related to the fetal frontal lobe sample (Fig. 5a). For comparison purposes, we also performed unsupervised hierarchical clustering using gene expression levels rather than TSS usage (Supplementary Fig. 13) and observed the resulting clustering of samples is similar to that in Fig. 5a.

Focusing on differences in TSS usage between fetal and adult samples, our analysis highlighted three brain disease-associated genes with differential TSS expression (Fig. 5b). For *ANK3*, which has been associated with bipolar disorder and schizophrenia³⁶, the T2 (exon 1e³⁷) TSS was more frequently used than the T3 (exon1b³⁷) TSS in all samples, except the adult frontal lobe, consistent with published findings³⁷. For *GPR56*, the T4 (e1m³⁸) TSS, which has been shown to be the highest expressed TSS in fetal human brain and required for normal human embryonic cerebral cortical development³⁸, was used more frequently in the fetal frontal lobe, *in vitro* neurons, and brain organoids, but not in the adult frontal lobe, in which T3 was used more often. For *RTN4* (also known as *NOGO*), we observed that the T3 (P2, NOGO-C³⁹) TSS, which was shown to be overexpressed in schizophrenia, was used more frequently in the adult frontal lobe, similar to published studies³⁹, but not in the other samples. For all three genes, TSS usage in adult frontal lobe was more similar to brain organoids than *in vitro* neurons (Fig. 5b), as might be expected given the more advanced development of the organoids³⁵.

More globally, *in vitro* neurons, brain organoids, and fetal frontal lobe used an upstream (more 5') TSS rather than a downstream (more 3') TSS significantly more often (Bonferroni adjusted P value < 0.05) than adult frontal lobe (Fig. 6a,b). For this comparison, we computed a scaled average peak position for all genes and datasets, which were compared between samples using a Paired-Wilcoxon signed-rank test (Online Methods). TSS usage was most significantly different between *in vitro* neurons and adult frontal lobe (Bonferroni adjusted P value = 0.0002) with significant differences also observed between fetal and adult frontal lobe (Bonferroni adjusted P value = 0.0013) and between brain organoids and adult frontal lobe (Bonferroni adjusted P value = 0.023, Fig. 6b). We observed this trend also using FANTOM5 CAGE data from similar samples²⁷ (Fig. 6b). Moreover, the number of TSS per gene was significantly higher (Bonferroni adjusted P value = $2.1*10^{-32}$) in the adult fetal lobe than the *in vitro* neuron or organoid samples from this study as well as in the adult FANTOM5 brain samples compared to the fetal FANTOM5 brain samples (Fig. 6c).

DISCUSSION

We compared six methods for 5' RNA-Seq by a comprehensive set of quality measures. CAGE performed best overall (Figs. 2 and 3), though other methods might be deployed when less RNA is available. GRO-cap should be considered for identification of eRNA TSSs (Supplementary Tables 5 and 6) but performed worse for mRNA TSSs due a higher rate of false positives (Fig. 3), particularly in genomic regions annotated as introns (Supplementary Table 1). While many peaks identified by GRO-cap could be "real" based on corroborative evidence (Supplementary Fig. 11), it is difficult to judge. Furthermore, GRO-cap is limited to fresh samples and relies on the TAP enzyme, which is no longer commercially available. The methods also vary in the associated time and cost of materials and kits (Supplementary Table 8). The per-sample cost is lowest for STRT and highest for Oligo capping. The amount of time and number of steps per library for each method is lowest for STRT and NanoCAGE-XL and highest for CAGE.

We aimed to test a fully representative set of the existing methods for 5' RNA-Seq, but excluded some from our comparisons (Supplementary Note 5).

A key question was which methods were best for annotating TSSs in a sample without previous annotation. Beyond finding that CAGE performed best, we explored how DNase-Seq and H3K4me3 ChIP-Seq could refine these assignments (Fig. 4). We found that DNase-Seq and H3K4me3 are insufficient without CAGE to reliably identify TSSs. Moreover, DNase-Seq, but not H3K4me3, does provide additional specificity beyond CAGE. Of course, it is not possible to know with full certainty whether any given TSS identified by these methods is correct. For standard RNA-Seq, DNase-Seq and, to a lesser extent, H3K4me3, are more valuable in identifying true TSSs based on the annotation, though there are still more false positives with standard RNA-Seq than CAGE even with the corroborative data (Supplementary Fig. 12). ATAC-Seq⁴⁰ could be substituted for DNase-Seq as the former is simpler with lower input requirements.

Finally, we addressed the question of TSS usage, which is critical for studying disease related genetic variation in non-coding regions, focusing on its relevance to assessing the

faithfulness of disease models. Human pluripotent stem cells (hPSCs) offer an excellent tool to model human disorders in the lab^{41, 42}, and are particularly relevant for brain-related disorders for which other models are limited^{43, 44}. We compared TSS usage *in vitro* neurons and brain organoids derived from hPSCs with post-mortem brain samples (Figs. 5 and 6) as it is important to understand how faithfully these models represent actual brain tissue at a transcriptional level. Our results provide a resource for future studies and add to the existing literature in this field^{27, 45}. Brain organoids were more similar than *in vitro* neurons to adult post-mortem brain samples by our TSS usage measures (Figs. 5a and 6b) – suggesting that the former might be a better model by these criteria. Some of the observed differences are likely explained by post-mortem brain having cell types not found in these *in vitro* models and potentially changes in RNA abundance due to death-associated cellular responses to hypoxia and other factors⁴⁶.

