Sites of transcription initiation drive mRNA isoform selection

Graphical abstract

Highlights
- A quantification of Drosophila and human nervous system full-length mRNA isoforms
- 3’ end site selection is coupled to alternative TSS usage
- Dominant promoters drive alternative polyadenylation through p300/CBP
- Conserved 5’-3’ couplings regulate tissue-specific functions

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In brief
Where an mRNA transcript starts determines where it ends. Epigenetic marks characterize “dominant promoters” that constrain start and polyadenylation sites, driving tissue-specific preferential expression of transcript variants.
SUMMARY
The generation of distinct messenger RNA isoforms through alternative RNA processing modulates the expression and function of genes, often in a cell-type-specific manner. Here, we assess the regulatory relationships between transcription initiation, alternative splicing, and 3' end site selection. Applying long-read sequencing to accurately represent even the longest transcripts from end to end, we quantify mRNA isoforms in Drosophila tissues, including the transcriptionally complex nervous system. We find that in Drosophila heads, as well as in human cerebral organoids, 3' end site choice is globally influenced by the site of transcription initiation (TSS). "Dominant promoters," characterized by specific epigenetic signatures including p300/CBP binding, impose a transcriptional constraint to define splice and polyadenylation variants. In vivo deletion or overexpression of dominant promoters as well as p300/CBP loss disrupted the 3' end expression landscape. Our study demonstrates the crucial impact of TSS choice on the regulation of transcript diversity and tissue identity.

INTRODUCTION
Variation at each step of pre-messenger RNA (mRNA) synthesis impacts the coding and non-coding content of the mature transcript. Alternative splicing (AS) and alternative polyadenylation (APA) generate mRNA isoforms that differ in their coding sequence (CDS) or the length of their 3' untranslated region (3' UTR), thereby contributing to proteome diversity and fine-tuning gene expression. Alternative 3' UTRs, through distinct sequence and structure elements that dictate interactions of the transcript with microRNAs and RNA-binding proteins (RBPs), regulate the encoded protein's abundance, localization, and integration into protein complexes. APA modulates protein function in a context-specific, gene-specific, or cell-type-specific manner and is critically involved in a variety of cellular processes; indeed, numerous human diseases including cancer and neurological disorders are associated with APA deregulation. 3' UTR genetic variants contribute to a substantial number of phenotypic traits and disease heritability, making APA a possible actionable target for therapeutic intervention.

The tissue- or context-specific regulation of APA is mediated through the activity of effectors such as transcription factors or RBPs. For example, in animals from flies to humans, the neuron-specific ELAV/Hu proteins inhibit splice and proximal polyadenylation (poly(A)) site usage to mediate the formation of neuronal 3' UTRs. Depending on cellular context, transcription elongation and termination factors interact with the cleavage and

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Figure 1. An accurate, comprehensive, full-length *Drosophila* transcriptome

(A) Combined isoform assembly (CIA) experimental and computational workflow. Long-read sequencing was performed on three *Drosophila* tissues: adult heads, embryos at the developmental time points 14–16 h after egg laying (AEL) and 18–20 h AEL, and adult ovaries. Transcript size selection was performed to optimize recovery of neuronal transcripts. The final transcriptome assembly was built on full-length reads, i.e., those that spanned an entire mRNA transcript isoform from experimentally validated 5' end to experimentally validated 3' end. Individual reads are represented as straight lines spanning different regions of the gene.

(B) BluePippin size selection considerably increased ONT cDNA read length (top) and optimized recovery of neuronal transcripts, whose length (bottom) exceeds the coverage range of LRS experiments without size selection (gray).

(C) Nucleotide composition profile of LRS reads at the 3’ end cleavage site for CIA full-length reads, compared with 3’ ends of discarded reads.

(D) Distribution as a function of transcript length of novel and previously annotated isoforms in the CIA transcriptome assembly dataset.

(E) Venn diagram showing the number of transcript isoforms (and genes) identified in each tissue in the CIA dataset, scaled by the number of isoforms. Data from different embryonic time points were pooled in (E) and (F).

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polyadenylation (CPA) machinery to enhance or inhibit 3' end processing.\textsuperscript{3–10} The gene-specific regulation of APA is less well understood. Alternative 3' UTR formation in individual mRNAs was shown to depend on sequence elements located in promoters or enhancers.\textsuperscript{11,12} Several studies provide evidence of a physical connection between transcription start sites (TSSs) and poly(A) sites (PASs); RBPs pervasively associate with promoter regions, as does the CPA machinery.\textsuperscript{13–16} Moreover, DNA methylation and CTCF recruitment influence APA,\textsuperscript{17} and gene loops affect alternative 3' end processing in yeast,\textsuperscript{18} indicating a possible role for chromatin looping in 3' end site selection. Together, such observations suggest that transcription regulation at promoters may be functionally coupled with APA; however, whether TSSs globally influence the selection of PASs remains unknown.\textsuperscript{19}

The main challenge in determining the regulatory links that mediate the choice of transcription initiation, splicing, and termination sites has been the ability to correlate different regions of a single transcript to one another—in particular, the 5' end and the 3' end of the same mRNA molecule, which typically lie several kilobases (kb) apart. Long-read sequencing (LRS) technologies now allow for full delineation of individual mRNA isoforms: in a single read, transcript coverage can be achieved from 5' to 3' end.\textsuperscript{20,21} LRS has been successfully used for the discovery of novel transcripts from repetitive regions, detection of novel splice variants, identification of interactions between alternative promoters and splicing of promoter-proximal exons, and for the identification of coupling events in feature pairs including TSSs, exons, and PASs.\textsuperscript{22–28} Short-read sequencing and LRS of nascent RNAs have shed light on intertwined co-transcriptional processes\textsuperscript{29,30} and demonstrated, for example, the influence of splicing dynamics on CPA efficiency.\textsuperscript{31,32} indicating a widespread interdependency between alternative transcription and RNA processing. However, so far, technologies have failed to resolve the link between 5' ends and 3' ends. Transcript isoform sequencing approaches that concurrently determine the start and end sites of individual RNA molecules, although well suited for determining transcript boundaries and their combinations,\textsuperscript{33} have not been employed to quantify couplings between 5' and 3' ends. Major limitations have indeed precluded the systematic analysis of the regulatory relationship between transcription initiation and termination. LRS read distributions typically peak at 1–2 kb in length, resulting in truncations, underrepresentation of long isoforms, and 5' or 3' sequencing biases.\textsuperscript{22,34} As a result, due to the incomplete representation of full-length mRNA isoforms, it has not been possible to quantify the contribution of different TSSs of the same gene to the expression of distinct 3' ends.

Here, we analyze the co-occurrence of mRNA features at the isoform level in the Drosophila nervous system, which is characterized by a particularly diverse transcriptome. We used multiple LRS approaches and developed a framework to accurately assess and quantify mRNA isoform usage, including the definition of true PASs. Our data demonstrate coupling between transcript 5' ends and 3' ends. We identify “dominant” promoters that, characterized by a unique epigenetic signature, outcompete cognate promoters to drive the expression of alternative, usually more distal, 3' ends. Promoter dominance is widespread in Drosophila brains and human cerebral organoids and constitutes a major mechanism to regulate 3' end site choice during transcription to generate select 5' UTR–3' UTR combinations in mature mRNAs.

**RESULTS**

A combined isoform assembly reflects the Drosophila transcriptome

To examine regulatory links between transcription initiation, exon usage, and APA in Drosophila, we first developed a comprehensive LRS isoform annotation approach (Figure 1A). In order to span the maximum range of the coding transcriptome, we used adult brains—the animal tissue with the greatest mRNA isoform diversity and where mRNAs reach their most extreme lengths\textsuperscript{35,36}—as well as embryos at different developmental stages (14–16 and 18–20 h after egg laying [AEL]), and adult ovaries (Table S1). Critically, we size-selected mRNAs (enriching for transcripts >3 kb) using Sage Science BluePippin. We performed Oxford Nanopore Technologies (ONT) cDNA sequencing as well as Pacific Biosciences (PacBio) Iso-seq.\textsuperscript{23} Both LRS approaches use reverse transcription on polyadenylated RNAs and PCR amplification followed by sequencing through a nanopore (ONT cDNA) or single-molecule real-time (SMRT) technology (Iso-seq).

Internal priming and RT template switching cause misidentification of 3' ends in most short-read and LRS approaches.\textsuperscript{21} To avoid these artifacts, we applied ONT direct RNA sequencing (DRS)\textsuperscript{17} and full-length poly(A) and mRNA sequencing (FLAM-seq).\textsuperscript{38} two independent LRS methods that detect the very end of poly(A) tails, and we defined the RNA cleavage site with nucleotide resolution. For a high-precision, high-coverage annotation of Drosophila TSSs, we used the Eukaryotic Promoter Database (EPD), a library of RNA polymerase II (RNA Pol II) promoters for which the TSSs were determined experimentally, usually by cap analysis of gene Expression (CAGE) or global run-on (GRO-cap).\textsuperscript{39} We found it crucial to only consider high-quality reads that span entire mRNA isoforms, from 5' end to 3' end. We assembled reads from each of the sequencing methods individually using full-length alternative isoform analysis of RNA (FLAIR).\textsuperscript{40} Each assembly was refined to retain only transcripts with a TSS represented in the EPD, and whose 3' end fell within a FLAM-seq or DRS cluster (Figures 1A, S1A, and S1B), thereby filtering out close to two thirds of all putative transcripts (Tables S1–S3). The remaining transcripts were assembled into a combined isoform assembly (CIA). We detected transcripts with mean read lengths over 4 kb and obtained high full-length coverage of long and ultra-long transcripts typical of the nervous system (Figures 1B, S1C, and S1D). Gene expression estimates

(F) CIA annotation tracks of detected *pumilio* mRNA isoforms in each tissue. Isoforms common to multiple tissues are depicted in gray. Boxes and lines represent exons and introns, respectively. Some introns (dashed lines) are not drawn to scale. TSSs and PASs are represented by arrows and gray stripes, respectively, in the gene model. Replicates per tissue: ONT cDNA: heads, n = 6; embryos 14–16 h, n = 3; embryos 18–20 h, n = 3; ovaries, n = 3. FLAM-seq and Iso-seq: heads, n = 3; DRS: heads, n = 1, embryos 14–16 h, n = 3; ovaries, n = 3. See also Figures S1 and S2 and Tables S1–S3.
from CIA transcripts were highly consistent with those assessed by short-read mRNA-seq in each tissue. In contrast, gene expression estimates assessed from nanopore sequencing on non-size-selected transcripts or DRS displayed substantial deviations from the gold standard method (Figure S1E), showing that size selection, rather than biasing toward longer transcripts, allowed for a better representation of tissue transcriptomes.

To assess the quality of full-length reads, we analyzed CIA 5' ends and 3' ends. 5' end pile-ups of ONT cDNA reads coincided with TSSs annotated in the EPD in 80% of cases; non-overlapping pile-ups fell within distal gene regions, usually 3' UTRs, and lacked distinctive TSS features such as RNA Pol II ChIP-seq and ATAC-seq peaks (Figures S1F–S1H), indicating high accuracy of Drosophila 5' end annotation in the EPD. CIA 3' ends harbor the characteristic, defined nucleotide composition at the cleavage site, whereas filtered-out 3' ends display noisy A-rich distributions reminiscent of sites of internal priming (Figure 1C). Strikingly, 3' ends unique to the Ensembl reference globally displayed a noisy nucleotide distribution, indicating that many reference 3' ends are mis-annotated (Figures S1I–S1L). We conclude that our stringent DRS- and FLAM-seq-guided filtering effectively identified false 3' ends. Thus, we generated a Drosophila mRNA isoform atlas, with 59,970 high-confidence, full-length transcripts. This CIA atlas that represents differential expression and poly(A) tail length of each mRNA isoform in heads, ovaries, and embryos can be accessed at https://hilgerslab.shinyapps.io/ciaTranscriptome.

