Gene expression

McSplicer: a probabilistic model for estimating splice site usage from RNA-seq data

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Associate Editor: Anthony Mathelier

Received on August 5, 2020; revised on January 20, 2021; editorial decision on January 21, 2021; accepted on January 25, 2021

Abstract

Motivation: Alternative splicing removes intronic sequences from pre-mRNAs in alternative ways to produce different forms (isoforms) of mature mRNA. The composition of expressed transcripts gives specific functionalities to cells in a particular condition or developmental stage. In addition, a large fraction of human disease mutations affect splicing and lead to aberrant mRNA and protein products. Current methods that interrogate the transcriptome based on RNA-seq either suffer from short-read length when trying to infer full-length transcripts, or are restricted to predefined units of alternative splicing that they quantify from local read evidence.

Results: Instead of attempting to quantify individual outcomes of the splicing process such as local splicing events or full-length transcripts, we propose to quantify alternative splicing using a simplified probabilistic model of the underlying splicing process. Our model is based on the usage of individual splice sites and can generate arbitrarily complex types of splicing patterns. In our implementation, McSplicer, we estimate the parameters of our model using all read data at once and we demonstrate in our experiments that this yields more accurate estimates compared to competing methods. Our model is able to describe multiple effects of splicing mutations using few, easy to interpret parameters, as we illustrate in an experiment on RNA-seq data from autism spectrum disorder patients.

Availability and implementation: McSplicer source code is available at https://github.com/canzarlab/McSplicer and has been deposited in archived format at https://doi.org/10.5281/zenodo.4449881.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Through alternative splicing (AS), a single gene can produce multiple mRNA transcripts, or isoforms, that combine exons in alternative ways. Approximately 95% of human multi-exon protein-coding genes undergo alternative splicing (Pan et al., 2008), creating a remarkably complex set of transcripts that give specific functionalities to cells and tissues in a particular condition or developmental stage.

RNA sequencing (RNA-seq) is routinely used in genome-wide transcript analysis. This technology produces short reads from which existing methods infer and quantify RNA splicing, broadly, in one of two different ways. Methods either analyze full-length transcripts or focus on individual splicing events. Transcript assembly methods such as StringTie (Pertea et al., 2015), CIDANE (Canzar et al., 2016) and CLASS (Song and Florea, 2013) aim to identify the set of expressed full-length transcripts which in principle provides a complete picture of all splicing variations, see e.g. transcript τ₁–τ₅ in Figure 1. The transcript assembly problem is, however, ill-posed (Lacroix et al., 2008) and error-prone especially for complex genes expressing multiple transcript isoforms (Hayer et al., 2015). Event-based methods, therefore, focus on local splicing patterns such as the classical exon skipping event denoted in Figure 1, without a prior attempt to assemble or quantify full-length transcripts. The relative abundance of different splicing outcomes that can potentially be shared by multiple transcripts, can then be quantified using a simple metric such as percent spliced in (PSI) (Venables et al., 2008).

A notable exception is SUPPA (Alamancos et al., 2015) which derives PSI values from quantified transcript abundances. Event-based methods differ in the complexity of the units of AS they quantify. In the simplest case, methods such as MSO (Katz et al., 2010), SUPPA, ASGAL (Denti et al., 2018), SpliceGrapher (Rogers et al., 2012) and SplAdder (Kahles et al., 2016) identify one
of the canonical types of AS, such as exon skipping, alternative 5' and 3' splice sites, intron retentions and mutually exclusive exons (see Supplementary Fig. S1). In Figure 1, this definition would include the two simple exon skipping events between t1 and t2 and between t3 and t4, and mutually exclusive spliced exons in t1 and t4, clearly underestimating the full AS complexity across t1–t5.

Compared to these simple types of splicing events, complex events involve multiple alternative splice sites or exons and according to Vaquero-Garcia et al. (2016) constitute at least one-third of AS events observed in human and mouse tissues. Methods such as JUM (Wang and Rio, 2018), MAJQ (Vaquero-Garcia et al., 2016) and the method proposed in Oesterreich et al. (2016), therefore consider AS units that generalize simple events to more complex patterns. They quantify the relative usage of an arbitrary number of introns that share a common splice site. Since these AS units capture only the common endpoints of alternative splicing patterns, such methods need to quantify two AS units for a single exon skipping event (Fig. 1). LeafCutter (Li et al., 2018) and Whippet (Stern-Weiler et al., 2017) add further introns to AS units. At the extreme end, Whippet enumerates all possible transcript fragments that combine overlapping events and estimates their relative abundance using an EM algorithm similar to full-length transcript quantification methods such as kallisto (Bray et al., 2016).

