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Numerous recursive sites contribute to accuracy of splicing in long introns in flies

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Abstract

Recursive splicing, a process by which a single intron is removed from pre-mRNA transcripts in multiple distinct segments, has been observed in a small subset of Drosophila melanogaster introns. However, detection of recursive splicing requires observation of splicing intermediates that are inherently unstable, making it difficult to study. Here we developed new computational approaches to identify recursively spliced introns and applied them, in combination with existing methods, to nascent RNA sequencing data from Drosophila S2 cells. These approaches identified hundreds of novel sites of recursive splicing, expanding the catalog of recursively spliced fly introns by 4-fold. A subset of recursive sites were validated by RT-PCR and sequencing. Recursive sites occur in most very long (>40 kb) fly introns, including many genes involved in morphogenesis and development, and tend to occur near the midpoints of introns. Suggesting a possible function for recursive splicing, we observe that fly introns with recursive sites are spliced more accurately than comparably sized non-recursive introns.

Author summary

The splicing of RNA transcripts is an essential step in the production of mature mRNA molecules, involving removal of intron sequences and joining of flanking exon sequences. Introns are usually removed as a single unit in a two-step catalytic reaction. However, a small subset of introns in flies are removed via splicing of multiple distinct consecutive segments in a process known as recursive splicing. This pathway was thought to be quite rare since intermediates of recursive splicing are seldom detected. In this study, we developed three new computational approaches to identify sequence reads, read pairs and patterns of read accumulation indicative of recursive splicing in Drosophila melanogaster cells using data from sequencing of nascent RNA captured within minutes after transcription. We used these methods to identify hundreds of previously unknown sites of recursive splicing.
recursive splicing, occurring commonly in fly introns longer than 40 kb and often in genes involved in morphogenesis and development. We observed that recursive splicing is associated with increased splicing accuracy of long introns, which are otherwise often spliced inaccurately, potentially explaining its widespread occurrence in long fly introns.

Introduction

RNA splicing is a crucial step in the mRNA lifecycle, during which pre-mRNA transcripts are processed into mature transcripts by the excision of intronic sequences. Introns are normally excised as a single lariat unit. However, some introns in the Drosophila melanogaster genome are known to undergo recursive splicing, in which two or more adjacent sections of an intron are excised in separate splicing reactions, each producing a distinct lariat [1,2]. Recursively spliced segments are bounded at one or both ends by recursive sites, which consist of juxtaposed 3’ and 5’ splice site motifs around a central AG/GT motif (with “/” indicating the splice junction) [1,3]. This mechanism appears to be restricted to very long Drosophila introns [3,4]. However, because recursive splicing yields an exon ligation product identical to that which would have been produced from excision of the intron in one step, the genome-wide prevalence and function of recursive splicing have been difficult to ascertain [3,4].

Recursive splicing was initially observed in the splicing of a 73 kb intron in the Drosophila Ultrabithorax (Ubx) gene, where the intron is removed in four steps through intermediate splicing of the 5’ splice site to two microexons and one recursive site before pairing with the proper 3’ splice site [1]. Bioinformatic searches for recursive sites predicted a couple hundred possible recursive sites in Drosophila, predominantly in introns larger than 10 kb [3], but sites in only four introns, all from developmentally important genes (Ubx, kuzbanian (kuz), outspread (osp), and frizzled (fe)), could be experimentally validated [1–3]. Biochemical characterization showed that recursive splicing is the predominant processing pathway for splicing of these introns, which are generally constitutively spliced [1–4]. More recently, an analysis by Duff and coworkers of all ~10 billion RNA-seq reads generated by the Drosophila ModENCODE project identified 130 recursively spliced introns in flies [4]. Using this larger catalog of recursive sites, they confirmed that recursive splicing is a conserved mechanism to excise constitutive introns, requires canonical splicing machinery, and only occurs in the longest 3% of Drosophila introns [4]. Similar analyses of mammalian RNA-seq datasets have resulted in the identification of just a handful of recursively spliced introns, mostly in genes involved in brain development, despite the greater abundance of long introns in vertebrate genomes [5].

The scarcity of validated examples suggests that recursive splicing is quite rare, even in Drosophila. However, the transient nature of recursive splicing intermediates makes it difficult to detect evidence for recursive splicing using standard RNA-seq data. Support for recursive splicing has come from RNA-seq reads that span a junction between a known splice site and a putative recursive splice site internal to an intron, or from observation of a sawtooth pattern of reads resulting from the splicing out of recursive segments [4,5]. Previous studies using polyA-selected RNA-seq data— which derive predominantly from mature transcripts— had limited ability to detect such evidence. However, nascent RNA sequencing, which profiles pre-mRNA transcripts shortly after they are transcribed, should enable much more efficient capture of reads from intermediates of splicing, including recursive splicing. Using such data should allow for more unbiased and systematic discovery of recursive splicing.

To globally detect transient splicing intermediates indicative of recursive splicing, we applied novel computational approaches to high-throughput sequencing data from short time
period metabolic labeling of RNA. This approach detected about four times as much recursive splicing as had been previously observed. This expanded catalog of sites and associated analyses suggests a function for recursive splicing in improving splicing accuracy.

Results

Pre-mRNA splicing can initiate immediately after transcription of an intron is completed, and can occur in as short a time as one or a few seconds [6–9]. Since recursive splicing involves the splicing of intermediate intronic segments, it may begin soon after the transcription of the first intronic recursive site. Thus, to have the greatest chance of capturing recursive splicing intermediates, it is essential to capture nascent transcripts as soon as possible after transcription, before introns have been fully spliced. Here, we used nascent RNA sequencing data from our recent study, which used incorporation of a metabolic label to isolate RNA at short time points after transcription [9]. The experimental approach to collect these data involved 5, 10, or 20 min labeling with 4-thiouridine (4sU) in *Drosophila* S2 cells and 4sU biotinylation to selectively isolate nascent RNA, followed by RNA sequencing with paired-end 51 nt reads [9]. These data were complemented by steady state RNA-seq data representing predominantly mature mRNA (Methods). The progressive labeling strategy used for these data results in isolation of transcripts that initiated during the labeling period, in addition to transcripts that were elongated during this period but initiated prior to the addition of the label [9]. While this likely does not significantly bias the distribution of fragment lengths sequenced, there is an overall 5' to 3' bias of reads across the entire transcript.

We hypothesized that this high-coverage nascent RNA data would more readily identify recursive sites and better characterize the prevalence of recursive splicing. For this purpose, we used a computational pipeline to detect three key signatures of recursive splice sites (Fig 1).

First, we used a custom python script to search for splice junction reads derived from putative recursive sites (RatchetJunctions), as previously described (Methods; Fig 1A) [4,5]. Ratchet junction reads contain a segment adjacent to an annotated 5' or 3' splice site juxtaposed to a segment adjacent to an unannotated intronic recursive site, providing direct evidence for the presence of a recursive splicing event.