We identified over 30,000 previously undescribed mRNA isoforms. Novel splice variants harbored canonical splicing signals and therefore likely arose from new combinations of known splice sites. In contrast, nearly 9,000 isoforms were characterized by unannotated 3' end sites (Figures S2A–S2E). Strikingly, isoform novelty drastically increased with transcript length, especially in heads and embryos, two tissues that contain neurons (Figures 1D and S2C), confirming the improved detection of long isoforms of neuronal mRNAs. CIA mRNA isoforms originate from 11,310 genes, 5,020 of which were found to be expressed in all three analyzed tissues. Interestingly, over 80% of these genes are expressed at least one identical isoform in all three tissues; although most genes expressed in heads were also expressed in other tissues, most CIA isoforms (35,170 out of 59,970) were found exclusively in head samples (Figures 1E, 1F, and S2F). We sequenced neural tissues much more deeply than ovaries and embryos (Table S1), which contributed to, but did not solely account for, the disproportionate representation of brain isoforms (Figure S2G). Our data are consistent with the neural-specific splicing pattern complexity described by modENCODE and further illuminate the astonishing isoform diversity of the nervous system.

We next investigated ultra-long mRNAs (>5 kb) of the nervous system more closely. Compared with ovaries and embryos, 3' UTRs disproportionately contribute to transcript length in head tissue (Figure S2H), consistent with the nervous-system-specific 3' UTR lengthening seen in multiple animal models. Moreover, nervous system transcripts display surprisingly long poly(A) tails, with their size increasing with transcript length (Figures S2I and S2J). This trend in flies has also been described in human cells and C. elegans, and it suggests a conserved coupling between distal PAS selection and tail length, possibly reflecting the result of distinct turnover kinetics and a potential role for long poly(A) tails in the protection of ultra-long transcripts.

**Coupling between transcript 5' ends and 3' ends**
The CIA transcriptome now allows us to quantify the co-occurrence of distinct co-transcriptional events in full-length mRNA isoforms. We focused on the analysis of regulatory relationships between transcription initiation and transcription termination. First, we categorized genes based on the number of identified TSSs and PASs in the CIA dataset (Figure 2A). We found that genes with alternative TSS usage (ATSSs) undergo APA disproportionately often, and vice versa (Figures 2B and S2K); moreover, 3' end diversity increases as a function of TSS number, and vice versa (Figures S2L and S2M). This could suggest that ATSSs have evolved to drive the production of distinct 3' ends. To study couplings between TSSs and PASs, we quantified the differential use of 3' ends as a function of the 5' end with which they are associated. We term a "5'-3' isof orm," a combination of 5' end and 3' end, i.e., a co-occurrence of any 5' end and 3' end in the same full-length CIA transcript. Importantly, many of the 5'-3' isoforms we detected in our sensitive LRS approach may have resulted from unproductive transcription and represent "noise" rather than biologically relevant isoforms. To eliminate these isoforms, we used an expression cutoff of >2 transcripts per million (TPM). We found over 16,000 5'-3' isoforms, almost 7,000 of which were novel (Figure S2N). We subsampled ONT cDNA reads and assessed the number of identified 5'-3' isoforms for each fraction and for different expression categories. Above cutoff, we reached near-saturation of 5'-3' isoform detection, even for genes with multiple TSSs and multiple PASs (ATSS-APA genes) (Figure S2O), strongly suggesting that our analysis faithfully represents the 5'-3' isoform landscape in Drosophila tissues.

**TSSs drive the selection of tissue-enriched 3' end sites**
To assess whether APA is driven by the use of distinct TSSs, we first asked whether tissue-specific 3' end expression is associated with tissue-specific 5' ends. Ovaries and heads constitute the two tissues at the extremes of the APA spectrum, with shifts toward proximal and distal PAS selection, respectively. We calculated differential 3' end and 5' end expression between the two tissues to identify "nervous-system 3' ends" and "ovary 3' ends," and we then assessed differential 5'-3' isoform expression in genes expressed in both tissues (Figure S2P). We discovered that for over half of all ATSS-APA genes, at least one 5'-3' isoform is enriched in one tissue compared with the other, representing a significant 5'-3' link (Figure 2C; Table S4), and distinct TSSs are specifically associated with 3' ends with differential expression between the two tissues (Figure 2D). Moreover, almost half of all nervous-system 3' ends were specifically expressed from a nervous-system TSS, and vice versa (Figure S2Q). In genes with several significant 5'-3' links, we observe, almost always, a pattern of bidirectionality in which one 5'-3' isoform is enriched in heads while the other is enriched in ovaries (Figures 2E, 2F, and S2R). Our results show that ovary- and head-specific PAS usage is linked to the alternative use of TSSs and suggest that TSSs influence PAS selection.
To functionally test this hypothesis in vivo, we used the CRISPR transcriptional activator (CRISPRa) system, in which a catalytically dead Cas9 (dCas9) fused to the VPR activation domain can be recruited to the upstream TSSs of individual genes by single-guide RNAs (sgRNAs). We tested all “bidirectional” genes for which a sgRNA strain was available (53 genes) and in which the upstream TSS was head- or ovary-enriched (23 TSSs, Figure 2G; Table S5). TSS activation failed in ovaries for all tested sgRNAs except one; in heads, we obtained significant gene activation for six nervous-system TSSs. In all cases, activation of the nervous-system TSS caused a specific increase in the expression of the linked, nervous-system 3’ end (Figures 2H and S2S). Notably, induction of the Malvolio (Mvl) nervous-system TSS in ovaries caused the ectopic expression of the linked, nervous-system 3’ end, demonstrating that specific TSS activation is sufficient to drive atypical 5’-3’ isoform expression (Figure 2I). Our data thus show that the site of transcription initiation drives head-specific 3’ UTR usage, thereby crucially contributing to the establishment of the distinct 3’ UTR landscape of the nervous system.

Co-expression of multiple 5’-3’ isoforms in neuronal cell types

The coordination between tissue-specific TSSs and APA could be mediated by tissue-specific trans-factors; for example, the
A

stai gene model

stai nanopore long reads

Act5C gene model

Act5C nanopore long reads

Proximal PAS

Distal PAS

TSS1

TSS2

B

LATER pipeline

Select ATSS-APA genes

Frequency of TSS linkage for each PAS

PAS1 PAS2 PAS3

Multinomial test

Identify significant biases

Promoter dominance score

C

TSS bias in ATSS-APA genes

No significant TSS bias

Promoter dominance

Log10(expected frequency) vs Log10(observed frequency)

D

Single TSS genes

Dominant promoter genes

AAUAAA None

AAUAAA None

E

ATSS-AS genes

AS-APA genes

ATSS-APA genes

Proportion of genes

0 0.5 1

Significant link

No link

F

TSS-PAS link

exon-PAS link

Dom P

Proportion of genes

0 0.5 1

Also exon-PAS link

Also TSS-PAS link

No additional link

G

Genes with AS-APA linkage

AS-APA linkage score

Other AS-APA genes

Dom P genes

H

Dom P

Proximal Distal

Deletion

orb TSS contributions to PAS expression

Dom P Proximal Distal

Legend:

orb

Control

Normalized expression

Dom P Proximal Distal

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pan-neuronal RBP ELAV promotes APA of individual genes in a TSS-dependent manner. To explore the regulation of co-transcriptional processing independently of the cellular environment, we investigated 5’-3’ links at the gene level in a single tissue—the brain—in which ATSS and APA are particularly abundant. Since APA isoform usage displays cell-to-cell heterogeneity, and some 3’ ends can be specific to certain cell populations, we assessed whether the 5’-3’ links that we identified in Drosophila heads tend to be expressed in the same cell, or whether on the contrary, distinct isoforms are exclusive to different cell types. Using the Drosophila brain atlas, we evaluated every CIA 3’ end at the single-cell level and quantified the co-occurrence of different 3’ ends of the same gene in each of the 177 cell types described in the dataset. We found that the majority of ATSS-APA genes are co-expressed as several APA isoforms in most cell types, and we did not detect a general trend of mutually exclusive 3’ end isoform expression within the brain (Figures S3A–S3C). We conclude that differential usage of TSSs and PASs can occur within the same cell type, independently of tissue-specific or cell-type-specific factors. Hence, we can use the nervous system 5’-3’ isoform dataset to probe PAS preference within the same cell populations.

**Global bias of 3’ end site selection depending on the TSS**

The identification of full-length gene isoforms of ATSS-APA genes in heads revealed that in many cases (e.g., stai), distinct PASs were preferentially associated with specific TSSs, while for other genes (e.g., Act5C) there was no such bias (Figure 3A). We set out to assess whether the competitive use of PASs is regulated at the site of transcription start. To discern regulatory links transcriptome-wide between transcription start and 3’ end formation, we developed the computational framework long-reads-based alternative termination estimation and recognition (LATER) (Figure 3B). For all ATSS-APA genes, for a given PAS, we calculated the frequency of association of each TSS with the expression of the associated 3’ end (Figures 3A, 3B, and 3D). We defined two modes of 3’ end site selection in ATSS-APA genes: “TSS-unbiased,” in which the association frequencies of distinct TSSs with a given 3’ end did not significantly differ; and “promoter dominance,” in which one TSS was disproportionately associated with the expression of a specific 3’ end. Strikingly, deviations from the expected proportions were the rule rather than the exception, with most (55%) ATSS-APA genes displaying promoter dominance in at least one tissue (Figures 3C, S3E, and S3F; Table S4).

Highly expressed genes displayed predominantly short 3’ UTRs, and stronger promoters were found to favor the selection of proximal PASs in reporter assays, consistent with the idea that high transcriptional activity enhances 3’ end processing on a first-come, first-served basis. In contrast, a fast RNA Pol II elongation rate correlates with the use of more distal PAS in yeast. However, we did not observe any significant difference in expression levels of isoforms from our identified dominant promoters (Figure S3G); importantly, full-length 5’-3’ isoform detection and categorization as dominant-promoter-isoform were not biased by read length for transcripts up to 10 kb long (Figures S3H and S3I). Therefore, transcript length or TSS strength cannot explain PAS selection in cases of promoter dominance. With the ability to quantitatively assess individual 5’-3’ isoforms, we demonstrate a global effect of TSS selection on differential 3’ end expression, causally linking transcription initiation to termination.

**Dominant promoters override strong poly(A) signals and constrain AS**

We asked whether dominant promoters showed a propensity to override well-defined rules of mRNA processing. For APA genes, differential 3’ end expression is thought to depend on PAS “strength”: unless specifically inhibited in trans, PASs containing the hexamer AAUAAA and variants thereof are rarely bypassed to produce a more distal 3’ end. For APA genes with a single promoter, the presence of the AAUAAA sequence was indeed a
predictor of proximal PAS usage in our dataset, and skipping of
the proximal PAS usually occurred in the absence of a poly(A)
signal. Strikingly, ATSS-APA genes with dominant promoters
showed the opposite trend; in fact, proximal PASs containing
AAUAAA were preferentially skipped in transcripts arising from
a dominant promoter (Figure 3D).