PSGInfer (LeGault and Dewey, 2013) quantifies alternative splicing based on Probabilistic Splice Graphs (PSGs). It assigns weights to the edges of a splicing graph (Heber et al., 2002) using parameters that describe the splicing process, rather than focusing on individual outcomes of the splicing processes such as local splicing events or full-length transcripts. The parameter estimates can then be used to estimate transcript and processing event frequencies. Motivated by the work by LeGault and Dewey (2013), we similarly aim to quantify alternative splicing by building a probabilistic model as a simple approximation to the underlying splicing processes. In contrast to PSG, however, our model employs the usages of annotated as well as novel splice sites across all expressed transcripts to describe a simplified splicing process that has generated the set of expressed transcripts. Traversing the linear ordering of all exons of a gene from 5' to 3', the usage of each splice site specifies the probability with which the site is used as donor or acceptor site. For example, the usage of acceptor s2 in Figure 1 indicates the abundance of transcripts t1, t2 and t3 that ‘use’ the acceptor relative to the total output t1–t3 of the gene. Our model assumes that splice site usages are independent of each other, which allows for a computationally more efficient estimation of parameters compared to PSGInfer.

This model by definition can generate complex splicing patterns that do not rely on any predefined simple or complex AS units as event-based methods like SpAdder, MAJQ or LeafCutter do. At the same time, splice site usages that capture simultaneous changes in multiple isoforms facilitate the interpretation of point mutations that disrupt splicing as is the case in many genetic disorders (Anna and Monika, 2018). Instead of attempting to quantify each one of multiple possible effects on intron or even transcript level, a reduced splicing site usage as computed by McSplicer may directly reflect the weakening of a splice site by a point mutation in the consensus splice site sequence that is responsible for these effects, as we illustrate in our experiments on RNA-seq data from autism spectrum disorder patients (Section 3.4).

Furthermore, our method simultaneously estimates the model parameters, i.e. splice site usages, using all reads mapped to a gene locus, often resulting in more accurate estimates compared to event-based methods that use only reads directly supporting their parameters. We demonstrate the improved accuracy of McSplicer compared to existing methods in our experiments.

2 Materials and methods

A typical RNA-seq analysis workflow that uses McSplicer to estimate the usage of splice sites consists of the five steps illustrated in Figure 2. After (A) mapping reads in an RNA-seq sample to a reference genome sequence using a read alignment tool such as STAR (Dobin et al., 2013) or HISAT (Kim et al., 2015), we (B) assemble reads to full-length transcripts using methods such as StringTie (Pertea et al., 2015) or CLASS (Song and Florea, 2013) to identify annotated as well as novel splice sites. Step (B) can be omitted and instead a curated catalog of known transcripts may be provided. In both cases, McSplicer does not rely on any transcript-level phasing of exons but uses the extracted splice sites and transcription start (TSS) and end sites (TES) to (C) partition a gene into contiguous, non-overlapping segments. Segments are defined as minimal subsequences of a gene’s exons and introns that are bounded by splice sites, TSS or TES. The example shown in Figure 2C contains six such segments. We count reads that overlap distinct combinations of such segments. The precise sequence of segments a mapped read overlaps defines its mapping signature (Canzar et al., 2016). Reads that map to the same signature are equivalent in terms of the splicing pattern they represent. From signature counts, i.e. the number of reads mapping to the same signature (see Supplementary Fig. S2 for an illustration), McSplicer estimates splice site usages in step (D). Splice site usages computed by McSplicer can be leveraged in (E) different types of downstream analyses, including the quantification of various types of splicing events.

In the following sections, we introduce McSplicer’s model and algorithm for the estimation of parameters in that model. A more detailed description of the model and algorithms is provided in Supplementary Section S2. In the technical description of our model, we refer to the exon boundaries at the 5' (acceptor) splice site and at the TSS as exon start sites, and to the 5' (donor) splice sites and TES as exon end sites. The description of our model is based on single-end reads which we apply to paired-end reads in Section 3.3. In the next section, we recapitulate the commonly assumed generative model of RNA-seq that also underlies the McSplicer model. For the sake of simplicity, we introduce the model based on individual observed reads and explain how parameters can be estimated from much fewer signature counts at the end of Section 2.3.