With the brain-related samples, we observed a genome-wide trend that relative TSS usage significantly varied with respect to upstream or downstream position within each gene (Fig. 6). Because the size of this effect seems to reflect the overall relatedness of the samples (Fig. 5a) and we identified this pattern in an unrelated CAGE dataset (Fig. 6b), we believe there may be a biological explanation underpinning this result though understanding its basis is beyond the scope of this study. To our knowledge, this phenomenon has not been observed previously, but studies of the 3' ends of transcripts have shown that upstream polyadenylation sites are preferentially used in proliferating cells⁴⁷, while the opposite has been reported in mammalian brain⁴⁸. Also, a recent paper showed unannotated TSSs were detected upstream of annotated, silenced promoters that may be relevant to aberrant gene expression in cancer due to hypomethylation of these upstream TSSs⁴⁹.

ONLINE METHODS

RNA samples.

We used the same batch of K-562 total RNA (Thermo Fisher Scientific) in experiments directly comparing the lab methods. For the CAGE, RAMPAGE, and STRT methods, we used a second batch of K-562 RNA (Thermo Fisher Scientific) for comparison to assess method reproducibility. Both batches were high quality RNA, with RNA Integrity Number (RIN) scores of 8.6 and 8.8 as measured with a BioAnalyzer (Agilent).

For the brain-related samples, we used total RNA from each of eight human samples. From ScienCell Research Laboratories, we used brain vascular smooth muscle cell RNA (#1105), brain microvascular endothelial cell RNA (#1005), neuron early passage monolayer culture from fetal brain RNA (#1525), and astrocytes cultured for 6 days from a fetal donor RNA (#1805). From Biochain, we used fetal frontal lobe RNA (#R1244051-50) and adult frontal lobe RNA (#R1234051-50).

We differentiated HUES66 embryonic stem cells, which was created by the Harvard Stem Cell Institute, into neurons by overexpressing *NGN1* and *NGN2*³⁴, harvested the cells after 26 days, and isolated total RNA using the Quick-RNA MiniPrep kit (Zymo Research).

We prepared four brain organoids from a single batch at day 60 of differentiation from H9 embryonic stem cells as previously described 50 . We isolated total RNA using 1 ml Trizol (Thermo Fisher Scientific) according to the manufacturer's instructions and isopropanol precipitated with 1 μl GlycoBlue Coprecipitant (Thermo Fisher Scientific). We removed the GlycoBlue, which inhibited the CAGE process, by mixing with 30 μ mol LiCl (Sigma) in a total 12 μl and incubating at $-20^{\circ}C$ for 20 minutes following by centrifugation at $4^{\circ}C$ at maximum speed (17,000 \times g). We removed the supernatant, rinsed the pellet with 70% ethanol, and re-suspended the pellet in 8.22 μl H₂O. We repeated this process one more time until there was no visible GlycoBlue. RIN scores for these 8 samples were 6.8 to 9.7 as measured by BioAnalyzer.

All biospecimens were collected with informed consent. The generation of hES cells used in this study was approved by the institutional review boards (IRBs) of the providing institutions. Use of all de-identified biospecimens for sequencing at the Broad Institute was further approved reviewed by the Broad's Office of Research Subject Protection (ORSP), which determined that the research did not involve human subjects according to U.S. federal regulations (45CFR46.102f). This study complied with all relevant ethical regulations.

RNA spike-in controls.

We obtained 32 individual ERCC spike-in controls 23 (gift from Jennifer McDaniel and Marc Salit, NIST). We added a m7G cap structure to the 5' end of RNA molecules using the Vaccinia Capping System (New England BioLabs) following the manufacturer's protocol. We made a pooled capped ERCC spike-in mix with an average concentration of 36 pg/µl for each transcript. In addition, we prepared a second pooled, capped ERCC spike-in mix with only eight of the transcripts with an average concentration of 138 pg/µl for each transcript. We also made a third pooled original (uncapped) ERCC spike-in mix for all 32 transcripts with average concentration of 60 pg/µl for each transcript.

CAGE libraries.

For K-562 samples, we prepared libraries following a published protocol⁹ using 10 µg total RNA for three replicates ("Main-1, 4, and 6") from the same batch of RNA. In a second experiment, we prepared a library with a single replicate ("Repeat") from 5 µg of a different batch of K-562 RNA. For the brain-related samples, we prepared CAGE libraries from eight total RNAs from sources described above, using between 5 to 10 µg each based on availability.

For the Main-1, Main-4, and brain-related samples, we added pooled capped ERCC spike-in RNA to each sample in a ratio of 0.128 µl per µg sample input. For the Main-6 sample, we added 1.28 µl pooled, uncapped ERCC spike-in RNA. For the other replicate (Repeat), we added 1.0 µl pooled capped ERCC spike-in mix containing only eight transcripts.

RAMPAGE libraries.

We prepared the libraries following the published protocol 11 using 5 μ g K-562 total RNA plus 0.64 μ l pooled capped ERCC spike-in RNA with the following modifications. (1) We used a universal template switching oligo 5'-

TAGTCGAACTGAAGGTCTCCAGCArGrGrG-3' instead of the barcoded ones. (2) We used a random 15mer oligo with modified tag sequence for RT to allow index read later on in sequencing 5'-TAGTCGAACGAAGGTCTCCCGTGTGCTCTTCCGATCT(N)15. (3) In the final PCR, we used an 8-base barcoded reverse primer to add index for each individual library, 5'-CAAGCAGAAGACGGCATACGAGATxrefXXGTGACTGGAGT-3'. (4) We used a different custom sequencing primer (Supplementary Table 9). We also made a replicate with a different batch of K-562 RNA with 1.0 µl pooled capped ERCC spike-in mix containing only eight transcripts.

STRT libraries.