Next, we tested whether splicing plays a role in the observed
5′-3′ couplings, possibly representing the regulatory intermediate
between dominant promoters and 3′ end site selection. First,
we ensured that splice isoform coverage in long reads was suffi-
cient to assess exon-exon junction choice. Except for isoforms
identified with one single read, likely representing very rare or
aberrant variants, we reached saturation of splice isoform detec-
tion (Figures S3J and S3K). We developed long-reads-based AS
estimation and recognition (LASER), based on the same princi-
ple as LATER (Figure S3L), to identify disproportionate associ-
ation frequencies between distinct TSSs and exon-exon junc-
tions—“TSS-exon links”—as well as between exon-exon
junctions and PASs—“exon-PAS links.” Compared with TSS-
PAS links (promoter dominance), we identified surprisingly little
coupling between AS and APA, with significant links in about
10% of AS-APA genes (Figures 3E, S3M, and S3N; Table S4).
A significant link between AS and 3′ end site selection was seen
in about one-third of genes with a dominant promoter; for
example, stai exons 4 and 8 are near-mutually exclusively
associated with distinct PASs and their respective dominant
promoters (Figures 3A and 3F). This enrichment, but lack of sys-
tematic association of AS with APA led us to hypothesize that
exon-PAS couplings are a consequence, not a causal interme-
diate, of the influence of dominant promoters on co-transcriptional
processing. Indeed, we find that in ATSS-APA genes, exon-PAS
links almost always (88%) occur when transcription starts from a
dominant promoter. Moreover, exon-PAS links are significantly
weaker in the absence of a dominant promoter (Figures 3F and
3G). We conclude that in ATSS-APA genes, AS does not repres-
ent a necessary intermediate step for biased 3′ end selection
by dominant promoters, although it may influence APA in individ-
ual cases. Together, our findings indicate that sites of transcrip-
tion initiation direct APA independently of poly(A) signal strength
and also impose a constraint on other RNA processing events
such as splicing.

To functionally validate 5′-3′ links and verify that 3′ end choice is
mediated by dominant promoters in vivo, we generated the fly
mutant orbΔDP, in which the dominant promoter of the gene orb
was specifically deleted by CRISPR-Cas9-mediated gene editing.
Orb possesses two 3′ ends and three TSSs, with the first
TSS dominantly associated with the distal-most 3′ end (Fig-
ure 3H). In orbΔDP embryos, expression of the distal but not the
proximal 3′ end was massively depleted (Figure 3I). Our data
thus show that dominant promoters influence PAS selection and
can mediate skipping of canonical poly(A) signals to favor
more distal sites of transcription termination.

3′ end site selection through promoter dominance
impacts transcriptome identity and gene function

To assess the functional importance of the reported connection
between TSSs and PASs, we first sought to determine if it is
evolutionarily conserved. We found that 5′ UTRs transcribed
from dominant promoters and 3′ UTR sequences generated via
dominant-promoter-associated PASs (“dominant-promoter-3′
UTRs”) were more conserved than their non-dominant and un-
linked counterparts, respectively (Figures 4A and 4B). Following
the notion that functional interactions can be detected through
evolutionary couplings,55 we performed a mutual information
analysis56 to test whether 3′ end site regions and their dominant
promoters mutate jointly to maintain genetic interactions. We
calculated the co-evolution score for each pair of nucleotide posi-
tions within the gene stat. Strikingly, a cluster of high-scoring
nucleotide pairs could be identified between 3′ UTR sequences
and regulatory regions upstream of the linked dominant prom-
oter, but not the non-dominant promoter. Act5C, a gene with
no TSS bias, did not display any 5′-3′ co-evolution clusters (Fig-
ure 4C). We performed a more global analysis, selecting 100
ATSS-APA genes (top and bottom 50 by promoter dominance p
value), and scored, for each gene, co-evolution clusters in
nucleotide pair matrices between 5′ end regions (TSS — 1 kb)
and the 3′ end region (3′ UTR). We found that co-evolution scores
were significantly higher for dominant promoters, compared with
other TSSs; most dominant-promoter genes, but not TSS-unbi-
ased genes, showed strong co-evolution between 5′ end and
associated 3′ end sequences (Figures 4D and 4E). Our results
show not only that sequences generated directly (5′ UTRs) or
indirectly (linked 3′ UTRs) from dominant promoters are
conserved but also that evolutionary pressure maintains the
link between them.

We next computationally predicted the consequence of
disrupting TSS-PAS links and the ensuing 3′ end mis-selec-
tion. In Drosophila heads, differential 3′ end site selection by
dominant promoters results in a change in protein-CDS, 3′
UTR lengthening, and 3′ UTR shortening in 40%, 42%, and
18% of cases, respectively. A substantial amount of regulatory
3′ UTR sequence is gained through dominant-promoter-
mediated 3′ UTR lengthening (Figure S4A); we sought to quan-
tify the influence of dominant promoters by computing the
occurrence, in either 3′ UTR isoform, of potential binding sites
for neuronal RBPs and microRNAs highly conserved and
enriched in fly heads, since these are more likely to exert a
functionally relevant effect on target mRNAs.57 Interestingly,
putative binding motifs for miR-277, a microRNA involved in synapto-
genesis with a possible role in neurodegeneration,58,59 were the
most impacted by dominant-promoter-mediated 3′ UTR
lengthening (Figure S4B). In addition, dominant-promoter 3′
UTRs were enriched in putative binding sites for RBPs well
known for specialized neuronal roles, such as pumilio (Pum) and
alan shepard (Shep), as well as for mir-2279, a poorly
expressed and conserved microRNA that is nonetheless pre-
dicted to target neural pathways related to axonal projections
(Figures S4C–S4F). This indicates that dominant-promoter-
associated 3′ UTR sequences function in the regulation of the
encoded protein in an isoform-specific manner; our ana-
lyses predict that disruption of conserved TSS-PAS links
causes a widespread mis-selection of 3′ end sites, resulting in
loss of tissue-specific protein isoforms and 3′ UTR- medi-
ated regulation by microRNAs and RBPs, strongly suggesting
that regulation through dominant promoters is functionally
relevant for animal fitness.
A combination of epigenetic features defines the chromatin environment of dominant promoters

One possible interpretation of the observed 5'-3' coupling is that dominant promoters possess a characteristic that subjects the nascent transcript to modified rules of co-transcriptional processing. Splicing and 3' end cleavage have been shown to be influenced by the presence of particular chromatin elements at the sites of transcription initiation and termination, respectively.60

We set out to identify whether dominant promoters possess a common regulatory feature that mediates coupling between TSS and PAS. We analyzed ChIP-seq data generated in Drosophila heads (modENCODE61) to assess the in vivo location of over 40 histone marks, histone variants, and transcription factor binding sites. We found that promoter regions of ATSS-APA genes, while not displaying any notable enrichment in RNA Pol II or common repressive or active chromatin marks, were strongly depleted for the histone variant H2A.Z. Conversely, acetylation of histone H3 at lysine 18 (H3K18Ac), a histone mark associated with gene activation and transcriptional priming in developmental transitions,62 was specifically enriched around the TSS of ATSS-APA genes (Figure S5A).

We grouped TSS regions genome-wide according to H2A.Z and H3K18ac ChIP-seq signal, which generated five clusters of distinct H2A.Z and H3K18 patterns. Cluster 1 and cluster 2 were characterized by H2A.Z depletions concomitant with H3K18Ac enrichment. Strikingly, those two clusters included significantly more dominant promoters than the other three clusters (Figures 5A and 5B; Table S4), suggesting that H2A.Z depletion and H3K18Ac enrichment are common characteristics of dominant promoters. Next, we assessed transcription factor binding at the TSS and linked 3' end of dominant promoter genes in fly heads, using the ReMap 2022 database.63 We found coupled enrichment of 20 factors at both transcription initiation and termination sites of these genes (Figures S5B–S5D); most...
interestingly, the highly conserved acetyltransferase Nejire (Nej, also known as p300 or CREB-binding protein, CBP) was the factor most frequently found at dominant promoters and at their associated 3' end (Figures 5C and 5D). Fly and mammalian CBP promote the proper deposition of H3K18Ac, the histone mark we found enriched around dominant promoters. Together, our data thus indicate that dominant promoters of ATSS-APA genes are characterized by a specific epigenetic landscape, partially established by the presence of CBP.

p300/CBP mediates dominant-promoter-driven 3' end site selection
To test whether CBP is instructive for the selection of alternative PASs, we performed mRNA-seq and assessed 3' end usage in two independent CBP mutants. We used 14- to 16-h embryos, a stage at which maternally deposited CBP was depleted but embryos still showed a normal gross morphology. The absence of zygotic CBP caused a widespread impairment of the embryonic 3' end landscape: 21% of all expressed APA genes displayed a change in 3' end site selection, characterized by a significant upregulation or downregulation of RNA expression downstream of the proximal PAS, compared with upstream regions (Figure 5E). Strikingly, affected genes are those that display, in wild-type flies, CBP ChIP signal at both the TSS and the associated PAS (clusters 1 and 2), whereas APA was largely unaffected in genes where CBP signal was only found at the TSS (cluster 3. Figure 5E: Table S4). PAS shifts were more frequent and more pronounced in dominant-promoter genes compared with TSS-unbiased genes (Figures 5F–5H), demonstrating that p300/CBP mediates, at least partially, dominant-promoter-driven 3' end site selection. In contrast, mutation of one of three other factors we had found enriched at the TSS and PAS of dominant promoter genes—Enhancer of zeste (E(z)), Deformed (Dfd), and Posterior sex combs (Psc)—had little to no effect on PAS usage (Figure S5E). We propose that in addition to CBP, other factors are involved in the promoter-mediated regulation of APA, both globally and on a gene-by-gene basis. Such factors may include chromatin modifiers, AS regulators, and transcription factors.

TSS influence on isoform choice is a conserved regulatory mechanism
To assess whether TSS-mediated PAS selection is conserved in mammals, we performed an LRS-based analysis in human cerebral organoids, an in vitro model of the human brain. Coupling FLAM-seq with ONT cDNA sequencing and size selection, we generated an organoid CIA dataset including many novel long mRNA isoforms and defined highly accurate 5’-3’ isoforms in ATSS-APA genes (Figures 6A and S6A; Tables S1–S3). Since FLAM-seq identified only 16,840 3' end sites, we performed 3' end sequencing (3'-seq) and predicted further confident 3' end sites based on the nucleotide composition of FLAM 3' ends, thereby substantially expanding the 3' end database (see STAR Methods). Similar to Drosophila, in human organoids the presence of ATSSs was associated with APA (Figure 6B). We applied LATER to the human dataset and found that over a third of ATSS-APA genes display a TSS bias, in which 3' end choice is influenced by the promoter (Figures 6C and S6B; Table S4), in many cases mediated by skipping of the proximal canonical poly(A) signal (Figures 6D and 6E). The lack of ChIP-seq data from human neural tissue prevented us from identifying a clear TSS signature of dominant promoters, as we did in Drosophila. However, we performed a transcription factor enrichment analysis using the ReMap 2022 database and found that factors displaying an association with APA, such as FOXA1 and p300/CBP, were enriched at dominant promoters and/or linked 3' ends also in human cells (Figure S6C). We conclude that dominant promoters apply a conserved transcriptional constraint on isoform choice, often mediating the usage of more distal PASs. The epigenetic signatures at these sites may have evolved to aid in the recruitment of transcription and processing factors—including p300/CBP—that execute this program, which is determined at the time of transcription initiation.