Second, we developed a new computational tool, RatchetPair, to identify read pairs that map to distant genomic sites in a manner such that presence of intervening recursive splicing can be inferred from the size distribution of inserts in the sequenced library (Methods; Fig 1A). Unlike ratchet junction reads, recursive junction spanning read pairs do not pinpoint a specific recursive site. Instead, a recursive site is inferred based on the empirical distribution of fragment lengths and genomic sequence information. To do so, we adapted the GEM algorithm [10], originally designed to infer protein binding sites from ChIP-seq data, to assign a probability that each read-pair was indicative of a recursive site in a given region (Methods). This modified GEM algorithm was run with all read pairs and splice junction reads pooled together to derive the empirical distribution of fragment lengths.

Third, we developed the first automated software, RatchetScan, for inference of recursive sites from sawtooth patterns in read density (Fig 1B). This type of pattern is an expected product of co-transcriptional recursive splicing and has been associated with many recursive introns [4,5,11]. Briefly, assuming that RNA is spliced shortly after transcription elongation past the recursive splice site or 3' splice site, the splicing of recursive segments during transcription of subsequent sequences will result in a sawtooth distribution of reads across the intron with recursive sites commonly located near the right-hand base of each "tooth". It is important to note that these approaches do not differentiate between unproductive splicing
(followed by degradation of the intron-containing transcript) and productive splicing of the full intron.

RachetScan predicts the locations of recursive sites in three distinct steps. First, RNA-seq data was processed to summarize read density in each sub-intronic region (S1A Fig). We then developed a Markov Chain Monte Carlo- (MCMC-) based inference algorithm to detect presence of sawtooth patterns in introns. This algorithm is suitable for efficient exploration of complex intronic read patterns encountered when considering a variable number of possible recursive splice sites in each intron. We considered all nucleotides as potential recursive splice sites, rather than only focus on sites at the center of strong juxtaposed recursive motifs.
allowing us to independently use sequence information to assess the false-positive rate of our method. Our RachetScan algorithm is initiated with a randomly chosen state, consisting of a set of proposed recursive sites in the intron (S1B Fig). In each round, a new state is proposed by perturbing the current state, with three classes of perturbations: (1) a new recursive site is added; (2) a recursive site is removed; or (3) a recursive site location is locally shifted, each with defined probabilities. Using a scoring function and transition rules (detailed in Methods), the algorithm decides to either accept the new proposed state or maintain the current state. This procedure was iterated over $10^7$ rounds and the current state was sampled every 50 rounds, where the number of samples recorded in each state is proportional to the probability that the intron is best fit by the model corresponding to that state. Finally, recursive sites are predicted based on the output of the inference algorithm and sequence information (S1C Fig; S2 Fig). This approach does not infer the order of splicing of recursive segments (but see below).

Combining these three approaches and using reads pooled across all replicates and labeling periods, our analysis detected 539 candidate recursive sites in 379 fly introns (S1 Table). From this set, we curated a set of 243 “high confidence” recursive sites in 157 introns (identified by at least 2 methods with greater than 5 supporting junctions or read pairs or a sawtooth FDR of 5% and visual inspection of read densities), and a “medium confidence” set of 296 sites (identified by at least 1 method with greater than 5 supporting junctions or read pairs or a sawtooth FDR of 20%; Fig 2A; Methods). Approximately 60% of our high-confidence sites (144 sites) were identified using all three approaches. Overall, 98 introns contained multiple recursive sites, with up to seven high-confidence sites observed in a single intron. For instance, intron 1 of the tenascin major (Ten-m) gene contains five recursive sites, two of which were previously unknown (Fig 1C). Of the recursive sites previously reported by Duff and colleagues, 124 occurred in genes expressed in S2 cells. Our approach detected 119 (96%) of these known sites, as well as 126 novel high confidence sites and 294 novel medium confidence sites (Fig 2A), thus increasing the number of recursive sites defined in this cell type by ~4-fold (S2C Fig). For three recursive segments, we were also able to detect reads that spanned the intronic lariat resulting from the second step of splicing (S2 Table; Methods). For 13 sites in 3 recursively spliced introns, we performed RT-PCR validation experiments using primers flanking recursive segments, followed by sequencing (Methods). These experiments validated 8 previously identified sites and 5 novel recursive sites in nascent RNA from Drosophila S2 cells, including 3 sites in an ~55 kb intron of Tet that was not previously known to be recursively spliced (S3A Fig, S3D Fig; S3 Table). Both the high confidence and the medium confidence candidate recursive sites exhibited a strong juxtaposed 3'/5' splice site motif (S4 Fig). The greater numbers of sites detected by our approach (2–4 times more sites in this cell type), using less than 1/20 th as many reads as used by Duff and colleagues, affirms the potential of nascent RNA analysis for identification of recursive splice sites.

Using this updated catalog of recursive sites, we observed that many very long introns (> 40 kb in length) have recursive sites, with 63% of such introns containing at least one high-confidence recursive site, and an additional 7% containing medium-confidence site(s) (Fig 2B). This observation suggests that recursive splicing is the prevalent mechanism by which very large fly introns are excised. We assessed the sensitivity of our detection pipeline by running it on subsamples of reads ranging from 0.1% to 100% of the total reads (Fig 2C). The shape of the resulting curve tapered off at higher coverage levels but never plateaued: new recursive sites were still being detected as read depth increased from 50% to 100% of sequenced reads and therefore would likely increase further at higher read depths. A somewhat higher proportion of recursive sites were detected in high-expressed genes (TPM > 20) than low-expressed genes (TPM ≤ 20). However, subsampling of the reads mapping to high-
expressed genes to levels comparable to those observed for low-expressed genes resulted in a substantially lower fraction of recursive sites at each depth, suggesting that recursive splicing is more prevalent in low-expressed than high-expressed genes (Fig 2C). Together, these data suggest that the true fraction of very long introns that contain recursive sites may be substantially higher than our observed fraction of 63–70%, i.e. that recursive splicing is likely present in almost all very long fly introns.
Recursive splice sites can be required for the processing of long introns [3]. However, it is possible that most recursive sites are functionally neutral, and that mRNA production is not impacted by their presence. The size of our dataset enabled us to examine four properties of recursive sites that could help to distinguish between these possibilities: sequence conservation; distribution in the fly genome; distribution within introns; and efficiency of splicing. In each case, the patterns observed suggest that recursive sites often have functional impact.

Both high and medium confidence recursive sites exhibited twice the level of evolutionary conservation observed in and around control AGGT motifs in long introns (Fig 3A), implying strong selection to maintain most or all of these sites. Recursively spliced introns were enriched in genes involved in functions related to development, morphogenesis, organismal, and cellular processes, with stronger enrichments observed for genes containing high-confidence recursive sites (Fig 3B; S4 Table). Both of these observations are consistent with results from a previous study based on a smaller sample of recursive introns [4].