We synthesized cDNA from 10 ng K-562 total RNA plus 0.64 μl 1/500× dilution of pooled original, uncapped ERCC spike-in RNA following a published protocol¹², combined with reverse transcription-PCR conditions based on SMART-Seq2⁵¹ and the following modifications. (1) We used a 5' biotin blocked dT oligo that contains a SalI restriction site (shown as underlined), 5'-/5Biosg/CTACACGACGCTCTTCCGATCTGTCGACT(30)VN-3'. (2) We used a 5' template switching oligo (TSO) containing an Illumina adaptor sequence, 5'-CUACACGACGCUCUUCCGAUCUNNNNNGGG - noting all bases are RNA. We made this switch because this oligo is compatible with HiSeq2500, MiSeq, and NextSeq sequencers (Illumina), while the original one ¹² requires a custom sequencing primer with an annealing temperature that is only suitable for the HiSeq2000. (3) We used a PCR primer with sequence compatible to dT oligo and 5' TSO, 5'-CTACACGACGCTCTTCCGATCT-3', for cDNA amplification. We then eliminated the polyA/T end of the double stranded cDNA by mixing with 1x CutSmart buffer, 10 units of Sall (New England BioLabs) and heating at 37°C for 60 minutes. We purified this product by using 0.7x volume AMPureXP SPRI beads (Beckman Coulter Genomics) following vendor protocol. We made the sequencing library with a modified NexteraXT (Illumina) protocol⁵² with the following modifications in addition. (1) We used 0.125 ng cDNA in ½ volume of a standard NexteraXT reaction. (2) We used the modified Nextera Index 1 primer, 5'-CAAGCAGAAGACGCATACGAGATxrefXXGTCTCGTGGGCTCGGAGA*T*G-3' with phosphorothioate bonds (denoted by *) and inverted end bases for protection; the 8 "X" bases indicate in-line index sequences that enable pooling samples. In the same experiment, we also prepared a library with the same quantities of capped ERCC spike-in RNA and K-562 RNA. We also made a replicate with a different batch of K-562 RNA.

In addition, we constructed STRT libraries using a similar protocol as above with 10 ng, 100 ng, 1 μ g, 5 μ g and 10 μ g K-562 total RNA from the same batch as above together with the same proportion of uncapped ERCC spike-in RNA. We made the following modifications to accommodate the higher inputs. (1) For the 5 μ g and 10 μ g inputs, we used 4 times as much the volume for the RT reaction. (2) We used 10 μ l each of the 10 times diluted first strand cDNA from the 1 μ g RNA input, 12.5 times diluted ones from the 5 and 10 μ g RNA inputs, into the 50 μ l of cDNA PCR amplification reaction. (3) We used a 50 μ l reaction volume that contained 40 units of SalI to remove the polyA/T end of the double-stranded cDNAs generated in all the higher input RNA libraries.

Oligo Capping libraries.

We prepared the library following published protocols 16,53 , using 40 μ g K-562 total RNA plus 5.12 μ l pooled capped ERCC spike-in RNA with the following modifications. (1) We used 0.5 μ l glycogen (Roche) as the carrier instead of ethachinmate in the ethanol precipitation step. (2) We used KAPA HotStart Ready Mix (KAPA Biosystems) instead of GeneAmp PCR kits (PerkinElmer) for PCR amplification. (3) We selected 250 to 600 bp instead of 150 to 250 bp PCR products for sequencing.

NanoCAGE-XL libraries.

We prepared a library using 7.5 µg K-562 total RNA plus 0.96 µl pooled capped ERCC spike-in RNA following the published protocol¹⁴ with the following modifications. (1) We used a TSO containing 6-base barcode (marked as xref) followed by 6-base spacer, 5′-TAGTCGAACTGAAGGTCTCCAGCAxrefGCTATArGrGrG. (2) We used a modified custom sequencing primer (Supplementary Table 9).

Standard RNA-Seg library.

We prepared a library using 1 μ g K-562 total RNA plus 2 μ l of 1:100 diluted ERCC spike-in mix 1 (Ambion) using the TruSeq RNA-Seq kit (Illumina) with the following modifications. (1) The rRNA was depleted using the RNase H method¹⁹ instead of using oligo (dT) selection. (2) We eluted the rRNA depleted RNA from the SPRI beads using EPF buffer from the TruSeq kit and heated at 70°C for 10 minutes. (3) We used a different set of barcoding indices rather than those in the TruSeq kit for the ligation and final PCR steps.

Sequencing

Libraries were sequenced with either HiSeq2500, MiSeq, or NextSeq machines (Illumina), as detailed in Supplementary Table 9. We used paired-end sequencing for some 5' end libraries to aid with understanding method performance, but this is not generally required with the possible exception of RAMPAGE. The NanoCAGE-XL library was sequenced with a second unrelated library and spiking in 30% PhiX library and loading at a reduced concentration (7 pM) to overcome monotemplate issues with libraries prepared with this method.

Additional 5' RNA-Seq datasets

For datasets previously generated by other groups, we downloaded the relevant fastq files (Supplementary Table 10) and used them in our method comparisons. For the STRT data²⁹, we picked 100 random mouse hippocampus cells from the single cell dataset and combined them together into one fastq file before processing.

Read processing and alignment

For 5' end RNA-Seq, we processed reads by trimming an appropriate number of bases depending on the lab method used (Supplementary Table 1). We aligned reads to either the human genome (Gencode v19) or mouse genome (mm10) with STAR⁵⁴ (version 2.4.2a) using two-pass mode, and the softclip option to trim reads and otherwise default parameters. We only used read 1, which was derived from the 5' end of the transcript, for these analyses,

except when we tested the use of read 2 with the RAMPAGE peak calling pipeline (see below). We generated basic alignment and performance metrics, using CollectAlignmentSummaryMetrics and CollectRnaSeqMetrics in Picard Tools (https://github.com/broadinstitute/picard). We analyzed the reads for both these metrics and peak calling analyses, but used BAM files generated by STAR with the hardclip option for the former because Picard did not recognize bases as trimmed with the softclip option.