DISCUSSION
Over the past decades, a rich body of work has described coupling mechanisms that coordinate transcription with splicing67,68; for example, a role for promoter identity, RNA

Figure 5. TSSs exert promoter dominance through specific chromatin signatures
(A) Heatmaps representing ChIP-seq signal at TSS ± 2 kb genome-wide for the histone variant H2A.Z and the histone mark H3K18Ac. Genes are grouped by k-means clustering, using both H2A.Z and H3K18Ac signal. On average, ATSS-APA genes measure 20,010 kb from TSS to distal PAS.
(B) For each cluster, the proportion of dominant promoters (in ATSS-APA genes) is shown. ***p < 2.2e–16 and *p = 0.006 (two-tailed Fisher’s exact test).
(C) ChIP-seq peak enrichment analysis of the TSS and PAS of dominant promoter isoforms. Factors significantly enriched (adj. p value < 0.01) at both TSS ± 150 nt and PAS ± 150 nt are shown, ranked by the ratio of total peaks that map to dominant promoters.
(D) CBP ChIP-seq signal and full-length reads representing distinct 5’-3’ isoforms of the gene Calnexin (Caln). The dominant promoter, its associated PAS, and long reads of the corresponding 5’-3’ isoform are colored in red. CBP ChIP-seq data from Drosophila heads are shown as a log2 ratio normalized to input.
(E) Enrichment of CBP ChIP-seq signal at transcript TSSs (+2 kb) and proximal PASs (+2 kb) and RNA expression of distal 3’ UTRs in CBP mutants (two independent alleles, nej3 and nej77), compared with control embryos. mRNA-seq heatmaps and profile plots display 0.5 kb upstream of the proximal PAS (prox) and the distal 3’ UTR downstream (distal, scaled region). Genes are grouped into three clusters by k-means clustering, using both CBP ChIP-seq signal at proximal PASs and mRNA-seq signal in nej3 and nej77 mutants. mRNA-seq was performed on RNA extracted from 14- to 16-h AEL embryos in four biological replicates for each genotype.
(F) mRNA-seq signal tracks of the gene chicadex (chic), whose distal PAS selection depends on p300/CBP. Dominant promoters for each PAS are indicated in the respective color. ChIP-seq data from Drosophila heads are from modENCODE.
(G) 3’ end selection change in CBP mutant embryos (nej77), calculated as the change in mRNA expression of the distal transcript regions, compared with control embryos, for PASs linked to a dominant promoter (Dom P) and those with no TSS bias. ***p = 6.7e–8 (two-tailed Student’s t test).
(H) Proportion of ATSS-APA genes with (Dom P) or without (no TSS bias) dominant promoters in which 3’ end selection was significantly affected (p < 0.05, Wald test) in CBP mutant embryos (nej77). ***p = 3.5e–11 (two-tailed Fisher’s exact test).
See also Figure S5 and Table S4.
Pol II kinetics, and transcription factors was demonstrated in defining splice site choice. In comparison, our knowledge on links between transcription initiation and APA was very limited. In this work, we provide an integrated view of mRNA features and their association in individual transcripts. Our data will serve as a useful resource to study alternative RNA processing, poly(A) tail lengths, RNA modifications, and the interrelation of these features in a tissue-dependent manner. Our finding that 3' end site selection depends on TSS choice has broad implications for the study of gene expression and its role in disease. It is well established that the use of distinct 3' end sites contributes to important gene expression programs, including those involved in developmental transitions, tissue identity, and the cell cycle; APA deregulation is associated with numerous human pathologies, most notably cancer. We hypothesize that the regulation of isoform expression by the use of ATSSs is a central mechanism to ensure tissue function and identity.

Given the pattern of bidirectional 5'-3' isoform production we found when comparing tissues, it is evident that both 5' ends with 3' ends may represent a cellular strategy to ensure the co-occurrence of particular 5' UTR and 3' UTR elements in the same mRNA molecule. Post-transcriptional gene regulation including mRNA localization, stabilization, and translation depends not only on the sequence and structural elements found in 5' and 3' UTRs, but also on 5'–3' communication, either through physical proximity mediated by the concomitant binding of RBPs to both RNA ends (closed-loop model) or through indirect interactions. Hence, dominant promoters may act to enhance these intramolecular interactions to regulate mRNA expression.

At dominant promoters, H2A.Z depletion, indicative of high transcription rates, frequent chromatin interactions, and lower nucleosome definition synergizes with the enrichment of the active histone mark H3K18Ac, which was shown to help prime genes for activation during developmental transitions; such increased chromatin accessibility at the TSS and PAS may enhance 5'-3' coupling and the controlled differential expression of distinct mRNA isoforms. CBP may also link 5' and 3' ends independently of its established role in H3K18Ac.

**Figure 6.** Dominant promoters drive PAS selection in human brain organoids

(A) Organoid CIA assembly pipeline. The distribution of novel and previously annotated isoforms as a function of transcript length is indicated (n = 3).

(B) Proportion of genes that undergo APA in each TSS category. **p < 0.001 (two-tailed Fisher’s exact test).

(C) Identification of TSS biases in human brain organoids. The plot was generated as in Figure 3C. 38% of ATSS-APA genes show a significant bias (p < 0.01, chi-squared test with Monte Carlo simulation and Benjamini-Hochberg correction).

(D) Representative example of a gene with promoter dominance in brain organoids. Full-length reads represent distinct 5'-3' isoforms of the gene CAST. The dominant promoter, its associated PAS, and long reads of the corresponding 5'-3' isoform are colored in red in the gene model. Pie charts represent the contributions of each TSS to the expression of each 3' end.

(E) PAS usage when the canonical poly(A) signal AAUAAA is found within a 50-nt window of the most proximal PAS of the gene. Proximal and distal denote PASs found in the proximal 20% or distal 80% of the 3' UTR, respectively. **p < 0.001 (two-tailed Fisher’s exact test).

See also Figure S6 and Tables S1–S3 and S4.
deposition; concomitant binding of CBP molecules at the TSS and PAS could facilitate an intragenic loop, a mechanism that was proposed to connect transcription initiation with PAS choices. Additionally, we hypothesize that CBP mediates the recently recognized influence of distal cis-regulatory elements on APA, possibly by binding to enhancer RNAs (eRNAs), an interaction that stimulates histone acetylation and transcription of target genes. Gene topology may further distinguish the regulation of neuronal ATSS-APA genes. In mouse brains, “melting” chromatin states and distinct chromatin contact patterns were seen in long genes associated with specialized neuronal processes, and it is possible that such topological constraints contribute to 5’-3’ coupling. We propose that dominant promoters, by residing in a chromatin environment that dictates specialized regulation through enhanced protein interactions and possibly gene looping, promote communication between the transcription and RNA processing machineries. Interestingly, dominant promoters display typical characteristics of promoters of developmental genes, including lower nucleosome occupancy, CBP binding, and H3K18Ac. “Developmental core promoters” were previously defined as TSSs regulated by “developmental enhancers” that play a defining role in development-, tissue-, or context-specific gene regulation, in contrast with “housekeeping promoters.” Our results in the context of prior literature are therefore consistent with a model in which developmental genes employ specific epigenetic regulation evolved to ensure robust and highly regulated interactions not only between enhancers and promoters but also between promoters and PASs to dictate gene expression.

Coupling 5’ ends to 3’ ends of transcripts represents a conserved principle in the regulation of gene expression, with broad relevance, as APA affects mRNA coding potential, localization, stability, and translation to achieve context-specific modulation of developmental genes. The universal impact of alternative mRNA processing in the etiology of disease has been highlighted by the substantial association found between APA-altering SNPs in 3’ UTRs with human phenotypic traits and diseases, which can be further probed using variant expression-aware annotations and large LRS datasets of human tissues. Linking 5’ ends to disease-relevant mutations in 3’ UTRs will close an important gap in our understanding of genetic disease mechanisms, aid in the identification of disease-associated mutations in the full-length context in which they are deleterious, and may provide a platform to target variant-associated diseases.

Limitations of the study
We centered our analyses on the nervous system as a whole, as opposed to considering the complexity of its many different cell types. As a consequence, for genes with extreme isoform diversity and highly cell-type-specific isoform expression, only relatively abundant isoforms passed our stringent detection cutoff. Therefore, we expect that many functionally relevant mRNA isoforms went undetected. Our study uses BluePippin size selection prior to nanopore LRS. Although gene expression calculations from these data were highly consistent with those obtained through mRNA-seq, in individual cases, longer transcripts may be overrepresented. Full-length mRNA coverage from nanopore long reads substantially declined in transcripts exceeding 10 kb in size. Although we excluded isoforms exceeding that limit from quantitative analyses, they are still depicted in the CIA atlas, where they may be underrepresented, compared with significantly shorter mRNAs. Finally, the transcription factor binding analysis on human TSSs conducted with the ReMap 2022 database used ChIP-seq data from a variety of human cells: the results shown in Figure S6 likely incompletely represent binding in cerebral organoids.

STAR METHODS
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  - Motif enrichment in dominant-promoter-associated 3’ UTRs
We actively worked to promote gender balance in our reference list. While citing references scientifically relevant for this work, we also
acknowledge that one or more of the authors of this paper self-identifies as a gender minority in their field of research or within their geographical location. One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in their field of research.

We support inclusive, diverse, and equitable conduct of research. We worked to actively promote gender balance in our reference list.

The authors declare no competing interests.

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AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research. We worked to ensure sex balance in the selection of non-human subjects. One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in their field of research or within their geographical location. One or more of the authors of this paper self-identifies as a gender minority in their field of research. While citing references scientifically relevant for this work, we also actively worked to promote gender balance in our reference list.

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REFERENCES


### STAR★METHODS

#### KEY RESOURCES TABLE

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**Oligonucleotides**

Oligonucleotides used for RT-qPCR

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**CRISPR guide RNAs**

| **CRISPR guide RNAs** | **STAR Methods** | N/A |

**Software and algorithms**

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Valérie Hilgers (hilgers@ie-freiburg.mpg.de).

Materials availability
All plasmids and fly strains generated in this study are available from the lead contact without restriction.

Data and code availability
- All LRS and RNA-seq data have been deposited at NCBI Gene Expression Omnibus (GEO) and are publicly available as of the date of publication. Accession numbers are listed in the key resources table.
- All original code has been deposited at Zenodo and is publicly available. DOIs and GitHub links are listed in the key resources table.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Drosophila melanogaster
Experiments in this study used male and female (in equal amounts, except for experiments using ovaries) Drosophila melanogaster embryos and adult flies. Flies were raised at 25 °C. The CIA reference transcriptome was built using w1118 flies (Bloomington stock number 5905). Flies mutant for p300/cbp/nej (nej3 and nejEP1179),101,102 Enhancer of zeste (E(z)731),103 Posterior sex combs (Psch27 and Psce22),104 Deformed (Dfd1)105 and Spps (SppsG8810)106 were obtained from the Bloomington Drosophila Stock Center. We used CRISPR/Cas9-mediated genome editing following the procedure described in Port and Bullock107 to generate the orb dominant promoter deletion orbDP. Two guide RNAs (GAGAGAGCTCTACATCAGC, CGCCACGGCGTGCAACGCTG) targeted the orb promoter region, generating a 1.9 kb deletion beginning 40 bp upstream of the annotated TSS. All embryo injections were performed by Bestgene, Inc. Recessive lethal mutations were kept in heterozygosis over GFP-balancer alleles. In CRISPRa, to induce the expression of tissue-specific TSSs, TRiP-OE lines from the Transgenic RNAi Project46,108 were used. Flies expressing single guide RNAs (sgRNAs) targeting the upstream TSS of genes of interest (sgRNA, example genotype: y[1] sc[*] v[1] sev[21]; P[y[1] +t7.7 v[1] +t1.8]= TOE.GS02080)attP40) were crossed with flies expressing, under control of tubulin-Gal4, a catalytically dead Cas9 (dCas9) fused to the VP64 activation domain (Tub>dCas9-VPR, genotype: w; UAS:dCas9-VPR; tub-Gal4/SM5, TM6B). All fly strains are listed in the key resources table.