2.1 A generative model for RNA-seq reads

Consider the RNA-seq reads that mapped to a given gene. Reads are derived from one end of each of N fragments and each read has length L. We assume that each fragment is independently generated from one of the possible transcripts allowed by our model (see next Section). In this section, we describe a generative model for the sequence of the nth read RN. The probability of RN can be written as

$$P(R_n) = \sum_t P(R_n | T_n = t) P(T_n = t),$$

where Tn represents the transcript from which Rn was generated. Following models in Li et al. (2010) and LeGault and Dewey (2013), we assume that the probability of generating Rn from a
transcript $t$ is proportional to the product of the (effective) length of the transcript, $l(t)$, and the relative abundance of the transcript, $u(t)$:

$$P(T_n = t) = \frac{l(t)u(t)}{\sum_{t'}l(t')u(t')}.$$  \hspace{1cm} (2)

The effective length of a transcript denotes the number of possible start position of a sampled read (Trapnell et al., 2010). We introduce $B_n$ that denotes the start position of $T_n$ leading to

$$P(R_n|T_n = t) = \sum_{b=1}^{L(t)} P(R_n|B_n = b, T_n = t)P(B_n = b|T_n = t).$$ \hspace{1cm} (3)

Making the simplifying assumption that $R_n$ was generated uniformly across transcript $t$, we have

$$P(B_n = b|T_n = t) = \frac{1}{l(t)}.$$ \hspace{1cm} (4)

$$P(R_n|B_n = b, T_n = t) = 1$$ if $R_n$ is identical to the sequence of length $l$ starting at a position $b$ in transcript $t$, and this probability is 0 otherwise.

### 2.2 McSplicer: an inhomogeneous Markov chain to model the relative abundance of transcripts

We propose a new model for the relative abundance of transcripts expressed by a gene, denoted by $w(t)$ in the previous section. Suppose we have obtained in step (B) in the McSplicer workflow (Fig. 2) exon start sites, $s_1, \ldots, s_{M_1}$ and exon end sites, $e_1, \ldots, e_{M_2}$, ordered by their occurrence in forward direction of a given gene. Here, we do not include the start site of the first exon and the end site of the last exon, since the former is treated differently in our model (see below) and the usage of the latter is always equal to 1 in our model. All exon start and end sites partition the gene into non-overlapping segments $X_1, \ldots, X_{M}$, where $M = M_1 + M_2 + 1$ and each segment is defined by a region enclosed by splice sites or transcription start or end sites that occur consecutively along the genome (see Figs 2C and 3). We introduce a sequence of hidden variables, $Z = (Z_1, \ldots, Z_M)$, where $Z_i$ is a binary indicator for whether the $r$th segment $X_i$ is transcribed ($Z_i = 1$). Then, a particular transcript can be represented by a sequence of states for $Z$, as illustrated for transcripts $t_1, t_2, t_3$ in Figure 3. Thus, we can model the relative abundance of transcripts by modeling the probability of $Z$.

We use an inhomogeneous Markov chain to model the probability of the sequence of hidden variables, $Z = (Z_1, \ldots, Z_M)$. Specifically, the initial probability is given by

$$P(Z_1 = 1) = \pi,$$ \hspace{1cm} (5)

where $\pi$ represents the proportion of transcripts that contain the first segment. We model the transition probability from $Z_i$ to $Z_{i+1}$ for $i = 1, \ldots, M - 1$ as follows. If two consecutive segments $X_i$ and $X_{i+1}$ are separated by an exon start site $s_m$, then

$$P(Z_{i+1} = 1|Z_i = 0) = p_m$$ \hspace{1cm} (6)

$$P(Z_{i+1} = 1|Z_i = 1) = 1.$$ \hspace{1cm} (7)

If they are separated by an exon end site $e_m$, then

$$P(Z_{i+1} = 0|Z_i = 1) = q_m$$ \hspace{1cm} (8)

$$P(Z_{i+1} = 0|Z_i = 0) = 1.$$ \hspace{1cm} (9)

That is, if the current segment is transcribed ($Z_i = 1$), the splicing process ignores an exon start site (Equation 7), but it considers the potential usage of an exon end site $e_m$ and decides to use it, i.e. end the exon, with its usage probability $q_m$ (Equation 8). On the other hand, if the current segment is not transcribed ($Z_i = 0$), the splicing process ignores an exon end site (Equation 9), but it uses an exon start site $s_m$ with its usage probability $p_m$ (Equation 6). The parameters $P(Z_{i+1} = 0|Z_i = 0) = 1$ and $q = (q_1, \ldots, q_{M_2})$ represent probabilities of using the corresponding exon start and end sites, respectively, given that each site is considered for potential usage. Throughout the rest of this work, we refer to these usage probabilities simply as usages. Supplementary Table S1 shows the relative abundances defined by the proposed model for the three transcripts presented in Figure 3. A more detailed description is provided in Supplementary Sections S2.1–S2.3.