For 5' vs 3' end coverage, we sampled 20 million reads from each bam file, except NanoCAGE-XL, for which we used all reads. We performed two types of analyses – one using the entire gene including intronic regions and the other using only the exonic regions. In both cases, we used only the position of the most 5' end of the read ("read start"). For the analysis of the entire gene, we only used genes greater than 500 bp in length and with Transcripts Per Million (TPM) > 1 as estimated with RSEM. We divided each gene into 100 equal sized bins, and totaled the percent of read starts on the same strand as the gene in each bin on a per gene basis using bedtools⁵⁵ intersect and post-processing in R. We averaged these percentages over all genes, and plotted the results with ggplot⁵⁶. For analysis using only the exonic regions, we used a similar approach, except for genes with more than one transcript, we selected the one with the highest TPM for each gene and limited our analysis to transcripts with greater than 500 bp in length and TPM > 1.

For ERCC spike-in analysis, we used the same trimmed reads, except as noted in Supplementary Table 1. We aligned to a version of all 92 of the ERCC spike-in RNA sequences that includes all sequences at the 5' ends (Supplementary Table 11) using BWA⁵⁷ and a custom Picard module to parse the aligned BAM file to calculate the coverage at each base of the ERCC spike-in reference sequences.

For standard RNA-Seq data, we aligned untrimmed reads with Tophat⁵⁸ (version v1.4.1) using default parameters except mate-inner-dist set to 300 and mate-std-diff set to 500, followed by Cufflinks⁵⁹ (v2.2.1) using the default settings.

Peak calling

To identify TSSs, we used the Paraclu²⁴ peak caller, applied to randomly sampled reads as follows. To compare between methods, we randomly sampled 20 million reads from aligned K-562-derived BAM files using a custom shell script, with the exception of NanoCAGE-XL, for which we used all 6,407,741 reads. For other parts of the analysis (reproducibility, MCF-7, mouse hippocampus, human brain-related data, FANTOM5 data) we sampled to different numbers of reads, as detailed below. We sampled only reads from each BAM file that passed platform / vendor quality checks and were flagged as being a primary alignment. For 5' end methods, 20 million reads are only from read 1, but for standard RNA-Seq, they are from both reads 1 and 2.

Post-processing of ParaClu peaks

We annotated peaks called by ParaClu with several scores to indicate the confidence for increased read density in that region, which were used to remove low-confidence peaks. The ParaClu caller adopts a hierarchical approach to peak calling and calls peaks within peaks²⁴. For the purpose of assessing TSS discovery accuracy, we were less concerned with internal

peak structure, and therefore merged all overlapping peak regions using the bedtools⁵⁵ 'merge' function. We set the scores for the aggregate peak to the maximum ParaClu score over the set of peaks to be merged. All peak regions wider than 300bp or narrower than 3bp were removed.

To indicate the confidence level for a given peak, ParaClu provides three metrics, D – the 'density rise', a measure of the fold change between maximum and minimum read density, and an indicator of signal strength., P – the minimum number of positions within the peak covered by reads, and S– the total number of reads mapping to the region within the peak. Low-confidence peaks could then be removed by setting threshold values D_{\min} , P_{\min} and S_{\min} for these three values, respectively, and removing peaks that do not pass all three cutoffs. To identify optimal values for these parameters, we tested all combinations of (integer) values for D_{\min} in [0,10], P_{\min} in [0,20], and S_{\min} in [0,180], and assessed peak-calling performance using the F_1 score (see below). To ensure that each 5' RNA-Seq method was analyzed optimally, we repeated this procedure to identify the best parameters for each method, and at the different read depths required for all the comparisons in this study: 20 million for 5' end RNA-Seq lab method comparisons, 7 million for CAGE reproducibility, 5 million for RAMPAGE and STRT reproducibility, and 13 million for brain-related samples, MCF-7, and mouse hippocampus samples. The final filtering parameters used for each dataset are found in Supplementary Table 12.

We assessed the reproducibility of TSS discovery by using bedtools⁵⁵ 'intersect' to compute the pairwise overlap between ParaClu peaks called using BAM files generated from four CAGE replicates. We also compared overlaps for pairs of replicates for RAMPAGE and STRT.

Peak calling from standard RNA-Seq data

To identify TSS using full-length RNA-seq reads, we merged aligned reads from two replicates using samtools⁶⁰ and sampled 20 million reads as described above. With expressed transcripts identified using Cufflinks⁵⁹, we annotated the region within 100 bp of the 5' end of each identified transcript as a TSS peak.

Tag cluster identification

To identify TSSs in enhancer regions and understand narrow vs broad peaks, we used an alternative peak calling method²⁷ that aims to find tag clusters (TCs). For each read, we defined the starting location of that read as the start of a TSS and then merged all such TSS starts within 20 bp on the same strand to get TCs. We discarded TCs with less than 3 reads supporting them or longer than 300 bp. We used CapFilter (see 'additional filtering steps' below) for methods that add an extra G (CAGE, RAMPAGE, NanoCAGE-XL and STRT). We defined broad peaks as TC's > 10 bp in width and narrow peaks as those 10bp.