Longer introns might contain more recursive sites purely by chance. Indeed, while the majority of recursively spliced introns had just one recursive site, the number of sites increased roughly linearly with intron length (Fig 3C). However, the positioning of recursive sites within introns was significantly biased away from a random (uniform) distribution. Instead, recursive sites in introns with only one such site tended to be located closer to the midpoint of the intron than expected by chance (Kolmogorov-Smirnov \( P = 0.003; \) Fig 3D). Furthermore, the first recursive site in introns with two or three such sites tended to be located approximately 33% and 25% of the way from the 5’ end of the intron, respectively (Fig 3E). The distribution of recursive sites within introns suggests that they are positioned so as to break larger introns into “bite-sized” chunks of intermediate size (typically ~9–15 kb in length; S5A and S5B Fig) rather than at random locations that would more often produce much longer and much shorter segments. Recursively spliced introns were also enriched in first introns, which are longer than non-first introns, relative to subsequent introns in fly genes (hypergeometric \( P < 0.05 \)).

To ask whether recursive splicing contributes to the efficiency of processing of very long introns, we evaluated the order and timing of recursive splicing events (Methods). We observed a steady increase in the proportion of exon-exon junction reads relative to recursive junctions across the time course, reflecting the progress of splicing (Fig 4A). Among recursive junction reads, we observed far higher counts of reads spanning the 5’ splice site and the recursive site (RS), relative to RS-RS or RS-3’ splice site junctions, consistent with recursive segments being predominantly excised in 5’ to 3’ order (Fig 4A; S5C Fig). This order of splicing is consistent with recursive splicing occurring co-transcriptionally. Using targeted RT-PCR amplification of segment combinations in nascent RNA from 3 recursively spliced introns, we were only able to detect products spanning the 5’ splice site and recursive site (S3B Fig). Surprisingly, we did detect a product spanning the recursive site and 3’ splice site for the third recursive site of the Ten-m intron in steady-state cDNA (S3C Fig; validated by sequencing, S3D Fig), indicating that splicing of downstream recursive segments can sometimes occur before splicing of initial segments. Finally, we also detected one read that spans a lariat resulting from a RS-RS junction (S2 Table), as well as four reads (for three junctions) that span lariats resulting from the excision of recursive introns in one segment (5’-3’ junction). The observation of these reads indicates that these introns are not always recursively spliced, though we note that these lariats are from introns that are much shorter than typical recursive introns (1.7–2.5 kb).

Previously we developed a framework for estimating rates of splicing from nascent RNA sequencing data across different labeling periods [9]. Here, we adapted this approach to estimate the splicing half-lives of individual recursive segments (Methods; S5D Fig; S5 Table),
which have a mean length of 9.1 kb (S5A Fig). Recursive segment half-lives were the slowest for the first segment in the intron, with faster half-lives for successive segments (S5E Fig). Overall, recursive segments had 1.5-fold longer half-lives than non-recursive introns of the same lengths (Fig 4B; Mann-Whitney $P = 1.5 \times 10^{-9}$). Estimating the mean splicing half-life of a recursive intron as the maximum of a set of exponentials (to approximate the waiting time to splice all recursive segments), we found that recursive introns are spliced more slowly than non-recursive introns of similar size (S5F Fig; Mann-Whitney $P < 2.2 \times 10^{-16}$), consistent

Fig 3. Characteristics of recursive sites in Drosophila introns. (A) Conservation of sequences around all detected recursive sites, with average phastCons scores for medium-confidence recursive sites (yellow), high-confidence sites (gold), and random AG|GT sites in introns increasingly larger than 1kb (grey). (B) Enrichment of significant gene ontology categories among genes with any recursive site (yellow) and high confidence sites (gold), where gene ontology terms are broken down into their umbrella categories. (C) Full intron length distributions for introns (y-axis) with varying numbers of recursive sites (x-axis). (D) Relative positions of recursive sites within introns for random sites chosen from a uniform distribution (grey) and single recursive sites in an intron (dark tan). (E) Distributions of the fractional distances (y-axis) of the first recursive segment for introns with increasing numbers of recursive sites (x-axis).

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Fig 4. Recursive sites aid in efficient splicing of long *Drosophila* introns. (A) The percentage of splice junction reads (mean percentage across recursive introns; x-axis) that span the exon-exon boundary (5’ splice site–3’ splice site; dark blue), a 5’ splice and recursive site (light blue), two recursive sites (gold), and a

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Sites of recursive splicing in flies
with the larger number of biochemical steps involved in recursive versus non-recursive splicing of an intron.

To ask whether recursive splicing occurs while the intron is continuing to be transcribed, we calculated the ratio of the half-life of the first segment to the estimated time needed to transcribe the remainder of the intron (Methods). For 49% of recursively spliced introns, the first segment half-life is shorter than the time to transcribe the full recursive intron (Fig 4C), implying common co-transcriptional splicing in about half of cases. We observed that longer recursive introns were more likely to be spliced co-transcriptionally.

The accuracy of splicing is likely to be at least as important as its speed, since splicing to an arbitrary (incorrect) splice site will most often produce an mRNA that is unstable or encodes a protein that is aberrant or nonfunctional. As a simple measure of potential splicing errors, we tallied the fraction of nascent RNA reads (from the 5 minute labeling period) that spanned “non-canonical” splice junctions, involving pairs of intron terminal dinucleotides other than the three canonical pairs “GT-AG”, “GC-AG” and “AT-AC” that account for ~99.9% of all known fly introns. For the bulk of non-recursive introns (most of which are < 100 nt in length), the frequency of such non-canonical splicing was negligible (Fig 4D, black curve). However, for non-recursive introns with lengths matching the much more extended lengths of recursively spliced introns, potential splicing errors were much more frequent (Fig 4D, gray curve), suggesting that the fly spliceosome loses accuracy as intron length (and the number of possible decoy splice sites) increases. Notably, recursive introns had ~37% fewer non-canonical junctions compared to similarly sized non-recursive introns (Fig 4D, gold curve, Kolmogorov-Smirnov P = 0.015). Therefore, presence of recursive splice sites may increase the accuracy of splicing, perhaps at the expense of splicing speed.

**Discussion**

Analysis of intermediates can provide insight into otherwise hidden biochemical pathways. Here, application of new computational approaches to nascent RNA sequencing data, which is highly enriched for splicing intermediates, enabled us to identify about four times more recursive sites in the *Drosophila* genome than were known previously. The surprisingly widespread occurrence of recursive splicing raises questions about what functions it may serve.

A priori, this pathway might improve the speed or accuracy of splicing, or might impact regulation. Our analyses suggest that recursive splicing does not in fact increase splicing rates, and may actually slow splicing somewhat, likely because of the additional steps involved. However, we observe that the *Drosophila* splicing machinery appears to make a relatively high rate of errors in the splicing of longer introns, and that presence of recursive sites may substantially improve splicing accuracy. In splicing of a non-recursive 30 kbp intron, the 5’ splice site is synthesized about 20 minutes before its correct partner 3’ splice site, creating a long window during which splicing can only occur to incorrect 3’ splice sites, likely contributing to the higher error rate seen for long fly introns. Presence of a recursive site may help to organize the processing of the intron, keeping the splicing machinery associated with the 5’ splice site engaged
in a productive direction and avoiding engagement with decoy 3’ splice sites. It was previously observed that masking a recursive splice site in a zebrafish cadm2 intron does not change the overall splicing of the intron but reduces cadm2 mRNA levels [5]. This observation could be explained if the recursive site promotes accurate splicing and prevents unproductive splicing pathways that result in unstable products targeted by RNA decay pathways such as nonsense-mediated mRNA decay.