### 2.3 Parameter estimation and uncertainty quantification

We use an EM algorithm to compute the maximum likelihood estimates for the model parameters $\Theta = (\pi, p, q)$, that is $\Theta := \arg\max_{\Theta} P(R_1, \ldots, R_N|\Theta)$. The complete log likelihood in the EM algorithm involves $P(R_n, B_n = b, T_n = Z|\Theta)$ for $b \in \{1, \ldots, M\}$. The number of reads mapping to distinct combinations of segments in this example, only the start of the first exon and the end of the last exon are bounded by TSS and TES, respectively, the remaining exon start and end sites correspond to splice sites. (D) Estimate splice site usages using McSplicer. (E) Leverage splice site usages in various kinds of downstream analyses, such as the quantification of different types of alternative splicing events.
\{1, \ldots, l(Z)\} (Supplementary Section S2.4). By combining the generative model and the McSplicer model in the previous two sections, \(P(R_a, B_a = b, T_a = Z|\Theta)\) can be written as

\[
P(R_a|B_a = b, T_a = Z)P(B_a = b|T_a = Z)P(T_a = Z|\Theta) = \frac{1}{l(Z)} \sum_b l(Z|\omega_a(Z)) = \frac{1}{l(Z)} \sum_b l(Z|\omega_b(Z)) \tag{10}
\]

if \(R_a\) is identical to the sequence of length \(L\) starting at position \(b\) in transcript \(Z\). Otherwise, this probability is 0. The details of the application of the EM algorithm to the proposed model are provided in Supplementary Section S2.4. The EM algorithm uses several quantities that we compute using dynamic programming, see Supplementary Section S2.5. Also, all quantities required in our EM algorithm can be computed using only signature counts (Supplementary Section S2.4), so the input to McSplicer are the signature counts rather than individual reads.

We quantify the uncertainty of our estimator \(\Theta\) using bootstrapping. Specifically, let \(c = (c_j)_{j=1}^J\) represent the signature counts over \(J\) signatures defined for a given gene, where the total signature count equals the total read count in the gene, i.e. \(\sum_j c_j = N\). We draw \(B\) independent bootstrap samples, \(c_1, \ldots, c_B\), from a multinomial distribution:

\[
c_j \sim \text{multinomial}(\frac{c_j}{N}, \ldots, \frac{c_j}{N}) \tag{11}
\]

Then, we compute \(B\) bootstrap estimators, \(\Theta^1, \ldots, \Theta^B\), by applying our EM algorithm to each bootstrap sample and use them to approximate the sampling distribution of our estimator \(\Theta\). In this paper, we quantify the uncertainty of \(\Theta\) using a confidence interval computed from the approximated sampling distribution. Other types of uncertainty quantification could easily be obtained from the bootstrap estimators.

### 2.4 Simulated datasets and evaluation

We used Polyester (Frazee et al., 2015) to simulate reads from a human transcriptome with abundances estimated from a real RNA-seq experiment (GEO accession GSM3094221) using RSEM (Li and Dewey, 2011). Based on these ground truth expressions, we simulated datasets with varying sequencing depth commonly observed in practice, including 20 million, 30 million and 75 million reads of 100 bp length. Following the same strategy as Soneson et al. (2016), we randomly selected a set of 1000 genes with at least two expressed transcripts and sufficiently high ground truth expression (gene-level percent spliced in) above 500. Among splice sites for which parameters estimated by compared methods have the same meaning (comparable splice sites, introduced in Section 3), we exclude from the analysis constitutive ones with true usage 1 and splice sites that are not used by any of the expressed transcripts (usage 0). That is, only splice sites that are alternatively used by expressed transcripts are considered.

From the ground truth abundance of transcripts, we calculate the true usage of a splice site as the relative contribution of transcripts using a given splice site to the total expression of a gene (see Supplementary Section S2.6.3). We quantify the accuracy of splice site usages inferred by each method by using the Kullback-Leibler (KL) divergence, defined in Supplementary Section S2.6.4. All code and data necessary to reproduce the results of this simulation study are available at https://github.com/canzarlab/McSplicer.

### 3 Results

We assess the performance of McSplicer in comparison to existing state-of-the-art methods on both simulated and real RNA-seq datasets. Simulated data allow to compare estimates to a known ground truth of expressed transcripts and thus known quantities of alternative splicing events. On the other hand, simulated data cannot fully capture the complexity of datasets generated in real RNA-seq experiments. Note that exon start and end sites whose McSplicer estimators can correspond to splice sites but also to transcription start and end sites (see Section 2.2). In the following, however, we restrict the evaluation to the usage of splice sites since transcription start and end sites cannot be reliably estimated from short-read RNA-seq data alone.