Peak calling for enhancers

To identify putative eRNAs in K-562 datasets, we adapted a previously published approach³³. In particular, we took all intergenic TCs and discarded TCs that overlapped another TC on the opposite strand. We paired reverse and forward stranded TCs, where the

reverse stranded TC occurred within < 400 bp of the forward stranded TC, and merged overlapping pairs together avoiding overlap between the reverse and forward stranded TC in each pair. We then filtered out all merged pairs where either the reverse or forward TC had much higher coverage than the other. More specifically, we only kept merged pairs with:

$$-.8 < \frac{Number\ Reads\ Reverse\ TC-Number\ Reads\ Forward\ TC}{Number\ Reads\ Reverse\ TC+Number\ Reads\ Forward\ TC} < .8$$

We used the middle point of each of these merged pairs as the center of the putative enhancer, and extended by 200 bp on either side to generate the putative enhancer. We compared these putative enhancers to public H3K27ac ChIP-Seq (peaks downloaded from: http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeBroadHistone/wgEncodeBroadHistoneK562H3k27acStdPk.broadPeak.gz), DNase-seq (see Corroborative data), and enhancer region datasets (ENCFF687ZGE) from ENCODE²⁸ and the DENdb database⁶¹ (http://www.cbrc.kaust.edu.sa/dendb/) using the bedtools⁵⁵ intersect function with default parameters. Similarly, we compared the intergenic peaks identified with ParaClu to each of the above datasets using the same approach to test if the peaks overlapped with eRNA.

Identification of TSS initiator sequences

For identifying the dinucleotide sequences at the start of TSSs ("-1" and "+1", where +1 is the position of the first transcribed base in a transcript and -1 is the position directly before it in the genome), we used a modification of previously published approaches^{62, 63}. In particular, we took all TCs within 100 bp of an annotated gene and with at least 10 reads mapping to them, and located the position in each TC with the largest number of reads starting there. We used this location as the putative start site (the +1 location) for the TC. Note that for methods that add an extra G (CAGE, RAMPAGE, NanoCAGE-XL and STRT), we shifted the putative start site over by one base pair in the 3' direction. We used bedtools⁵⁵ to extract the +1 and -1 base for each putative TSS from the human genome and created logos using the ggseqlogo package in R⁶⁴.

Additional filtering steps

We evaluated three specialized filtering programs: CapFilter¹⁴, strand invasion⁶⁵, and the RAMPAGE¹⁰ second read filter (downloaded from http://megraw.cgrb.oregonstate.edu/software/CapFilter, https://academic.oup.com/nar/article/41/3/e44/2902349/Suppression-of-artifacts-and-barcode-bias-in-high#supplementary-data, http://labshare.cshl.edu/shares/gingeraslab/www-data/dobin/ENCODEpipelines/ENCODE_longRNApipeline_GIT/DAC/rampagePeakCaller.py). For CapFilter and strand invasion, we slightly modified the code to allow applications beyond NanoCAGE-XL – the lab method for which they were originally developed. We applied CapFilter to CAGE, RAMPAGE, NanoCAGE-XL, and STRT. In each case, we began by calling peaks as above and then applied CapFilter to the resulting peaks. This led to a smaller set of final peaks. We tested CapFilter with various settings of the thresholding parameter (which corresponds to the percentage of G's in the first position), and found that a threshold of 20% seemed to be optimal for all experimental methods (Supplementary Fig. 2).

We applied the strand invasion filter⁶⁵ to the NanoCAGE-XL and STRT data. This filter removes potentially artefactual reads that are the result of strand invasion, where the PCR primer primes at sequences in the cDNA similar to the template-switching oligonucleotide (TSO) rather than in the TSO. By checking for matches between the TSO and the sequence upstream of read 1, such events can be identified and filtered out. Using all the NanoCAGE-XL aligned reads and 20 million sampled STRT aligned reads from the BAM files, we applied the strand invasion filter, varying the maximum allowed edit distance between the TSO and the upstream sequence, before calling peaks with ParaClu. We tested varying this parameter between 0 and 7, both with and without CapFilter. We found that this filter did not improve on CapFilter when used in combination with CapFilter (Supplementary Fig. 3a) and did not include strand invasion in our main analysis.

We compared the RAMPAGE peak caller¹⁰ with and without second read filtering to the Paraclu results. Because the RAMPAGE filter did not improve performance CapFilter (Supplementary Fig. 3b), we did not include it in our main analysis.

Accuracy assessment

In order to estimate the accuracy of each method, we looked at the number of peaks that overlapped known TSSs based on the UCSC annotation (see Corroborative data). True positive (TP) peaks were defined as those that overlapped at least one annotated TSS, and false positive (FP) peaks as those that did not overlap any TSSs. Intergenic peaks did not overlap any genes according to the UCSC annotation. False negative (FN) peaks were defined as all annotated TSSs, which were (1) within any gene expressed with TPM >1 (quantified using RSEM 66 (v.1.2.7) from 5' RNA-Seq data), (2) overlapped by at least one DNase-Seq peak (see Corroborative data), and (3) did not overlap any of the peaks called by ParaClu. We calculated the ROC curve for each method based on the peak scores output by Paraclu. We calculated sensitivity, precision, and F_1 score according to these formulas.

$$Precision = \frac{TP}{TP + FP}$$

$$Sensitivity = \frac{TP}{TP + FN}$$

$$F1 = 2 \frac{Sensitivity*Precision}{Sensitivity*Precision}$$

In all cases, a 100bp tolerance was considered an overlap, implemented using bedtools 'window'.

For mouse-derived data, we modified the expression estimation pipeline, so that instead of using RSEM with its standard settings to quantify expression when defining false negatives, we first mapped the reads to the mm10 UCSC transcriptome using STAR and then applied RSEM to the resulting bam file. In order to produce a bam file that RSEM could use, STAR

was run with the options: --quantMode TranscriptomeSAM, --alignIntronMax 1, --alignIntronMin 2, --scoreDelOpen -10000, and --scoreInsOpen -10000.