Recursive sites may also participate in splicing regulation. A previous study of a handful of recursively spliced introns in humans identified RS-exons that are initially recognized during recursive splicing via an “exon definition” model of splice site recognition [5], while an alternative “intron definition” pathway has been proposed for recursive splice site recognition in flies [4]. An exon definition model would require presence of a 5’ splice site downstream of each recursive site. Consistent with this model, we observed that recursive segments following recursive sites are enriched for strong 5’ splice site motifs relative to first recursive segments and relative to non-recursive introns matched for length (Fig 4E). Use of an exon definition pathway in the initial steps of splicesome assembly might also contribute to splicing accuracy, with the downstream 5’ splice site helping to specify the recursive site [9]. It could also produce alternative mRNA isoforms containing an additional exon [5].

Exon definition of recursive segments through transient RS-exons requires that the recursive site first be recognized as a 3’ splice site and subsequently as a 5’ splice site for splicing of the subsequent segment (assuming that simultaneous recognition of an RS in both modes is sterically prohibited). For this ordered recognition to occur (and for sequential splicing of recursive sites generally), binding of dU2AF/U2 snRNP must outcompete binding of U1 to the RS prior to its splicing to the upstream exon. Consistent with this expectation, the 3’ splice site motifs of RS are very strong, stronger than non-recursive 3’ splice sites, and they have higher information content than RS 5’ splice sites (S6 Fig).

Developmental genes are enriched for long introns, which are more likely to be recursively spliced, but explanations for this pattern remain murky. It is possible that intron length is used to tune the timing of expression of these genes relative to the rapid embryonic cell cycle [12,13]. Alternatively, long introns may be needed to accommodate large transcriptional enhancers or complex three-dimensional organization of these gene loci related to their dynamic transcriptional regulation, or to facilitate alternative splicing. Thus, it is unclear whether recursive splicing is a feature of developmental genes or exists to facilitate the splicing of long introns that independently persist in developmental genes. In addition to producing unstable mRNAs, splicing errors may also produce stable mRNAs that encode aberrant protein forms, including dominant negative forms. Perhaps recursive splicing has been selected for in these genes to improve splicing accuracy and avoid production of aberrant developmental regulatory proteins at critical stages to improve the robustness of development.

**Methods**

**RNA-seq data analysis**

We used RNA-seq data from our recent study of splicing kinetics in Drosophila S2 cells (GEO GSE93763; [9]). These data included 3 independent replicates of S2 cells labeled for 5, 10 and 20 minutes with 500 μM 4-thiouridine, isolation of labeled RNA, and library preparation using random hexamer priming following ribosomal RNA subtraction. Separation of total RNA into newly transcribed and untagged pre-existing RNA was performed as previously described [14,15], where 4sU-labeled RNA was selectively biotinylated and captured using streptavidin coated magnetic beads. cDNA for two independent biological replicates of “total” RNA were prepared using an equal mix of random hexamers and oligo-dT primers from unlabeled S2
Cells [9]. Libraries were sequenced with paired-end 51 nt reads (100 nt reads for the “total” RNA samples), generating an average of 126M read pairs per library. Reads were filtered and mapped to the *Drosophila melanogaster* dm3 reference assembly as described in [9].

Gene expression values (TPMs) in each replicate library were calculated using Kallisto [16] and the transcriptome annotations from FlyBase *Drosophila melanogaster* Release 5.57 [17].

**Identifying sites of recursive splicing**

We used three features of recursive sites found in our nascent sequencing data to identify recursive sites: (1) splice junction reads derived from putative recursive sites (“RachetJunctions”), (2) recursive-site spanning pairs, specifically read pairs that map to sites flanking putative recursive segments such that the fragment length can only be accounted for by the presence recursive intermediate (“Rachet Pair”), and (3) a sawtooth pattern in intronic read density (“RachetScan”). Details of the computational and statistical methods for each of these approaches and our pipeline for recursive site detection are described below.

Out of the full set of recursive sites that were identified across all three methods, we filtered down to a final set of sites with the following criteria: (1) in genes with TPM $\geq 1$ in the total RNA libraries, (2) in introns with at least 3 reads spanning the 5’ to 3’ splice sites (using the largest annotated intron), and (3) not overlapping with an annotated 5’ splice site in the that intron. We ran our final pipeline on reads pooled across replicates and labeling periods to increase detection power. This resulted in a total of 539 recursive sites identified by any method. High-confidence sites were identified by the criteria used by Duff *et al.* [4]. We wrote a script to plot the read density around putative recursive sites and manually filtered each site based on the presence of a recognizable sawtooth pattern. This resulted in the identification of 243 high-confidence sites.

Conservation of recursive sites was estimated using per nucleotide phastCons scores [18] from a 15-way *Drosophila* alignment downloaded from UCSC Genome Browser.

**RachetJunction:** Identifying splice junction reads from recursive intermediates. Splice junction reads that span putative recursive junctions provide direct evidence for recursive splicing (Fig 1A bottom). In order to identify such reads, we extracted the coordinates of annotated introns and exon-exon junctions from FlyBase *D. melanogaster* Release 5.57 and aligned the 4sU-RNAseq reads to the corresponding genome release using hisat2 [19]. We then used pysam [20] to extract reads with an upstream junction matching an annotated 5’ splice site and a downstream end mapping to an AGGT that is upstream of the downstream most corresponding annotated 3’ splice site.

**RachetPair:** Identifying recursive-site spanning pairs. In addition to splice junction reads, read pairs with one end on either side of a recursive splice junction—henceforth referred to as recursive junction spanning read pairs—provide evidence for recursive sites. We defined putative recursive junction spanning read pairs as read pairs with a first read aligning close upstream of an annotated 5’ splice site and a second read aligning to an intronic region more than 1000 nt downstream of the first read. Additionally, we filtered out read pairs than have an insert length of less than 1000 nt conditioned on completion of an annotated splicing event (excluding cassette exons with an AGGT at their 5’ end).

Unlike splice junction reads, recursive junction spanning read pairs do not immediately implicate a specific recursive site. Instead, a recursive site must be inferred based on the empirical insert length distribution and genomic sequence information. To do this, we adapted the GEM algorithm, which was originally used to infer protein binding sites from ChIP-seq data [10].
Our modifications to the algorithm and choices for parameters described in Guo et al. are as follows:

1. The probability of a read, \( r_m \), given that there is a recursive site at position \( m \), \( P(r_m|m) \), was defined as the probability of observing the implied insert length in the empirical insert length distribution.

2. The prior probabilities of each position being a recursive site, \( P_{1-30} \), were set such that \( \prod_i / \max(0, M(i) - 0.8) \), where \( M(i) \) is the motif score for position \( i \) as described below. This function was used to determine the prior probabilities that reflect the preference for strong motifs observed in the Duff et al. set of recursive sites [4].

3. Recursive splice junction reads were counted within the number of effectively assigned reads in the M-step. This ensured that sites with support from recursive junction reads are more likely to be recursive sites.