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Cell 186, 1–18.e1–e11, May 25, 2023 e5
Please cite this article in press as: Alfonso-Gonzalez et al., Sites of transcription initiation drive mRNA isoform selection, Cell (2023), https://doi.org/10.1016/j.cell.2023.04.012
Human cerebral organoids

iPSC-derived cerebral organoids were generated as described in Giandomenico et al., with some modifications. Briefly, after dissociation into a cell suspension with acutase, 6,000 cells were seeded per one well of 96-well plates in 100 μl of embryoid body medium (EBM: DMEM/F12, 20% Knockout replacement serum, 1x Glutamax, 1x MEM-NEAA, 2% ESC FBS, 50μM ROCK Inhibitor, 10 μM bFGF). On day four, the medium was replaced with EBM without bFGF and ROCK inhibitor. On day five, the medium was replaced with a neural induction medium (NIM: DMEM/F12, 1x N2 supplement, 1x Glutamax, 1x MEM-NEAA, 10μg/ml heparin solution). On day 7-9, the formed organoids were embedded into Matrigel (Coming, 356234) and kept in NIM for one day, and in 1:1 NIM: organoid differentiation medium (ODM: 1:1 DMEM/F12: Neurobasal, 1xN2 supplement, 1x B27- vitamin A supplement, insulin, 2-ME solution, Glutamax, MEM-NEAA) for one additional day, followed by four days in ODM. Next, the organoids were transferred to ultra-low attachment 6-well plates and cultured on an orbital shaker (85 rpm) in organoid maturation medium (OMM: 1:1 DMEM/F12: Neurobasal, N2 supplement, B27+ vitamin A supplement, insulin, 2-ME solution, Glutamax supplement, MEM-NEAA, Vitamin C solution, chemically defined lipid concentrate, BDNF, GDNF, cAMP, 1% Matrigel).

METHOD DETAILS

Sample collection for RNA analysis

For head transcriptomes, 3-day-old w1118 flies were collected and flash-frozen in liquid nitrogen and heads were homogenized in QiAzoL Lysis Reagent (QIAGEN 79306) for RNA extraction. For ovary transcriptomes, 3-day old w1118 virgin females were collected, and 20 ovaries per replicate were dissected and homogenized. For embryo transcriptomes, eggs from w1118 flies were collected for two hours on agar plates and aged for either 14h (14-16h AEL embryos), or 18h (18-20h AEL) at 25°C. 50 embryos per replicate were homogenized. For orbΔOP, nej1, nejFP1179, E[z]731, Psc[h27], Psc[c22], Dfd1 and Sps[G810] mutant analysis, eggs from mutant flies grown in heterozygosis with GFP-marked balancer chromosomes were collected for two hours on agar plates and aged for the appropriate amount of time at 25°C. Embryos were dechorionated following standard procedures and placed on a plate containing halocarbon oil. 20 to 30 mutant embryos were hand-scored according to morphology and against GFP signal, in at least three replicates. For the CRISPRa experiment, to obtain flies ubiquitously expressing dCas9 and a promoter-targeting sgRNA, tub>dCas9VPR virgin female flies were crossed with sgRNA males. Crosses were maintained at 25°C and parents were removed from the vial after two days. Eclosed progeny were aged for five days, selected against Tb and Cyo, and the heads and ovaries of five female flies per replicate were processed for RNA extraction. A sgRNA line targeting a non-Drosophila sequence was used as a control. Organoid RNA was prepared from 60-day-old cerebral organoids as described in Rybak-Wolf et al. Briefly, organoids were collected in TRIzol (Invitro-gen 15596026) and RNA was prepared with the Direct-zol RNA Miniprep kit (Zymo Research R2050) according to the manufacturer’s instructions.

RNA extraction and RT-qPCR

For all experiments, RNA was extracted using QiAzoL Lysis Reagent (QIAGEN 79306) according to the manufacturer’s instructions. Before library preparation, RNA integrity was analyzed using a 2100 Bioanalyzer (Agilent Technologies). Only RNAs with RQN values of 10 were used for all sequencing experiments. For RT-qPCR, 300 ng total RNA were used for reverse transcription with iScript gDNA Clear cDNA Synthesis Kit (Bio-Rad). RT-qPCR was performed in a LightCycler 480 II instrument using FastStart SYBR Green Master (Roche). RT-qPCR primer sequences are listed in Table S6. Before library preparation, RNA integrity was analyzed using a 2100 Bioanalyzer (Agilent Technologies). Only RNAs with RQN values of 10 were used for all sequencing experiments. For RT-qPCR, 300 ng total RNA were used for reverse transcription with iScript gDNA Clear cDNA Synthesis Kit (Bio-Rad). RT-qPCR was performed in a LightCycler 480 II instrument using FastStart SYBR Green Master (Roche). RT-qPCR primer sequences are listed in Table S6.

Short-read Sequencing (RNA-seq and 3′-seq)

Libraries for mRNA-seq were prepared from 3-day-old w1118 fly heads with 100 ng of total RNA using TruSeq Stranded mRNA Library Prep (Illumina 20020595) according to the manufacturer’s instructions. Libraries for total RNA-seq were prepared from dissected fly ovaries with 100 ng of total RNA using TruSeq Stranded total RNA Library Prep (Gold) (Illumina 20020599) according to the manufacturer’s instructions. Paired-end sequencing was performed using the NovaSeq6000 platform (Illumina) and 101-bp reads. mRNA-seq data from 14-16h AEL embryos and 18-20h AEL embryos are from Carrasco et al. Sequencing data were processed using the RNA-seq module from snakePipes, adding flags for –trim, -m “alignment-free,alignment”. Reads were mapped to the Drosophila melanogaster reference genome (Ensembl assembly release dm6), and the transcriptome reference annotation release-96 using STAR. 3′-seq libraries were prepared with 10 ng of total RNA using the QuantSeq 3′-seq Library Prep Kit REV (Lexogen) according to the manufacturer’s instructions. Paired-end sequencing was performed using the NovaSeq6000 platform (Illumina) and 101-bp reads.

Nanopore sequencing (ONT cDNA)

Nanopore sequencing was performed on 3-day-old w1118 fly heads, 14-16h AEL embryos, 18-20h AEL embryos, dissected fly ovaries, and human cerebral organoids. For generation of full-length CDNA libraries, polyadenylated RNA molecules were isolated from total RNA preparations using the NEB’s NEBNext Poly(A) mRNA Magnetic Isolation Module (NEB). Purified polyadenylated RNA molecules were used for library preparation using the cDNA-PCR Sequencing protocol (Oxford Nanopore Technologies). The following modifications were made to the protocol. To eliminate short reads from the final data, both input polyadenylated RNA molecules and cDNA molecules were cleaned upon further processing using AMPure XP beads (Beckman Coulter) using a
magnetic bead sample ratio of 0.4. To retain cDNA fragments > 3kb, the BluePippin device and appropriate separation DNA gel cassettes were used (Sage science). cDNA was amplified using 14 PCR cycles and 12 min extension time at 65°C. Libraries were sequenced on a MinION 1B or GridION sequencing device from Oxford Nanopore Technologies (R9.4.1). Reads were processed using guppy-5.0.7 (model: dna_r9.4.1_450bps_sup.cfg). Reads were aligned to the *Drosophila melanogaster* reference genome (Ensembl assembly release dm6) or to the *Homo sapiens* reference (GRCh38), and transcriptome reference annotation release-96 and release-91, respectively. For genomic reads, pairs were mapped using minimap2, with parameters “-minimap2 -ax splice -u f”. Alignment files were sorted and indexed using samtools v1.12. For transcriptome alignments, “minimap2 -ax map-ont -u f” was used.

**Nanopore Direct RNA sequencing (DRS)**

DRS was performed on 3-day-old *w*1118 fly heads, 14-16h AEL embryos, and dissected ovaries. Polyadenylated RNA molecules were isolated from total RNA preparations using the Dynabeads® mRNA Purification Kit (Invitrogen). Multiple poly-A+ pulldowns were pooled to reach 500 ng PolyA+ RNA input for library preparation using the Direct RNA sequencing kit (Oxford Nanopore Technologies). Libraries were sequenced on a MinION 1B or GridION sequencing device from Oxford Nanopore Technologies. Reads were processed using guppy-5.0.7 (model: rna_r9.4.1_70bps_hac.cfg).

**Iso-seq**

Iso-seq libraries were prepared using 500 ng total RNA from 3-day-old *w*1118 fly heads, processed with the Iso-seq express 2.0 workflow (PacBio) with 14 cycles of PCR amplification and size selection with the BluePippin system for transcripts larger than 3 kb according to the manufacturer’s protocol. After SMRTbell adapter addition, libraries were sequenced on three SMRTcells on a Sequel I PacBio sequencer. The raw data files were processed with SMRT Link v8 software to generate CCS fastq files. Data analysis was performed using the Iso-seq3 pipeline to generate consensus reads. Reads were mapped using STARlong to the *Drosophila melanogaster* reference genome (Ensembl assembly release dm6), and the transcriptome reference annotation release-96.

**FLAM-seq**

FLAM-seq libraries were prepared as described in Legnini et al. (extended protocol available at 10.21203/rs.2.10045/v1) using 4 µg total RNA from 3-day-old *w*1118 fly heads. Briefly, poly(A)-selected RNA was tailed using the USB poly(A) length assay kit (Thermo Fisher), cleaned up with RNAclean XP Beads (Beckmann Coulter) and reverse transcribed with SMARTScribe Reverse Transcriptase kit (Clontech). The resulting cDNA was purified with XP DNA beads (Beckmann Coulter), amplified by PCR with the Advantage 2 DNA polymerase mix (Clontech), and purified again using Ampure XP DNA Beads (Beckmann Coulter). After SMRTbell adapter addition, libraries were sequenced on 3 SMRT cells on a Sequel I PacBio sequencer. Reads were processed using the FLAMAnalysis pipeline (https://github.com/rajewsky-lab/FLAMAnalysis) with the *Drosophila melanogaster* Ensembl genome assembly and transcriptome reference annotation (release dm6).

**Comparison across LRS methods**

Calculations of transcript coverage per read were obtained by dividing the number of aligned nucleotides by the annotated transcript length. To compare gene expression estimates across long-read and short-read sequencing methods, a variance stabilizing transformation (VST) was applied using the DESeq2 function vst() on raw gene counts data from the different samples. The transformed data was used to compute a PCA using the DESeq2 function plotPCA() with standard parameters. Enrichments relative to TSS and PAS were computed by comparing the total number of reads mapping to TSS or PAS regions divided by the total number of reads assigned to the whole gene. Poly(A) signal enrichment was obtained by screening for motifs in a 20-nucleotide window of every PAS. Screening followed a hierarchical order based on known poly(A) signals and their strength, with the following rank: AATAAA, ATTAAA, AATATA, AAGAAA, AATACA, AATGAA, ACTAAA, CATAAA, GATAAA, TATAAA, TTTAAA. The positional probabilities per nucleotide were computed by counting the total number of times a given nucleotide was found in a given position per total number of nucleotides observed at a given position.

**Comparison of long read 5’ end pile-ups**

For the benchmarking of LRS putative novel TSSs, we used ONT cDNA datasets. The reads were trimmed to their most 5’ nucleotide, and peaks were called in windows of 50 nt. Only peaks with more than 30 counts per million were kept for comparison. Peaks were tested for overlaps against the Eukaryotic Promoter Database (EPD) using a window of 50 nt. Using the ChipSeeker package and a window of -150 to +150, non-overlapping 5’-pile-ups were annotated to features against the reference annotation (Ensembl assembly release dm6).