Corroborative data

H3K4me3 peaks were originally generated from ChIP-Seq as part of the ENCODE Consortium²⁸. We downloaded peaks from: http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeBroadHistone/wgEncodeBroadHistoneK562H3k4me3StdPk.broadPeak.gz.

The associated GEO Accession number is GSM733680. We obtained bam files for K-562 DNase-1 hypersensitivity peaks (DNase-seq) from the SRA database (SRR231254 and SRR231189), called peaks with Macs2⁶⁷ with the --nomodel option and default settings, and selected only peaks found in both replicates, by computing the intersection of the two replicates using bedtools⁵⁵ 'intersect'. For the other DNase-seq datasets, we downloaded bed files containing peaks from the ENCODE portal⁶⁸ (ENCFF408UYX for MCF-7; ENCFF630GRU for mouse brain).

TSS annotations

Annotated transcription start sites (TSS) were downloaded from the UCSC table browser, accessible at: https://genome.ucsc.edu/cgi-bin/hgTables and Gencode²⁵ annotated transcripts were downloaded from ftp://ftp.sanger.ac.uk/pub/gencode/Gencode_human/release_19/gencode.v19.annotation.gtf.gz and converted to a bed file containing TSS only by extracting the 5' end of all transcripts in the .gtf file using a custom shell script.

Coverage of peaks in brain-related samples

We sampled 13 million aligned reads from each brain-related dataset and called peaks with Paraclu and applied CapFilter as above. We combined the resulting peaks from all the samples into one file and merged with bedtools⁵⁵. This ensures that slight differences in peak calls between samples (due to inherent randomness in the data) do not affect downstream analysis. We removed all peaks that did not overlap at least one annotated TSS within a tolerance of 100 bp. We used the bedtools coverage command to find the number of reads covering each peak – resulting in a matrix with one column for each sample and one row for each (merged) peak. We also used bedtools to annotate each peak to include the name of the gene in which it was located.

Coverage of peaks in FANTOM5 data

We downloaded FANTOM5²⁷ BAM files from fantom.gsc.riken.jp/5. In total we downloaded 12 samples: iPS day 18 of differentiation to neurons⁶⁹, adult frontal cortex, adult and fetal occipital cortex, adult and fetal parietal cortex, and adult and fetal temporal cortex (Supplementary Table 10).

We processed the iPS and adult frontal cortex data through the same pipeline as our brain-related data, to generate a matrix with one column for each sample and one row for each (merged) peak. We used a similar pipeline for the remaining six samples, except instead of sampling 20 million reads, we sample 6 million reads for the parietal cortex sample and 10

million reads for the remaining four samples because there were fewer reads for these samples. In addition, we were unable to apply CapFilter to the FANTOM5 reads because they were already trimmed.

Differential TSS usage

To control for differences in overall gene expression, we computed the *relative usage* for each TSS i in gene j as $R_{ij} = 100 \frac{r_{ij}}{\sum_{i=1}^{k_j} r_{ij}}$, where r_{ij} is the number of reads within TSS peak

i in gene j and k_j is the number of TSSs in gene j. To identify differential usage, we considered all gene/ sample pairs with > 100 reads, and compared the relative usage (rounded to the nearest integer) for the set of TSSs in a given gene between each pair of samples using Fisher's exact test implemented using the R function fisher.test in the stats package with default parameters, including alternative="two.sided." More specifically, the test was performed on the 2 by k_j table with one row per sample and one column per TSS, where the table entries are taken to be the corresponding R_{ij} values (we use R_{ij} values instead of raw counts to help correct for number of reads per gene, resulting in a more conservative test statistic). We then took the minimum P value across all comparisons, Bonferroni corrected for the multiple comparisons per gene, was reported as the P value for differential usage across the samples within a gene. Note that we only compared genes with at least 100 reads mapped to them in the samples of interest. We corrected Fisher P values for multiple hypothesis testing using the Benjamini-Hochberg FDR correction, implemented using the R function p.adjust.

Testing upstream vs. downstream bias in TSS usage

To test if there was an upstream vs. downstream bias in TSS usage between different brain samples, we used the peak by sample matrix generated above. For each gene and each sample, we calculated an average normalized peak score. More formally, for a given gene, if that gene had k peaks 1, 2, ..., k (ordered from most to least upstream), and had n_{ij} reads covering the j-th peak in the j-th gene, the score for that gene was equal to:

$$score_{i} = \frac{1}{k} \sum_{j=1}^{k} j * n_{ij}$$

The larger this score, the more that downstream peaks are used.

We used a Wilcoxon signed-rank test (one-sided) to compare the average normalized peak scores for genes between samples. We correct for multiple testing with Bonferroni correction (a total of 20 tests were made—12 from our brain-related data, 8 from the FANTOM5 data).

Expression analysis

To compare estimated gene expression values between samples (Supplementary Fig. 10), we extracted TPM values from the RSEM results for each 5' method, as well as from standard

RNA-Seq. We log normalized these data before calculating the Pearson correlation coefficient, r.

The coloring in each comparison is based on a normalized density. For each pair of samples, this was calculated by removing all genes with TPM = 0 in either sample, and using a 2-D Gaussian kernel-based density estimate, using the kde2d function in the R MASS package⁷⁰. The density values were normalized to be between 0 and 1, to allow for a shared scale—this enables us to see the relative density around each gene.

Statistics

To compare TSS usage for a given gene between different conditions, we used a Fisher's exact test (see 'Differential TSS usage' for more details). For this test n=2 (one sample for each condition). In order to test for difference in number of TSS used and 5' biases between conditions we used a one-sided Wilcoxon signed-rank test (see 'Testing upstream vs. downstream bias in TSS usage' for more details), with 95% confidence intervals estimated using a Gaussian approximation. Again, this test uses n=2 (one sample for each condition). Multiple hypothesis testing was performed for all statistical tests using either Bonferroni or Benjamini-Hochberg FDR correction. All other statistics included in the paper are descriptive in nature.