4. The sparsity parameters, \( \alpha_s \), was defined as the number of assigned reads divided by 40.

5. The algorithm converged when prior probability did not change by more than \( 10^{-5} \) between iterations. Upon convergence, read pairs were assigned to a putative recursive site using the MAP estimate.

The modified GEM algorithm was run with all read pairs and splice junction reads pooled together.

**RachetScan: Identifying sawtooth pattern in recursive intron read density.** Recursively spliced introns contain a distinct “sawtooth” pattern due to the co-transcriptional nature of splicing. This is depicted in Fig 1B, where the horizontal lines represent elongating pre-mRNAs—with a uniform distribution of elongation distances across a population of cells over time—and the blacked out sections represent segments that have already been spliced out and degraded. Reads sequenced from the nascent RNA population will only be derived from the sections of RNA that have not yet been spliced and degraded, such that their density across the intron exhibits linear decay across each recursive segment.

We developed an algorithm to predict recursive splice sites from the presence of a sawtooth pattern in introns. Our algorithm consists of three distinct phases: pre-processing of the RNA-seq data, Monte Carlo Markov Chain based inference of the presence of a sawtooth pattern, and the prediction of recursive sites based on the output of our inference and sequence information.

RNA-seq read pre-processing (visualized in S1A Fig). We searched for the presence of a sawtooth pattern in the read distribution of all introns over 8 kb that had at least one spanning splice junction read in any sample. Empirical testing suggested our method displayed a high rate of false positives in introns under 8kb, likely due to regression over short segments being more sensitive to noise in read density. We removed regions annotated as exons using bedtools subtract. The number of read pairs aligning to each position were summed to obtain per base coverage counts, where read pairs straddling a given position were counted as a positive alignment.

In order to avoid erratic read coverage in repeat regions inhibiting our ability to perform meaningful regressions in later steps of the analysis, we masked the read densities in repeat regions and replaced the read counts in RepeatMasker annotated repeat regions [21] and the 100 flanking nucleotides with the median read density from the 900 nt flanking either side. This length was chosen because it was short enough that read densities in this range were comparable to those in the masked region, but long enough to avoid sensitivity to noise in read densities. To attain additional smoothing and reduce the time required to perform the
regressions in the next step of our analysis, we separated introns into 100 nt bins and calculated the average of each bin. Throughout the rest of our analysis, we represented the read density of each intron using arrays of these average values.

**Regression.** We performed linear regression on all sub-regions of each intron. We assumed that variance in read density at each position was proportional to the coverage level at that position, which is likely true since RNA-seq read coverage is intrinsically the sum of Bernoulli random variables. To calculate these regressions, we developed a function that made use of the Scipy stats weighted linear regression function \[22\] as a sub-process, such that:

**Algorithm:** Heteroscedastic Regression  
**Data:** A ← Array of RNA-seq data  
**Result:** slope, yInt, and weights for regression

```plaintext
Algorithm: Heteroscedastic Regression
Data: A ← Array of RNA-seq data
Result: slope, yInt, and weights for regression
nextW ← [1..1];
curW ← [0..0];
while |curW - nextW| ≤ 10^{-3} do
    curW ← nextW;
    slope, yInt ← regression(curW, A);
    for position ∈ intron do
        nextW[position] ← \frac{1}{yInt + \text{slope} \times \text{position}};
    end
end
return slope, yInt, nextW;
```

Note that \(|\text{curW} - \text{nextW}| \leq 10^{-3}\) checks whether all weights have changed by at most \(10^{-3}\).

**Monte Carlo Markov Chain (MCMC; visualized in S1B Fig).** We developed a Monte Carlo Markov Chain (MCMC) algorithm to detect the presence of a sawtooth pattern in each intron. Our algorithm is round based such that upon entering each round, we have an accepted state consisting of a set of proposed recursive sites in the intron. In each round, a new state is proposed by perturbing the current state. We use a scoring function and transition rules (defined below) to decide if we wish to accept this proposed state or continue with the current state. This procedure is iterated for \(10^7\) rounds and a sample of the current state is recorded every 50 states. The number of samples recorded in each state is proportional to the probability that the intron is best fit by the model corresponding to that state. Therefore, to attain probabilities that each state is the most accurate model, we normalize the number of samples recorded in each state by the total number of samples.

There are three classes of perturbations used to propose new states (depicted in S1B Fig):

1. A new recursive site was added probabilities 0.4 (visualized as transition 1 & 2).
2. A recursive site was removed with probability 0.4 (visualized as transition 4).
3. A recursive site was slightly perturbed with probability 0.2 (visualized as transition 3).

States are scored using a function taking into account how well the corresponding regression fits the observed RNA-seq read density as well as the number of free parameters in the model. The scoring function is based on the Bayesian Information Criteria (BIC), such that:

\[
\text{BIC}(M) = L \times \text{RSS}(M) + 2 \times (2N) \times \log(L)
\]

where \(\text{RSS}(M)\) is the weighted sum of squared deviations for all recursive segments, \(L\) is the intron length, and \(N\) is the number of recursive sites. Note that \(2N\) is the number of free parameters in the model, as each recursive segment is fit for its own slope and y-intercept. The score is then given by:

\[
\text{Score}(M) = e^{-\frac{\text{BIC}(M)}{2}}
\]
where T is a constant used to scale the magnitude of the scores. T = 5 was used for all analyses presented here. In order to constrain our algorithm to fit sawtooth patterns and note more general patterns in the read density, new states are only considered if, at each recursive site, the RNA-seq density predicted by regressions increased by at least 1.5 fold.

We use the standard transition rules for MCMC inference, which we outline here for convenience. If the score for the new state is lower than the score for the old state, the new state is deterministically adopted. Otherwise, the new state is adopted with probability \( \frac{\text{score}_{\text{new}}}{\text{score}_{\text{old}}} \). When the old state had zero recursive sites, this probability was divided by 2 to account for the imbalance in transition probabilities. We chose parameters for burn-in-time, number of iterations and sampling frequency that empirically resulted in consistent convergence across multiple runs of the algorithm. These values were: a burn-in of \( 10^5 \) iterations, sampling frequency of 50 iterations, a total of \( 10^7 \) iterations.

After all samples were collected, we calculated the probability that each position in the intron is a recursive site. For each position, we summed the occurrences of that position as a recursive site across all samples. Probability scores were then calculated for each position by dividing this sum by the total number of samples.

Peak Calling (visualized in S1C Fig). We predicted recursive sites from the MCMC probability scores in a two-step process. First, regions with probability above a given threshold (0.08) were recorded. Any of these regions within 500 nt of each other were merged. For each of these regions, a position potential function, \( P \), was defined as 1 inside the peak and flanked by a logistically decaying curve on either side. The logistic function is given by:

\[
f(x) = \frac{1}{1 + e^{-k(x-x_0)}}
\]

The parameters were set as \( x_0 = 500 \) nt from either end and \( k \) was set as 6/500 for the left flank and -6/500 for the right flank. The resulting distribution has values very close to zero at 1000 nt away from the peak and values of 0.5 at a distance of 500 nt. This distribution was chosen based on the empirical performance of the MCMC-based inference when compared to random. Each AGGT in the intron was then scored by the following equation and the maximum scoring AGGT was then reported as a putative recursive site:

\[
S(i) = P(i) * \max(M(i) - 0.8, 0)
\]

FDR Quantification. Shuffled peaks were produced to evaluate the false discovery rate of the sawtooth pattern identification pipeline. For each intron, the initially recorded regions of probability exceeding 0.08 were redistributed with uniform probability across the intron. The length and number of regions were maintained. The remainder of the peak calling procedure was then applied to obtain a null distribution of recursive probability peaks.