**Generation of the Drosophila Combined Isoform Assembly (CIA) database**

**Transcriptome assemblies**

For each tissue and method, all sequencing replicates were merged into a FASTQ file before assembly. Minimap2 was used to map Nanopore long reads with the “-ax splice -u f” option to the Drosophila dm6 genome indexed with the “-x 14” option. STARlong was
used to map Iso-seq and FLAM-seq data using the following parameters:

```
--outFilterMultimapScoreRange 20 --outFilterScoreMinOverLread 0.66 --outFilterMismatchNmax 1000 --winAnchorMultimapNmax 200 --seedSearchStartLmax 12 --seedPerReadNmax 100000 --seedPerWindowNmax 100000 --alignTranscriptsPerReadNmax 100000 --alignTranscriptSpPerWindowNmax 100000
```

The resulting BAM files were indexed and converted to bed12 files. FLAIR was then used to correct and collapse isoforms. During the FLAIR `correct` step, splice junction information from the respective RNA-seq datasets (short reads) was used to correct individual transcriptions. During the FLAIR `collapse` step, the Eukaryotic Promoter Database EPD was used to retain only reads with a supported TSS at their 5’ end, using “--max_ends 5” to allow for multiple 5’-3’ end identification. A minimum of three (Nanopore) or two (iso-seq/FLAM-seq) full-length reads were required for an isoform to be collapsed in the assembly. The resulting isoforms were annotated with SQANTI v1.2 to determine novel isoforms and structural categories, using an internal priming window of 50.

**Generation of a PAS database**

Assemblies were filtered for 3’ ends that likely originated from internal priming or truncation during library preparation. We used FLAM-seq and DRS data, as both of these methods allow for poly(A) tail detection, to perform poly(A) tail calling. Only reads containing a poly(A) tail were retained, and were trimmed to a single nucleotide preceding the poly(A) tail. Single nucleotide reads were clustered in 20-nt windows; clusters supported by at least two reads were included in the PAS database. The database includes only protein-coding transcripts.

**3’ end filtering and correction**

Individual assemblies from each method were corrected using the PAS database. The following filtering parameters were considered: 1) all isoforms overlapping a 3’ end in a window of 100 nt were retained, 2) 3’ ends found in the assembly more distal than the 3’ ends found in the database were retained only if they were within the reference annotation and contained an AATAAA signal. For 3’ end correction: 1) 3’ UTR bins were created using the PAS database, starting from the end of the open reading frame, between each consecutive PAS, to the most distal PAS. Isoform 3’ ends falling within the last bin of the 3’ UTR (between the two distal-most PASs) were corrected to the most distal bin, provided the isoform covered more than 10% of the last bin. Assemblies were merged first by tissue, using TAMA. Isomapping was merged if their difference was less than: 150 nt at the 3’ end, 50 nt at the 5’ end, and 10 nt at exon boundaries. After generating merged transcriptional organisms for each tissue, we combined transcriptional organisms to create the CIA assembly. All steps and pipelines used to create CIA can be found in: https://doi.org/10.5281/zenodo.7759448.

**Functional annotation of CIA transcriptome**

To generate an annotation of the CIA transcriptome at isoform-feature level, we used IsoAnnotLite version 2.7.3 with “--novel flag”, using precomputed files for *Drosophila melanogaster* and the CIA reference. Annotated transcriptome data were deposited at NCBI Gene Expression Omnibus (GEO). To explore and retrieve features from the CIA transcriptome, the R package TaiLoR is available at: https://doi.org/10.5281/zenodo.7759434.

**Generation of the human cerebral organoid CIA database**

Organoid CIA was generated using FLAIR and steps were identical to Drosophila CIA, with the following modifications. The FANTOM TSS database was used for FLAIR collapse. The organoid 3’ end database used organoid FLAM-seq data obtained from biological replicates of the RNA samples from which ONT cDNA data were generated. Short-read correction used organoid mRNA-seq data obtained from biological replicates of the RNA samples from which ONT cDNA data were generated. A minimum of three full-length reads were required for an isoform to be collapsed in the assembly. The same parameters were used for database building as for Drosophila CIA, except that clusters supported by at least one read were included in the PAS database. The assembled transcriptional organisms were assessed for novel isoforms as well as structural categories using SQANTI3v.1.2.

**Poly(A) tail length estimation**

For FLAM-seq datasets, poly(A) tail length estimation was performed using https://github.com/rajewsky-lab/FLAMAnalysis. For DRS datasets, poly(A) tail length estimations were performed using https://github.com/nanoporetech/pipeline-polya-ng. Lengths were summarized at gene level as the median poly(A) tail length per gene. At isoform level, tails were assigned to transcripts and summarized as median poly(A) tail length per transcript.

**Saturation analysis**

Saturation analysis was performed by pooling all ONT cDNA datasets from all tissues and randomly sampling different fractions from 1% to 100% from the raw read files using seqtkV1.2-r94. Then, the CIA framework was applied to each individual fraction. Results were summarized as a fraction of recovered compared to the full set.

**3’ end and 5’ end diversity calculation**

The diversity of 3’ ends per gene type was estimated with the Shannon and Simpson indexes using the R package vegan. To assess the regulatory relationship between TSS and PAS diversity, we computed the number of 3’ ends found in genes with increasing numbers of 5’ ends, and vice versa. The matrices of counts for both calculations were provided as input for both Shannon and Simpson index calculation using the function `diversity()`.
3' UTR length comparisons
Differential expression of 3' ends in heads and ovaries was computed using DEXSeq. The average length of the bins that were significantly differentially expressed was calculated and summarized per gene for each tissue.

Long-reads-based Alternative Termination Estimation and Recognition (LATER)
Quantification of 5'-3' isoforms
We counted 5'-3' isoforms using GenomicFeatures. Each ONT cDNA read was assigned to a TSS in a window of 50 nt and to a PAS in a window of 150 nt. Only the reads that mapped to both features were retained and considered full-length reads. Counts were summarized in 5'-3' isoforms, resulting in counts for each 5'-3' combination. For dominant promoter calculations, transcripts longer than 10 kb were not assessed due to lack of full-length coverage.

Calculation of TSS bias in APA-ATSS genes
A joint frequency matrix containing the reads of each 5'-3' isoform was summarized and subjected to multinomial testing with chi-squared test. We used Monte-Carlo simulation processing to obtain reliable estimates for the p-values and then corrected them using the Benjamini-Hochberg method. Only genes with at least two 5'-3' isoforms, each isoform defined by at least two full-length reads, were considered for the analysis. For Drosophila data, a gene was classified as transcriptionally biased with the p-value cutoff: adj. p-value < 0.1. For human brain organoid data, because it was supported by fewer reads, we used a more stringent p-value cutoff: adj. p-value < 0.01.

Calculation of TSS bias
Promoter dominance was estimated using two different metrics: TSS contribution and PAS contribution. TSS contribution represents the number of reads of a given 5'-3' isoform, divided by the total number of reads supporting the overall expression of that 3' end. PAS contribution represents the number of reads of a given 5'-3' isoform, divided by the total number of reads supporting the overall expression of that 5' end. A TSS was termed a “dominant promoter” if 1) the gene was classified as transcriptionally biased, 2) the TSS contribution exceeded 20%, and 3) the PAS contribution exceeded 60%. The R package LATER with a description of all processing steps can be found in: https://doi.org/10.5281/zenodo.7759430.

Quantification of differential 5'-3' isoform expression
5'-3' isoforms were quantified using the LATER counter and summarized as a counts table per pair. The table was provided to the DEXSeq framework for differential isoform usage, modeling each 5'-3' isoform as an exon feature within a gene group. To determine whether the changes in 5'-3' isoform expression originated from the TSS, the PAS, or both, differential gene expression was carried out individually for each TSS and PAS, then assigned to each 5'-3' isoform.

Long-reads-based Alternative Splicing Estimation and Recognition (LASER)
LASER quantifies the regulatory links between exons, 5' ends and 3' ends. Given that every read represents a full-length transcript, we assessed all features of each read to quantify the frequency of co-occurrence between features using multinomial testing.

Quantification of TSS-exon or 3'-exon associations
Reads were filtered to retain only full-length reads using the same parameters as in LATER. For each read, junctions were corrected using short-read sequencing and the reference annotation. Then for each read, a database was created containing all exon junctions as well as the 5' and 3' ends. Using this read to feature assignment, the total reads carrying the combination of a given 5' end with an exon-junction, or 3' end with a given exon junction were summarized.

Calculation of TSS-exon or 3'-exon biases
We created a database of exon junctions that considered only exons that are independent of 5' (alternative 1st exon) or 3' regulation (alternative last exon). Only genes containing more than one splice junction combination were retained. A joint frequency matrix containing the total number of counts per 5'-exon or 3'-exon pair was summarized and subjected to multinomial testing as in LATER. As a measure of bias strength, we summarized every residual of each tested combination using the sum of squares for each gene. To classify splicing events associated with links, we classified alternative splicing events in the CIA transcriptome using SUPPA. Using this annotation, exon junctions associated with the splicing events were extracted. Only junctions with an absolute residual change > 0.7 were considered biased. The R package LASER with a description of all processing steps can be found in: https://doi.org/10.5281/zenodo.7759428.

ChIP-seq data analysis
ChIP-seq data obtained from Drosophila head tissue (modENCODE) was analyzed. Fastq files were mapped and processed with snakePipes using DNA-mapping and the ChIP-seq workflow, adding flags for “–singleEnd and –fragmentLength 50”. Bigwig signal tracks were generated by computing the log2 fold change of each ChIP compared to the respective input. Heatmaps, gene profiles and clustering were generated using deepTools.

Analysis of transcription factor enrichment at TSSs
TSSs were generated from the CIA reference transcriptome using a 50 nt window. Enrichment of factors at TSSs was estimated using the ReMap2022 databases for Drosophila and Human and the package ReMapEnrich with, as background, ATSS-APA genes without a dominant promoter. Enrichment was determined with the cutoff: p-value<0.01.
Analysis of single-cell RNA-seq data using the CIA 3' end database

Raw data from the single-cell Drosophila brain transcriptome atlas were mapped using CellRanger. To generate the matrix of counts, the CIA 3' end database was provided as an input to the Sierra3 function CountPeaks(). Per isoform-cell counts were annotated to cell types and clustering information with PeakSeuratFromTransfer() using metadata from Davie et al.3 End expression was then summarized per cell type using the Seurat::AverageExpression() function, and normalized using the Seurat::NormalizeData() function with LogNormalize. 3' ends per cell type were considered expressed if they were represented by at least 0.1 normalized counts.

Conservation of 5' UTRs and 3' UTRs

PhasCons scores were retrieved using the GenomicScores R package for Drosophila using reference phastCons tree model for the 27 species.

Co-evolution analysis

We determined gene co-evolution maps at the single nucleotide level using pairwise mutational information between positions derived from 27 species alignment tracks from UCSC, with Drosophila melanogaster (dm6) as the reference sequence. For the genes stai and Act5C, we extracted multiple sequence alignments from -1.5 kb to the 3' end of the gene. The retrieved alignments were filtered using the refineMSA function from the ProDy package, keeping sequences with 60% gaps (parameter: rowocc = 0.4) and an identity level of 98% (parameter seqid=0.98), since the alignments spanned the entire gene, including introns. We used mutual information to estimate the probability that a given nucleotide change would be accompanied by another nucleotide change. We normalized the mutual information using the average product correction method (APC) and implemented in the ProDy python package. To perform a global analysis of co-evolution, we selected the top 50 dominant promoter and the bottom 50 (by p-value from the LATER analysis). We computed co-evolution using three regions of interest for each gene to reduce computational time: 1) TSS1 (-1kb), 2) TSS2 (-1kb), 3) the entire 3' UTR sequence. To extract the mutual information between each TSS region and the 3' UTR from the co-evolution matrix, we identified the local maxima of normalized mutual information using the function gsignal::findpeaks(x, MinPeakDistance = 2, MinPeakWidth = 2, MinPeakHeight = 0.2) of the R package gsignal. For every gene, we computed the sum of local maxima of the overlapping regions promoter/3'UTR. We classified genes as “co-evolving” when the sum of local maxima was in the top 50th percentile of the distribution of the sum of local maxima in the dataset. The code for all steps from extraction to processing and output is available at https://doi.org/10.5281/zenodo.7759440.