Life Sciences Reporting Summary

Further information on experimental design is available in the Life Sciences Reporting Summary.

Code availability

Custom computer code used to generate results that are reported in this study and central to its main claims is freely available at https://github.com/seanken/FivePrime.

Data availability

Sequence data for K-562 libraries are available at Gene Expression Omnibus GSE103486.

Sequence data for human brain-related samples are available at dbGaP under accession code phs001463.v1.p1.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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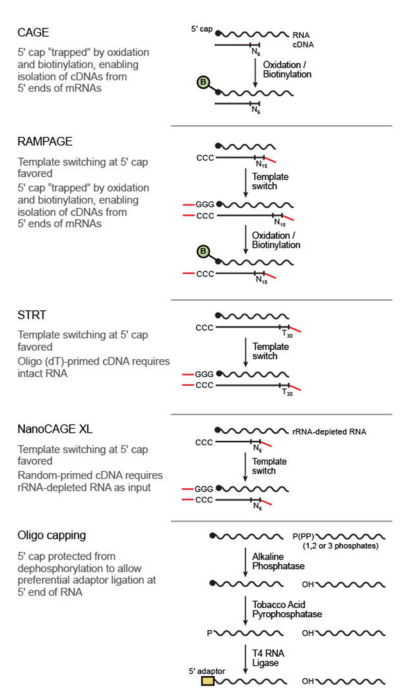


Figure 1. Methods for 5' end RNA-Seq.

Salient details for five protocols tested in this paper. Additional properties of these protocols can be found in Supplementary Table 8.

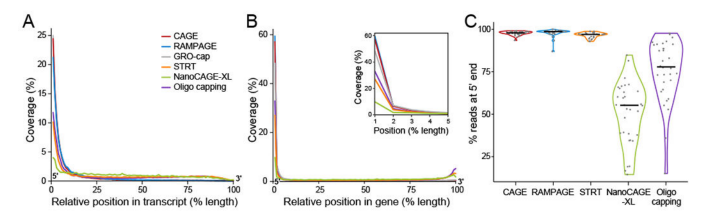
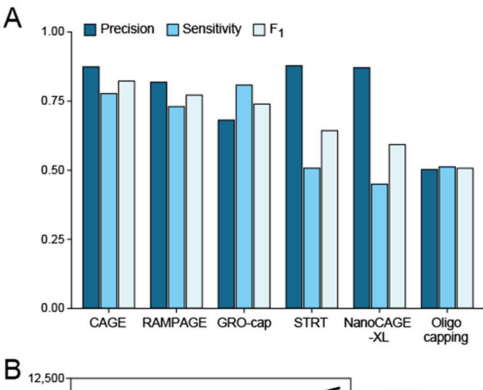


Figure 2. Read performance metrics for 5' end methods.

(**a,b**) Normalized coverage by position for endogenous transcripts. For each library, shown is the average relative coverage (*y*-axis) at each relative position along the transcripts' (**a**) or genes' (**b**) length (*x*-axis). Intronic regions are included in (**b**), but not in (**a**). Inset in (**b**) shows 5% closest to the 5' end of genes. (**c**) 5' end coverage for spike-ins. For each library, a violin plot shows the % of reads with alignment including position 10 from the 5' end of each of the 32 spike-in transcripts (*y*-axis). Median is shown as a black line. For libraries with replicates, data are shown for the "Main" library (Main-1 for CAGE; Online Methods). STRT data presented are for un-capped spike-in RNA, which performed better than capped spike-in RNA. Sample size for each method: n = 1 library per method.



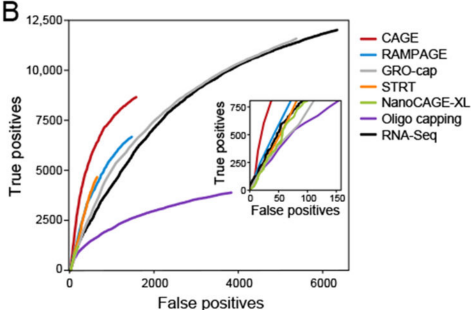


Figure 3. TSS peak performance metrics.

(a) Sensitivity, precision, and F_1 score (the harmonic mean for sensitivity and precision) – shown for each 5' end method based on the UCSC annotation (b) ROC curves for each lab method and standard RNA-Seq with inset showing highest confidence region. Sample size for each method: n = 1 library per method, except CAGE is a combination of 3 libraries.

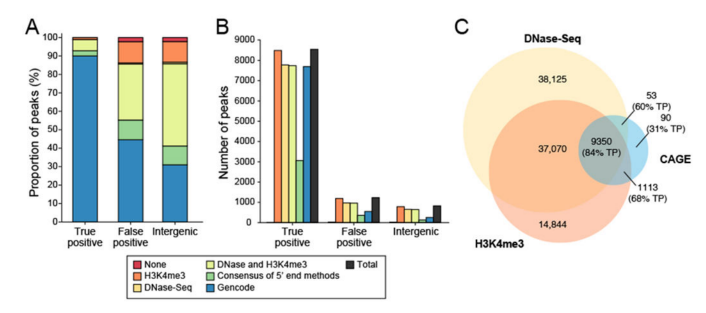


Figure 4. TSS discovery for unannotated CAGE peaks.