Motif scoring
We calculated position weight matrices (PWM) for the intronic portions of Drosophila 5' and 3' splice sites using all annotated splice sites. These weight matrices were then juxtaposed with the 3' splice site PWM followed by the 5' splice site PWM to create a recursive splice site motif PWM. Individual motif occurrences were scored using a normalized bit score [23]. The bit score for each motif occurrence is defined as the sum across the log probabilities for each nt being drawn from the motif. We calculated normalized scores by subtracting the minimum possible score and dividing by the range of possible bit scores.
Identification of reads from lariat junctions

We searched for reads crossing the 5'SS-branch point junction using code previously developed in our lab (https://github.com/jpaggi/findbtps). In short, our approach works by: (1) identifying reads that do not have a valid alignment; (2) splitting the unalignable reads just before the 7-mer best matching the consensus 5'SS motif; and (3) mapping this split read as a pair, requiring that the second segment align upstream of the first segment. We require the following features to be present: (1) each segment must be at least 15 nt long; (2) only 1 mismatch is allowed per segment; (3) segments must be separated by less than 1 Mbp; and (4) the pair has a unique alignment.

We then filtered the resulting alignments for cases where the second segment aligned immediately downstream of a 5'SS or recursive site and the first segment aligned within 100 nt upstream of a 3'SS or recursive site. Overall, we detected 323 5'SS-branch point junction reads across 319 introns. The putative branch points show a motif favoring an A at the branchpoint and a U at the -2 position, consistent with the human branchpoint consensus motif. We observed 7 5'SS-branchpoint junction reads from introns that we report to be recursively spliced. These counts are consistent with analysis by Duff et al., which identified 46 recursive lariat junction reads amongst 10.2 billion reads. If such reads occurred at the same frequency in our data, we would expect to observe 1.8 recursive lariat junction reads. All implicated branch points are adenosines. These 7 reads implicate a lariat associated with the following categories of splicing events (S2 Table): (1) 5'SS-RS: two reads associated with two unique junctions; (2) RS-RS: one read associated with one junction; and (3) 5'SS-3'SS: four reads associated with three unique junctions. The 5'SS-3'SS lariat junction reads suggest that recursive splicing is not always used for these introns. All three such junctions derived from introns of lengths far shorter than typical recursive introns (1762 nt, 1929 nt, and 2548 nt), suggesting that non-recursive splicing may compete with recursive splicing of introns in this size range.

PCR validation of novel recursive sites

Nascent RNA was isolated after 5 minutes of labeling with 4sU (as described above) and reverse transcribed to first-strand cDNA using ProtoScript II Reverse Transcriptase (M0368S, NEB) primed with random hexamers according to manufacturer’s protocol. cDNA was diluted 1:5 and 1μL was used as template for PCR reactions using primers designed to amplify recursive segments anchored by either the intronic 5’ splice site or intronic 3’ splice site (S3 Table). PCR amplification was performed using Taq DNA Polymerase (10342020, Invitrogen) for 40 cycles. PCR products were visualized on a 1.5% agarose gel relative to Azura PureView 50bp DNA ladder (AZ1155, Azura). PCR products were purified with a DNA Clean & Concentrator kit (D4033, Zymo) and Sanger sequenced to confirm the junction boundaries.

Estimating the true number of recursive sites

In order to assess the sensitivity of our recursive site detection pipeline, we subsampled our reads to various proportions of the total read coverage and re-assessed the number of recursive sites detected. To do so, we used the samtools view -s command [24] to subsample each fastq file from all samples to the following fractions: 0.1%, 0.5%, 1%, 2.5%, 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, and 90%. For each of these subsampled read sets, we re-ran the entire recursive site detection pipeline as described above to assess the number of recursive sites detected.

To assess the impact of gene expression levels on our power to detect recursive sites, we separated long introns into those from lowly expressed genes (TPM ≤ 20) and highly expressed genes (TPM > 20). Using the subset of reads mapping to these genes, we repeated the
subsampling procedure and entire recursive site detection pipeline described above to characterize the percentage of lowly or highly expressed long introns that have recursive sites.

Finally, to understand whether the lower proportion of lowly expressed introns that have recursive sites is due to technical or biological reasons, we subsampled reads from long introns within highly expressed genes to match the read distribution of a comparable number of long introns from lowly expressed genes. Specifically, we isolated all reads from long introns in highly expressed genes and used pysam [20] to randomly subsample these reads to match the distribution of reads from lowly expressed introns. Using only this subset of reads from highly expressed genes, we again repeated the subsampling procedure and the entire recursive site detection pipeline described above to characterize the percentage of highly expressed introns that have recursive when reads from these introns are subsampled to a lower read coverage.

Determining the order of recursive splicing

Previous studies have searched exclusively for recursive junction reads consistent with the 5' to 3' removal of recursive segments [4,5]. In order to determine if recursive splicing does indeed follow a 5' to 3' order, we quantified junction reads consistent with alternative orders of recursive splicing. These reads fall into two categories: junction reads between two intronic AGGTs and junction reads from an intronic AGGT to an annotated 3' splice site.

We constrained our search to combinations or recursive sites producing recursive segments of at least 1 kb. Nearly all recursive segments detected in our study were greater than 1 kb, thus adding this constraint mainly served to filter out spurious hits likely caused by alignment errors and unannotated splicing events. We considered all events with support from at least 3 uniquely aligning reads with recursive splice sites scoring above 0.85 in the scoring metric described above. Requiring at least three uniquely aligning reads matches the cutoff used for our previous analysis, where we found that recursive splice sites generally have strong motifs that score greater than 0.85.

These analyses produced thirteen candidate intronic AGGT to annotated 3' splice site recursive junction reads, and no candidate intronic AGGT to AGGT recursive sites. These candidate recursive splice sites were evaluated visually in a genome browser. Two of these sites corresponded to recursive splice sites detected by both methods in our study. One of these sites has sixty recursive junction reads supporting a 5' to 3' order, while only five junction reads support a 3' to 5' order. The second site has 829 and 13 junction reads for the 5' to 3' and 3' to 5' orders, respectively. All other candidate alternative ordering sites did not appear to be represent viable recursive site candidates, due to either a lack of sawtooth pattern, low intron expression, or extensive repeats complicating the alignment. These data suggest that recursive splicing overwhelmingly, but perhaps not always, proceeds in a 5' to 3' order.