Identification of differential poly(A) site usage

We identified differential poly(A) site usage, using the APA target caller with the parameters “min_distance = 100 padj < 0.05”.

Motif enrichment in dominant-promoter-associated 3' UTRs

To predict potentially relevant microRNA binding sites (i.e. with a higher likelihood to exert a functional impact on target mRNAs) in dominant-promoter associated, distal 3' UTRs, we used a subset of 65 microRNAs that were 1. highly conserved (node of origin: Diptera) and 2. well expressed in fly heads (at least 1000 cpm) from MirGeneDB v.2.1, collapsed them into 52 unique 7mer (2-8) seed sequences and computed the number of occurrences of their reverse complementary sequence in either proximal or distal 3' UTR isoforms for a set of 173 dominant-promoter genes for which the distal 3' UTR was uniquely associated with a dominant promoter. RBP enrichment in dominant-promoter-associated distal 3' UTRs was performed on the distal 3' UTR segments using the BSgenome.Dmelanogaster.UCSC.dm6 reference genome package in R. The FASTA files were submitted to the MEME suite server and the AME program was used to calculate enrichment. For the comparisons, proximal 3' UTR segments were used as control sequences. Motif scanning was performed using FIMO with a cut-off p-value < 0.0001, using the motif matrices from for RBP enrichment. For microRNA enrichment analysis, motif scanning used the miRbase v22 Single Species microRNA database for Drosophila melanogaster. To further assess the regulatory potential of these 6 microRNAs, we first confirmed that they are expressed in fly heads in MirGeneDB v.2.1, and of the only two resulting (poorly) expressed in fly heads, dme-mir-2279-5p and dme-mir-9388-5p, we used TargetScan Fly v7.2 to compile a list of predicted binding sites transcriptome-wide for mir-2279-5p. A Gene Ontology analysis was performed on the resulting gene list (mRNAs not expressed in heads were excluded) using DAVID (v2022q4). We defined microRNA targets as genes with a cumulative weighted context score less than -1. Head-expressed genes were used as the background. GO terms with a p-value less than 0.05 (after Bonferroni false discovery rate (FDR) correction) were considered significant.

3'-seq analysis

Reads were processed with fastp to remove poly(A) stretches and then mapped to the dm6 genome using STAR v2.6.1b with modified parameters (“--sjdbOverhang 74 --limitBAMsortRAM 60000000000 --alignIntronMax 1”). In order to eliminate the signal that may come from internal priming, any poly(A) sites overlapping with a strand-specific blacklist region that contained genomic positions with more than 70% As in a 10-bp upstream window were discarded. Regions with high A density within 250 bp of annotated transcription end sites were not included in the blacklist. The remaining single base pair poly(A) sites from all samples with a minimum coverage of 5 reads per sample were grouped, with sites within 15 bp merged into a single poly(A) cluster.
Random forest classification of 3’ ends
Using 3’-seq clusters, we extracted features from 3’ ends identified by FLAM-seq in human organoids. These features included: poly(A) signals at 20 nucleotides upstream from the identified 3’ end, the nucleotide content and annotated feature (e.g. 3’ UTR, 5’ UTR) of the 3’ end. We used these features to train a Random Forest model in R using the randomForest package. We created a training set based on FLAM-seq 3’ end clusters as our TRUE set and non-overlapping 3’ ends as the FALSE set. The model was trained using 1000 trees with 12 random variables set at each split (randomForest(ntree=1000, mtry=12)). The TRUE clusters obtained from classification were then used as a poly(A) database to correct human organoid assemblies. Pretrained models are available at: https://doi.org/10.5281/zenodo.7438383.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical parameters and tests are reported in the respective figure legends; software used is described in the STAR Methods section and in the key resources table.

ADDITIONAL RESOURCES

A Drosophila mRNA isoform atlas, depicting all CIA transcript isoforms identified and representing their differential expression in several tissues and developmental stages, is publicly available. https://hilgerslab.shinyapps.io/ciaTranscriptome.
Figure S1. An accurate, comprehensive, full-length Drosophila transcriptome, related to Figure 1
(A) Combined isoform assembly (CIA) workflow, and schematic of 3’ end correction and filtering. ONT DRS (in heads: ONT DRS and FLAM-seq) data were used to build a database of confident 3’ ends. The CIA assembly was performed with ONT cDNA and ONT DRS (in heads: ONT cDNA, Iso-seq, ONT DRS, and FLAM-seq) data using FLAIR and the Eukaryotic Promoter Database (EPD-new). Note that due to the low number of Iso-seq reads compared with ONT cDNA reads, Iso-seq reads contribute to CIA to a much lower extent. Since this assembly contains 3’ end artifacts, we filtered out any transcripts with 3’ ends not represented in the DRS/FLAM 3’ end database. Assembled transcript models were corrected with DRS/FLAM 3’ ends.
(B) Number of corrected transcripts per tissue (left) and average length of correction (right). Data from the two embryo datasets (14–16 h AEL and 18–20 h AEL) were pooled.
(C) Read lengths in each tissue with each LRS method. BluePippin size selection (red graphs, below) considerably increased full-length transcript coverage.
(D) Full-length transcript coverage per read for nanopore cDNA and PacBio Iso-seq in heads, before (left two graphs) and after (right two graphs) size selection. For each read, the fraction of the target transcript covered is shown; reads were grouped by the length of the target transcript.
(E) Principal-component analysis plot of gene expression across the samples (3 biological replicates each) generated using nanopore cDNA with and without size selection, nanopore direct RNA sequencing (DRS), and Illumina short-read mRNA-seq from three tissues. Data from the two embryo datasets (14–16 h AEL and 18–20 h AEL) were pooled. Note that LRS methods without size selection (ONT cDNA and DRS, blue and gray) cluster further from mRNA-seq expression estimates (black) than ONT cDNA with size selection (red).
(F) Cumulative plot representing the fraction of long-read 5’ ends that overlap with a TSS described in the Eukaryotic Promoter Database in a window of 50 nt, as a function of long-read 5’ end counts per million. A 5’ pile-up was defined as a cluster of >30 counts per million per window (dashed line).
(G) Pie chart representing the number and proportion of 5’ pile-ups that overlap (purple) with a TSS described in the Eukaryotic Promoter Database. Non-overlapping pile-ups (gray in the left pie chart) were assessed for the gene region of occurrence (right) as annotated in ENSEMBL.
(H) Cumulative enrichment plots of RNA Pol II ChIP-seq signal, H3K4me3 ChIP signal, and ATAC-seq signal detected at 5’ pile-ups (+2 kb) that overlapped (purple) or not (gray) with TSSs annotated in the Eukaryotic Promoter Database. ChIP-seq and ATAC-seq data from Drosophila heads are from modENCODE.
(I) Venn diagram describing the overlap of mRNA 3’ ends of LRS reads after filtering (CIA) with filtered-out 3’ ends (discarded) and Ensembl-annotated mRNA 3’ ends. 3’ ends detected by FLAM-seq or DRS represent CIA 3’ ends (purple); not-detected 3’ ends (gray) were discarded.
(J) Nucleotide composition profiles (spanning 200 nt, top) and sequence logos (spanning 40 nt, bottom) of LRS reads at the cleavage site for each denoted category of 3’ ends from our processing pipeline. Noisy, A-rich distributions are indicative of internal priming. The left and middle panel nucleotide distribution profiles are also shown in Figure 1 and reproduced here for side-by-side comparison with the Ensembl-only category.
(K) In each tissue, proportion of 3’ ends at which the indicated poly(A) signals were detected for each category (CIA or discarded [Dc]). Data from the two embryo datasets (14–16 h AEL and 18–20 h AEL) were pooled.
(L) In CIA transcripts, proportion of 3’ ends carrying a novel (purple) or a previously annotated (gray) 3’ end. CIA transcripts were categorized by poly(A) signal. Replicates per tissue: ONT cDNA: heads, n = 6; embryos 14–16 h, n = 3; embryos 18–20 h, n = 3; ovaries n = 3. FLAM-seq and Iso-seq: heads, n = 3; DRS: heads, n = 1, embryos 14–16 h, n = 3; ovaries, n = 3. Illumina TrueSeq mRNA-seq: each tissue, n = 2.
**Figure S2. Transcription start sites drive tissue-specific 3’ end expression, related to Figures 1 and 2**

(A) Number and proportion of novel (red) and previously annotated (gray) isoforms in the full CIA transcriptome assembly dataset across tissues.

(B) Proportion and number of newly identified features (red, novel) that contribute to newly identified isoforms. Most previously unannotated isoforms arise from the differential use of known (annotated) splice junctions with the canonical splice signal and from alternative 3’ end site usage (from 6,483 novel 3’ ends).

(C) Proportion of structural and quality annotation of novel transcript isoforms (SQANTI) categories found in the CIA transcriptome assembly as a function of transcript length. The number of transcripts in each category is indicated in parentheses. Transcript lengths were binned by kb.

(D) Type and percentage of splicing events found in CIA isoforms. Newly identified isoforms (red) showed no splice class bias, compared with previously annotated isoforms. A3, alternative 3’ splice site; A5, alternative 5’ splice site; AF, alternative first exon; AL, alternative last exon; MX, mutually exclusive exon; RI, retained intron; CE, cassette exon.

(F and G) Overlap of transcript expression across tissues considering all reads from all LRS approaches (F), or sub-sampling 4.3 million ONT cDNA reads (G). Isoforms from 14- to 16-h and 18- to 20-h AEL embryos were pooled in (G). 4.3 million reads were used for subsampling because it represents the smallest read number obtained for an individual tissue (ovaries). Isoforms were considered distinct if they differed by more than 10 nt at exon boundaries, 50 nt at the 5’ end, or 150 nt at the 3’ end.

(H) Contribution of coding sequence and 3’ UTR length to transcript length for long (>5 kb) transcripts across tissues. **p < 2.2e–16 (ANOVA). Data from the two embryo datasets (14–16 h AEL and 18–20 h AEL) were pooled.

(I) Distribution of genes by mean poly(A) tail length for each tissue. Data from the two embryo datasets (14–16 h AEL and 18–22 h AEL) were pooled.

(J) Increase in poly(A) tail length as a function of transcript length. Poly(A) tails of mitochondrial mRNAs were included for comparison.

(K) Proportion of genes that undergo ATSS in each PAS category. **p < 0.001 (two-tailed Fisher’s exact test).

(L and M) 3’ end diversity as a function of the number of TSSs per gene (L) and TSS diversity as a function of the number of 3’ ends per gene (M). The Shannon index is a measure of diversity that considers the relative abundance of different species (individual 5’-3’ isoforms) in a population (the sum of all 5’-3’ isoforms). To account for possible coverage biases, the analysis of the whole dataset (black line) was also performed in randomly sampled fractions of the pooled nanopore cDNA data (in grayscale).

(N) 5’-3’ isoforms across the Ensembl and CIA reference transcriptomes. 5’-3’ isoforms were considered distinct if they differed by more than 50 nt at the 5’ end or 150 nt at the 3’ end. Comparison after gene expression filtering (>2 transcripts per million (TPM)).

(O) Saturation analysis of 5’-3’ isoforms of ATSS-APA genes in the tissue pool, grouped by their expression in transcripts per million (TPM). Reads were randomly sampled in the indicated fractions and the assembly pipeline including 3’ end correction was performed in each fraction.

(P) Differential expression of 3’ ends in heads compared with ovaries, plotted as a function of the differential expression of the 5’ end associated with each 3’ end. Red represents 5’-3’ isoforms with a significant 5’-3’ link, i.e., a significant expression change for both the 3’ end and its associated 5’ end (|log2(FC)| > 0.5 and adj. p value < 0.05, Wald test, 3 replicates per tissue).