(a,b) Corroborative data for TSS peaks from CAGE. Shown is the proportion (a) and number (b) of peaks (y axis) with support from each corroborative data source (color legend) for peaks initially defined as 'true positive', 'false positive' and 'intergenic' based on the UCSC annotation. (a) Peaks were assigned to only one category of support based on their corroboration by Gencode annotation, consensus of four best 5' end methods, DNase-Seq, or H3K4me3 ChIP-Seq data in this order (e.g., a peak corroborated by Gencode is not listed in the other categories even if it has additional support.) (b) Peaks were assigned to as many corroborative categories as evidence supported. (c) TSS prediction with CAGE, DNase-Seq and H3K4me3 ChIP-Seq data. Numbers of peaks shown here in overlapping categories correspond to CAGE peaks for all overlaps involving CAGE peaks and DNase-Seq peaks in the overlap with only H3K4me3 ChIP-Seq peaks. For each subset of CAGE peaks, we also show the % true positives (TPs) out of all the CAGE peaks in that category. Areas not to scale.

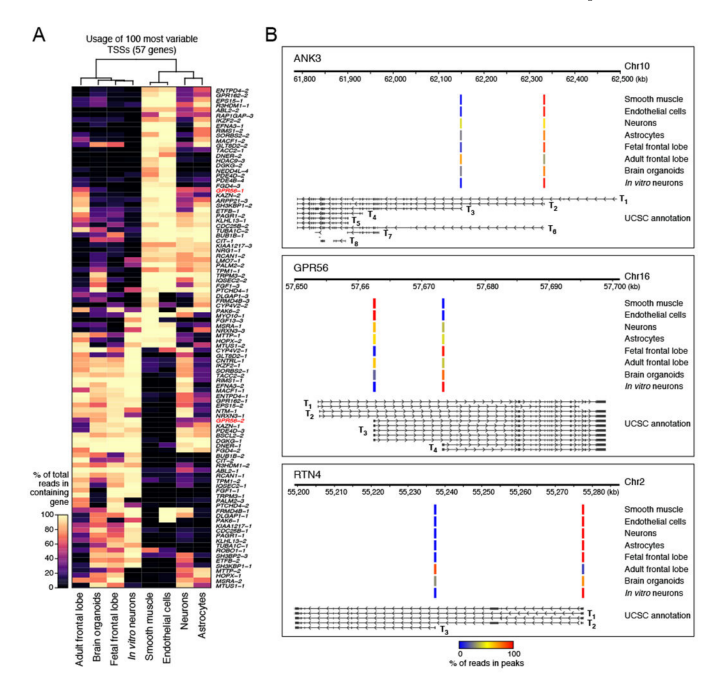


Figure 5. Differential TSS usage in brain-related samples.

(a) Most variable TSSs across brain-related samples. Shown are the top 100 most significantly differentially used TSSs across the samples (p < 0.001, Fisher's exact test) ordered by their variance. Sample size for each method: n = 1 library per sample. (b) Specific examples of differential TSS usage. For each gene, shown are the alternative transcripts and TSSs (T_i , bottom) and the scaled values of TSS usage (reads in a peak / all reads in peaks for a given sample) in each sample for each of the alternative TSSs.

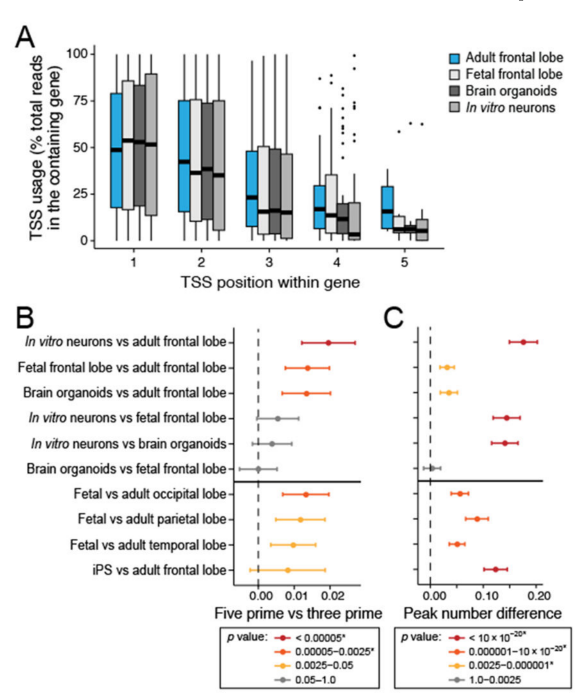


Figure 6. Adult brain samples preferentially use more downstream TSSs.

(a) Adult frontal lobe used downstream TSSs more often than fetal frontal lobe, brain organoids, and *in vitro* neurons. Numbering of TSS position within a gene starts from the 5' end. Box and whisker plot shows the relative TSS usage (*y*-axis) for all TSS, black bar indicates median value, box edges correspond to the 25th and 75th percentiles, while whiskers indicate a further 1.5*IQR where IQR is the interquartile range. Outliers outside this range are shown as dots. (b) Comparisons of sample pairs showing "younger" samples have more frequent upstream TSS usage in both this study and FANTOM 5. The *x*-axis is a

scaled, normalized difference of the average peak position in each dataset for all genes (Online Methods), with error bars representing 95% confidence intervals. (c) Comparisons of sample pairs showing "younger" samples use, on average, fewer TSS per gene in both this study and FANTOM 5. The x-axis is the average difference of the number of peaks active (defined as overlapping at least one read) in each dataset for all genes (Online Methods), with error bars representing 95% confidence intervals. For (b) and (c), the P values were calculated using a Wilcoxon signed-rank test (Online Methods) and an asterisk indicates a Bonferroni-corrected P value less than 0.05. The P values can be found in the source data spreadsheet file for this figure. For all panels, sample size for each method: n = 1 library per sample, except iPS FANTOM5 combines data for 2 replicate libraries.