Splicing rates in recursively spliced introns

We quantified splicing rates for each recursive segment independently by applying an approach for 4sU RNA-seq data that we previously described [9]. Specifically, we used reads that overlapped recursive sites and junction reads (split between either the recursive site and an annotated splice site, between two recursive sites, or between the 5' and 3' splice sites; as detailed in S5D Fig), as measures of uncompleted and completed segment splicing, respectively. The junction dynamics approach from Pai et al. 2017 [9] was applied to each set of reads to obtained a splicing half-life for each recursive segment. For full introns matched for length, we used splicing half-lives calculated in Pai et al. 2017 [9]. We estimated co- vs. post-transcriptional splicing of the first recursive segment by comparing the segment splicing half-life to the time to transcribe the remainder of the intron. Specifically, the time to complete intron
transcription was estimated as:

\[
3'\text{transcription} = \frac{\text{length of intron (nt)} - \text{length of first segment (nt)}}{1,500 \text{ nt/min}}
\]

and the splicing delay was calculated as the ratio of the first segment’s splicing half-life to the 3’ transcription time.

**Estimating the rate of splicing of full recursive introns.** To estimate the mean lifetime of a recursively spliced intron, we estimated the waiting time for all recursive segments to be spliced out by calculating the maximum of the set of individual exponentials from each segment. For one exponential, the mean lifetime is \( \tau = \frac{1}{\lambda} \) where \( \lambda \) is the coefficient from the exponential fit. There is an analytical solution for estimating the mean lifetime in situations where there are only two exponentials to be combined. Thus, we limited our analysis to recursive introns with only one recursive site, corresponding to the presence of two recursive segments (i.e. two exponentials). For these introns, the mean lifetime \( \tau_{\text{recursive}} \) can be calculated by:

\[
\tau_{\text{recursive}} = \frac{1}{\lambda_1} + \frac{1}{\lambda_2} + \frac{1}{\lambda_1 + \lambda_2}
\]

where \( \lambda_i \) is the exponential coefficient for the first segment and \( \lambda_j \) is the exponential coefficient for the second segment. To conservatively compare our recursive intron \( \tau_{\text{recursive}} \) values with the mean lifetimes of non-recursive introns, we added the time necessary for the first segment to be transcribed to \( \tau_{\text{recursive}} \), the rationale being that the first segment must be completely transcribed before the second can begin to be spliced. Assuming a 1.5 kb/min transcription rate, \( \text{txn}_{\text{seg1}} = \frac{l_1}{1500} \), where \( l_1 \) is the length of the first segment (in nucleotides).

**Estimating splicing accuracy**

We estimated the accuracy of splicing in *Drosophila* introns by identifying non-annotated junction reads with non-canonical splice site sequences within annotated introns within the nascent RNA reads from the 5 minute labeling period. To do so, we first re-mapped the raw 4sU-seq reads with the STAR v2.5 software [25], with the mapping parameter—outSAMattribute NH HI AS nM jM to mark the intron motif category for each junction read in the final mapped file.

The jM attribute adds a jM:Bc SAM attribute to split reads arising from exon-exon junctions. All junction reads were first isolated and separated based on the value assigned to the jM:Bc tag. Junction reads spanning splice sites in the following categories were considered to be annotated or canonical: (1) any annotated splice site based on FlyBase *D. melanogaster* Release 5.57 gene structures [jM:Bc,[20–26]], (2) intron motifs containing “GT-AG” (or the reverse complement) [jM:Bc:1 or jM:Bc:2], (3) intron motifs containing “GC-AG” (or the reverse complement) [jM:Bc:3 or jM:Bc:4], and (4) intron motifs containing “AT-AC” (or the reverse complement) [jM:Bc:5 or jM:Bc:6]. Junction reads with jM:Bc:0 were considered to arise from non-canonical non-annotated splice sites. We calculated the frequency of inaccurate splice junctions for each intron as a ratio of the density of reads arising from non-canonical non-annotated splice sites to the density of all junction reads from the intron.

**Calculating splice site scores**

We calculated the strength of splice sites using a maximum entropy model as implemented in maxEntScan [26] using 9 nucleotides around the 5’ splice site (-3:+6) and 23 nucleotides around the 3’ splice site (-20:+3). These models were optimized on mammalian splice site
preferences, but seem to be reasonable for Drosophila as well and have been used in gene prediction in fly genomes.

**Gene ontology analyses**

Gene Ontology enrichment analyses were performed using a custom script to avoid significant gene ontology terms with overlapping gene sets. Specifically, the script used the Flybase gene ontology annotation downloaded from the Gene Ontology Consortium website [27] and searches for the gene ontology term with the most significant enrichment of genes with recursively spliced introns (relative to a background of all genes with introns greater than 10,000 kb). Genes that belong to the most significant gene ontology term are then removed from the foreground and background sets of genes and the process is repeated iteratively until no genes are left in the foreground set. P-values are computed using a Fisher-exact test and then corrected using a Benjamini-Hochberg multiple test correction.

**Code availability**

Source code for our pipeline to identify recursive splicing sites is available at https://github.com/jpaggi/recursive.

**Supporting information**

**S1 Fig. RachetScan method for automated detection of sawtooth patterns indicating of recursive splicing.** (A) RNA-seq pre-processing steps to convert reads into an array of read densities: (1) summing the read coverage for each base-pair (top) and (2) replacing the read counts in annotated repeat regions and 100 flanking nt with median read density in 900 nt flanking regions (bottom) (B) MCMC algorithm infers probability that each position in intron is a recursive splice site, where upon entering each round with a previously accepted state, this state is perturbed to propose a new state and the new state is either accepted or rejected. The procedure is performed over $10^7$ rounds, with sampling every 50 rounds to obtain a probability that each base pair is a recursive site. (C) Sequence information is used in conjunction with MCMC-inferred probabilities to predict recursive sites. (TIFF)

**S2 Fig. Identifying sites of recursive splicing.** (A) The probability derived from the sawtooth MCMC model of a site being a recursive site for the final set of recursive sites (light orange), all sites with minimal support from any method (dark orange), and random sites placed down in the same introns (grey). (B) The sawtooth score (see Methods) for the final set of recursive sites (light orange), all sites with minimal support from any method (dark orange), and random sites place down in the same introns (grey). (C) Number of recursive sites (left) and high-confidence sites (right) identified by one of multiple identification pipelines, with the majority of recursive sites identified by both junction reads and sawtooth scores, as well as present in the Duff et al. dataset. (D) The gene expression levels of genes with recursive introns (TPM, y-axis) relative to the junction spanning read support for each recursive intron (read count, x-axis), showing the varying power to identify recursive sites with the sawtooth recursive method (orange), junction-spanning reads alone (blue), or both methods (black). (E) The cumulative distribution of distances between the recursive site identified with the sawtooth recursive method and the best matching recursive motif (orange) and random sites placed down in the same introns (grey) are significantly different. (PDF)
S3 Fig. Experimental validation of novel recursive sites. PCR products from recursive segments anchored to intronic 5’ splice sites (A) and intronic 3’ splice sites (B), for recursively spliced introns from three genes: Ten-m (left), Luna (middle), and Tet (right). PCR was performed on first-strand cDNA from nascent RNA from Drosophila S2 cells isolated after 5 minutes of labeling with 4sU and bands were visualized on a 1.5% agarose gel with a 50bp ladder. Schematics indicate the junctions across which the amplicons were designed. Junction boundaries were confirmed by sequencing for lanes 3, 5, 17, 18, and 19 in (A). (C) PCR products across recursive segments and intronic 3’ splice sites from first-strand cDNA from steady-state RNA for the Ten-m recursively spliced intron. The junction boundary was confirmed by sequencing for lane 4. (D) Representative sequence traces confirming the junction boundaries for two novel 5’-RS recursive splicing events (top and middle) and one RS-3’ event (bottom). Peaks delineate specific nucleotides, including A (green), C (blue), G (black), and U (indicated by a T, red). For each of these events, the full band was sequenced (164nt, 153nt, ad 159nt from top to bottom respective), however only 30nt around the junction is shown here for visualization purposes.