(Q) For all tissue-specific TSSs, or 3’ ends, number of associated 3’ ends or TSSs, respectively, that are also tissue-specific.

(R) Number of genes with two or more links in which the expression bias is opposite between the two tissues (bidirectional). Significant links were determined as: |log2(FC) (5’-3’ isoform expression)| > 1.5 and adj. p value < 0.01 (Wald test, 3 replicates per tissue).

(S) RT-qPCR quantification of the indicated transcript regions in flies in which dCas9-VPR was recruited to nervous-system TSSs for activation (purple). In control flies, dCas9-VPR was co-expressed with a non-targeting sgRNA (gray). Shown are five further genes for which TSS activation was successful in heads (in addition to Fatp1 shown in Figure 2): Malvolio (Mv), tout-velu (ttv), tramtrack (ttk), wunen (wun), and charlatan (chn). RNA levels were normalized to RpL32 mRNA, and levels in control flies were set to the value 1. Error bars represent mean ± SD of four biological replicates (five heads per replicate) for each genotype. *p < 0.05, **p < 0.01, ***p < 0.001 (one-tailed Student’s t test).
Figure S3. Dominant promoters drive PAS choices, related to Figure 3

(A) Cumulative plot representing the fraction of ATSS-APA genes as a function of the fraction of cells co-expressing more than one 3’ end across cell types with an expression of more than 0.1 normalized counts.

(B) Proportion of ATSS-APA genes in which two or more isoforms were found to be expressed in the indicated percentage of cells in single-cell RNA-seq data from the Drosophila Brain Atlas. For most genes (above the 0.5 proportion), most (over 50%) cells express two or more 3’ ends.

(C) t-distributed stochastic neighbor embedding (t-SNE) maps representing 3’ end expression in Drosophila brain cell types for the two representative genes Multiplexin (Mp) and stathmin (stai). Cells are colored according to their expression of either only the proximal (red), only the distal (blue), or both 3’ end isoforms (purple). Shown below are the gene model and 5’-3’ reads representing the expression of the detected 5’-3’ isoforms. Some introns (dashed lines in the gene model) are not drawn to scale.

(D) Schematic representation of how TSS and 3’ end contributions to 5’-3’ isoform expression were calculated. Full-length 5’-3’ reads were quantified and assigned to 5’-3’ isoforms. For a given 3’ end, the contribution of each 5’-3’ isoform to the expression of the 3’ end was calculated (pink), as well as for a given TSS, the contribution of each 5’-3’ isoform to the expression of the 5’ end (orange). A TSS is termed a dominant promoter for a 3’ end if the respective 5’-3’ isoform contribution has a contribution to 3’ end expression significantly higher (p < 0.1, chi-squared test with Monte Carlo simulation and Benjamini-Hochberg correction, also see E) than that of all other 5’-3’ isoforms for the same 3’ end.

(E) TSS bias in ATSS-APA genes assessed using multinomial testing in Drosophila heads. The observed vs. expected counts of 5’-3’ isoforms were used for multinomial testing (chi-squared test with Monte Carlo simulation and Benjamini-Hochberg correction, n = 3). Genes are represented as dots, ranked by p value and color-coded according to bias score (promoter dominance score: absolute value of residuals). Highest-ranked genes (220 genes in the brain) represent near-exclusive 5’-3’ combinations, as exemplified by stai.

(F) Promoter dominance and absence thereof (no TSS bias) shown on representative ATSS-APA genes with two TSSs and two PASs. The proportional contribution of the first TSS (red) and the second TSS (blue) to the expression of the proximal and the distal 3’ end of the same gene are indicated. Lines crossing signify TSS contributions that differ significantly between the PASs.

(G) Pie chart representing the percentage of dominant promoters that constitute the top expressed TSS of the gene in heads.

(H) Scatterplot showing the expression ratio between isoforms expressing the distal and proximal PAS, respectively, measured by long-read sequencing (ONT cDNA), in function of ratios measured by mRNA-seq (Illumina short reads). The ratios were calculated by estimating the ratio of normalized TPM (transcripts per million) assigned to proximal and distal 3’ ends in APA genes. Each dot represents a gene. The correlation coefficient (two-tailed Pearson correlation) is indicated for genes with a dominant promoter (promoter dominance) and TSS-unbiased genes (no significant TSS bias).

(I) Proportion of 5’-3’ isoforms by category, expressing the indicated types of coding sequence, as a function of coding sequence length. Coding sequences are categorized by length within the gene context and represent either the longest, shortest, or an intermediate CDS isoform. Coding sequences of a gene were considered of identical length (all same) if none differed by more than 200 nt. 5’-3’ isoforms are grouped into 5’-3’ isoforms with a dominant promoter (dominant) and 5’-3’ isoforms with no dominant promoter (not dominant).

(J and K) Saturation analysis of splice junctions (J) and splice combinations (K) in CIA transcripts, grouped by their expression in number of reads. Reads were randomly sampled in the indicated fractions and a junctions (J) or combinations (K) database was built for each fraction. Splice junctions are exon-exon junctions. Splice combinations are unique assemblies of consecutive exons for each gene. Exons containing, or upstream of, a TSS (first exon), or containing or downstream of a PAS (last exon), were excluded from the analysis.

(L) Long-reads-based alternative splicing estimation and recognition (LASER) framework to identify TSS biases in alternatively spliced (AS) genes (left), and splicing biases in alternatively polyadenylated (APA) genes (right). TSS-exon bias: for each splice junction of each ATSS-AS gene, the observed vs. expected frequencies of TSS-junction combinations were calculated to identify TSSs disproportionately associated with the junction (TSS-exon links). Exon-PAS bias: for each PAS of each AS-APA gene, the observed vs. expected frequencies of splice junction-PAS combination were calculated to identify splice junctions disproportionately associated with the PAS (exon-PAS links). Significant TSS-exon and exon-PAS links were identified by multinomial testing (p < 0.1, chi-squared test with Monte Carlo simulation and Benjamini-Hochberg correction) and assigned a linkage score (sum of squares of residuals). Splice junctions are exon-exon junctions.

(M) Genes in which alternative polyadenylation is linked to alternative splicing (exon-PAS links) or transcription start sites (TSS-PAS link: promoter dominance), or both. Intersections between the gene sets are depicted as connecting lines. The number of genes in each exclusive group is indicated. Only 81 genes with an exon-PAS link were identified outside of the ATSS-APA gene group, and only 21 within the ATSS-APA gene group that were not associated with a dominant promoter (TSS-PAS link).

(N) Type and number of alternative splicing events found in mRNA isoforms transcribed from dominant promoters: alternative 3’ splice site; alternative 5’ splice site; intron retention; mutually exclusive exon; cassette exon.
A. Upon use of dominant-promoter-associated 3’ ends:
Total 3’ UTR sequence gained: 817,365 nt
Total 3’ UTR sequence lost: 70,304 nt

B. Gained and lost 3’ UTR 7mers

C. RBP binding motifs gained or lost in dominant-promoter 3’ UTRs

D. RBP binding motifs enriched in dominant-promoter 3’ UTRs

E. Predicted microRNA sites enriched in dominant-promoter 3’ UTRs

F. Gene Ontology terms for miR-2279-3p predicted targets in heads (n=1023)

(legend on next page)
Figure S4. Functional impact of promoter dominance on transcriptome diversity and tissue identity, related to Figure 4

(A) 3’ UTR sequence length gained or lost by the predicted shift in PAS selection as a result of promoter dominance, for 173 dominant-promoter genes, in fly heads. “Gained” and “lost” refers to dominant-promoter-3’ UTRs associated with the distal and proximal PAS, respectively.

(B) Number of potential binding sites (7-mers) gained (blue) or lost (red) by the predicted shift in PAS selection as a result of promoter dominance, for a set of 65 highly conserved and highly expressed microRNAs (collapsed into 52 seed sequences), for 173 dominant-promoter genes, in fly heads. The total number of gained and lost sites and the 3’ UTR length difference between proximal and distal isoforms are indicated at the bottom.

(C) Number of binding motifs for the indicated RNA-binding proteins (RBPs) gained or lost by the predicted shift in PAS selection as a result of promoter dominance, for 173 dominant-promoter genes, in fly heads.

(D) RBP binding motifs enriched in dominant-promoter 3’ UTRs associated with the distal PAS, compared with 3’ UTRs associated with the proximal PAS.

(E) Predicted microRNA binding sites enriched in dominant-promoter 3’ UTRs associated with the distal PAS, compared with 3’ UTRs associated with the proximal PAS. MicroRNAs detected in Drosophila heads (MirGeneDB v2.1) are shown in red.

(F) Enriched gene ontology terms in 1,023 mRNAs expressed in heads that are predicted targets for dme-miR-2279-5p.
Figure S5. TSSs exert promoter dominance through specific chromatin signatures, related to Figure 5

(A) Heatmaps and cumulative enrichment plots of ChIP-seq signal at TSS ± 2 kb for RNA Pol II, the histone marks H3K4me3, H3K9Ac (typical for active promoters and TSSs of expressed genes), H3K27Ac, and H3K4me2 (active enhancer and TSS marks), the histone variant H2A.Z and the histone mark H3K18Ac genome-wide. Genes are grouped by CIA categories. ChIP-seq data from Drosophila heads are from modENCODE.61

(B–D) ChIP-seq peak enrichment analysis at the TSS and PAS of dominant promoter isoforms. Represented are factors significantly enriched (adj. p value < 0.05) at either the TSS ± 150 nt (B), the associated PAS ± 150 nt (C), and both (D), ranked by the ratio of total peaks that map to the TSS (B and D) or PAS (C).

(E) Enrichment (blue) and depletion (red) of distal 3' UTR RNA expression in the indicated mutants compared with control embryos. mRNA-seq heatmaps and profile plots display 0.5 kb upstream of the proximal poly(A) site (prox), and the distal 3' UTR downstream (distal, scaled region). For CBP and Psc, results from two
independent mutant alleles are shown. RNA was obtained from hand-sorted embryos at 16–18 h AEL in three biological replicates for each genotype (except CBP mutants: 14–16 h AEL, four replicates). Genes are grouped into three clusters by k-means clustering using both CBP ChIP-seq signal at proximal PASs and mRNA-seq signal in nej3 and nejEP1179 mutants. Heatmaps for CBP mutants, also shown in Figure 5, are reproduced here with a different color scale for side-by-side comparison with the other mutants. CBP, Dfd, E(z), and Psc were found enriched at the TSS of dominant promoters and their associated 3’ end; Spps was found enriched at the TSS of dominant promoters only.
Figure S6. Dominant promoters drive PAS selection in human brain organoids, related to Figure 6

(A) Venn diagram describing the overlap of 5'–3' isoforms in the Ensembl and CIA reference transcriptomes for human brain organoids (three biological replicates). 5'–3' isoforms were considered distinct if they differed by more than 50 nt at the 5' end or 150 nt at the 3' end. Comparison after gene expression filtering. Organoid CIA identified around 22,000 5'–3' isoforms.

(B) TSS bias in ATSS-APA genes assessed using multinomial testing in human brain organoids. The observed vs. expected counts of 5'–3' isoforms were used for multinomial testing (chi-squared test with Monte Carlo simulation and Benjamini-Hochberg correction). Genes are represented as dots, ranked by p value and color-coded according to bias score (absolute value of residuals).

(C) ChIP-seq peak enrichment analysis at the TSS and PAS of dominant promoter isoforms. Represented are factors significantly enriched (adj. p value < 0.1) at the TSS ± 150 nt (left), and at both the TSS ± 150 and its associated PAS ± 150 nt (right), ranked by the ratio of total peaks that map to the dominant promoter. Transcription factors and co-activators reported to influence 3' end site choice are in bold; homologs of p300/CREBBP, SMC, and TAF11 are in red. Data are from the ReMap database.