(TIF)

S4 Fig. Properties of recursively spliced introns. Sequence logo for all intronic AG|GT sites (top), medium-confidence recursive sites (middle) and high-confidence recursive sites (bottom).

(PDF)

S5 Fig. Rates of recursive splicing. (A) Distribution of lengths of recursive segments (nucleotides, x-axis) for medium-confidence recursive segments (yellow) and high-confidence recursive segments (gold). (B) Recursive segment length distributions (nt, y-axis) for introns with varying numbers of recursive sites (x-axis). (C) The number of splice junction reads (y-axis) spanning a 5’ splice site and recursive site (blue), two recursive sites (gold), and a recursive site and 3’ splice site (yellow) across the labeling periods (x-axis). (D) Junction reads used to estimate splicing half-lives for recursive segments (red lines), centered on 3’ recursive sites (red dots) for each segment. Incomplete splicing is estimated from intron-exon junction reads (pink bars). Completed splicing is estimated from a sum across split-junction reads between the 5’ splice site and recursive site (light blue bars), two recursive sites (orange bars), a recursive site and the 3’ splice site (yellow bars), and the 5’ splice site and 3’ splice site (exon-exon read, dark blue bars). Each segment’s splicing is informed by different types of junction reads dependent on the position in the intron, as drawn for an intron with three recursive segments. (E) Splicing half-lives (y-axis) for recursive segments with varying positions across the intron (x-axis), where on average, all segments in an intron tend to be spliced out at similar rates. (F) The distribution of mean life-times (y-axis) for recursively spliced introns (estimated by the maximum of exponentials from constituent recursive segment splicing rates, gold) relative to non-recursive introns chosen to match the length of the recursive introns (grey).

(PDF)

S6 Fig. Recursive splice site motif strength. Distribution of splice site strengths (maxEnt score, y-axis) across both 3’ splice sites (orange) and 5’ splice sites (blue) for recursive sites (right) and non-recursive introns matched for intron length (left). Significance is indicated such that **: P < 0.01 and ***: P < 0.001, with a Mann-Whitney U test.

(PDF)

Column 3 –TPM: Gene expression values calculated using kallisto (TPMs).
Column 4 –completed_splicing_junction_reads: Number of junction reads supporting completed splicing across the entire intron.
Column 5 –recursive_site: Coordinate for the recursive site.
Column 6 –method: Method used for identification of the recursive site, where “junction” indicates site identified by either RachetJunction or RachetPair, “sawtooth” indicates site identified by RachetScan, and “both” indicates site identified by both methods.
Column 7 –in_duff: Flag indicating the recursive site was identified in the Duff et al. study.
Column 8 –high_confidence: Flag indicating the recursive site was identified as a high-confidence site (1) or a medium-confidence site (0).
Column 9 –junction_reads: Comma-separated list of the number of junction reads (5′-RS) supporting the recursive site in each timepoint (combined across replicates) [5m, 10m, 20m, total].
Column 10 –spanning_read_pairs: Comma-separated list of number of spanning read-pairs supporting the recursive site in each timepoint (combined across replicates) [5m, 10m, 20m, total].
Column 11 –sawtooth_score: Sawtooth score for the recursive site, as defined in the Methods.
Column 12 –mcmc_probability: Probability of this site being a recursive site, as derived from the MCMC sampling procedure integral to the RachetScan method.
Column 13 –recursive_index: Recursive index for the recursive site, as defined in the Methods.
Column 14 –motif: Sequence found around the recursive site.
Column 15 –motif_score: Motif score for the recursive site, as defined in the Methods.
Column 16 –downstream_reads: Number of splice junction reads originating from the 3′ end of the exon.
Column 17 –intron_body_reads: Number of reads in the body of the intron.

S2 Table. Recursive branchpoints from recursive lariat spanning reads. Column 1 –intron coordinates: Coordinates of intron containing recursive site, with chr:start-end:strand.
Column 2 –junction type: Type of junction the lariat spans (5′SS-RS, RS-RS, 5′SS-3′SS)
Column 3 –junction coordinates: Coordinates of the junction that the lariat spans
Column 4 –branchpoint: Coordinate of the branchpoint identified by the lariat spanning read
Column 5 –sequence: Sequence overlapping the branch point (the putative branch point nucleophiles is at base 11)
Column 6 –timepoint: labeling period in which the lariat read was identified
Column 7 –read ID: Identifier of the lariat spanning read.

S3 Table. Primers used for experimental validation of recursive sites. Column 1 –PRIMER PAIR: Name of primer pair used
Column 2—Name: Name of individual primer (note, individual primers are repeated across different primer pairs; primers marked as “Duff” were used in Duff et al. 2015)
Column 3—Sequence: Sequence of individual primer
Column 4 –Expected amplicon size: Expected amplicon size if recursive segment has not yet been spliced out. Molecules where the recursive segment has been spliced will not be amplified with these primer combinations.

S4 Table. Gene ontology enrichment for all recursive sites and high-confidence recursive sites.
S5 Table. Summary statistics and information about rates of recursive spliced segments.

- **Column 1 –intron**: Coordinates of intron containing recursive site, with chr:start-end:strand.
- **Column 2 –gene**: FlyBase gene symbol for parent gene.
- **Column 3 –recursive_site**: Coordinate for the recursive site (or 3’ splice site in the case of the final segment of the intron).
- **Column 4 –segment_type**: Indicator whether the segment is spliced to a recursive site ("segment") or to the 3’ splice site ("threeess", in the case of the final segment of the intron).
- **Column 5 –segment_len**: Length of the segment (nucleotides).
- **Column 6 –segment_num**: Position of the segment relative to other segments in the intron.
- **Column 7 –three_length**: Length of the region from the recursive site (or 3’ splice site) to the polyA site of the transcript (nucleotides).
- **Columns 8–10 –ie_count_[timepoint]**: count of the intron-exon junction reads for each of the labeling periods (summed across three replicates per labeling period) - for recursive sites, this overlaps the recursive site, while for the final segment this overlaps the 3’ splice site.
- **Columns 11–13 –ee_count_[timepoint]**: count of the exon-exon junction reads for each of the labeling periods (summed across three replicates per labeling periods) - this includes junctions deriving from recursive intermediates, as outlined in the Methods.
- **Column 14 –halflife**: Half-life of the recursive segment computed using the junction dynamics approach described in Pai et al. 2017 (min).
- **Column 15 –txn_to_three**: Time to transcribe the remainder of the intron from the recursive site to the 3’ splice site (min).

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References