We compare the performance of McSplicer to PSGInfer, SplAdder, MAJIQ and StringTie. In Supplementary Section S2.6.1 we provide details on software versions and command line arguments used. PSGInfer quantifies alternative splicing using a generative probabilistic model, an idea that also motivated the approach taken in McSplicer. SplAdder was used in a large-scale study (Kahles et al., 2018) to detect and quantify alternative splicing events in nearly 9000 tumor RNA-seq samples. In a comparative benchmark analysis performed in Kahles et al. (2016) it showed a better performance than competing methods JuncBase (Brooks et al., 2011), rMATS (Shen et al., 2014) and SpliceGrapher (Rogers et al., 2012), from which, of course, general superiority cannot be concluded (Denti et al., 2018). Compared to SplAdder, which is limited to the detection of simple types of splicing events, MAJIQ introduced a novel approach that additionally captures more complex transcript variations. MAJIQ was shown in a recent benchmark (Mehmood et al., 2020) to compare favorably to existing state-of-the-art methods and the authors demonstrated in Vasque-Garcia et al. (2018) that MAJIQ also outperforms LeafCutter and rMATS.

StringTie, on the other hand, assembles and quantifies full-length transcripts from RNA-seq but was not specifically designed for the quantification of splice site usage. Nevertheless, splice site usage can be inferred from the abundance of the assembled transcripts and we include this approach as a baseline in our benchmark: In all experiments, McSplicer uses StringTie to construct the ex-intron structure in steps (B) and (C) of the workflow (Fig. 2), which potentially contains novel splice sites. In contrast to the inference of splice site usage from expressed full-length transcripts, however, McSplicer estimates the usage of the same set of splice sites using the EM algorithm described in the previous section.

Each method, however, uses a different set of parameters to quantify alternative splicing events. PSGInfer infers the weights of its constructed splice graph edges. SplAdder quantifies four canonical types of splicing events using the widely used \textit{percent spliced in} (PSI) metric. PSI denotes the ratio between the number of reads supporting one outcome of the event (e.g. the inclusion of an exon) over the number of reads directly supporting either of the two alternative outcomes. Similarly, MAJIQ computes the \textit{percent selected index} (\(\Psi\)) for each splice junction involved in a local splicing variation (LSV), which denotes its fractional usage. To ensure a meaningful comparison of splice site usages in McSplicer to edge weights from PSGInfer, PSI from SplAdder and \(\Psi\) from MAJIQ, we only consider splice sites for which the meaning of these four quantities, if defined, coincide. These comparable splice sites are obtained from alternative splicing events between two expressed transcripts such that all remaining transcripts expressed by a gene consistently support one of the two possible outcomes of the event. Note that comparable splice sites are defined based on transcripts expressed in a given sample. We define comparable splice sites more formally in Supplementary Section S2.6.2. For comparable splice sites of simple events, the four different parameters, i.e. splice site usage, edge weights, PSI and \(\Psi\), equally reflect the relative abundance of transcripts expressed by a given gene that use the splice site, or equivalently contain the corresponding exon. Analogously, \(\Psi\), edge weight and splice site usage are equivalent for comparable splice sites of complex events. We will therefore consistently refer to these different parameters in the following as splice site usage. From StringTie assemblies of full-length transcripts, estimates of splice site usage can directly be obtained from the relative abundance of transcripts using a given splice site. For an illustrative example of comparable and non-comparable splice sites see Supplementary Figure S3.

#### 3.1 McSplicer more accurately infers splice site usage than competing methods

In this section, we assess the performance of McSplicer on RNA-seq datasets simulated as described in Section 2.4. All methods but...
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Fig. 4. Accuracy of McSplicer and competing methods in quantifying the usage of variable splice sites from 50 million simulated RNA-seq reads. For each method, only splice sites in events that the method reports and quantities are considered. SplAdder is limited to the quantification of simple AS events.

Fig. 5. McSplicer leverages all RNA-seq reads mapped to a gene to improve the accuracy of splice site usage estimates. On the dataset with 50 million simulated reads, McSplicer achieves lower KL divergence from true splice site usages when considering all reads mapped to a gene locus at once (blue) compared to using only reads that overlap any of the event’s exons (pink). ES denotes exon skipping, A3SS alternative 3' splice site, A5SS alternative 5' splice site, IR intron retention and CMPLX complex events.

PSGInfer were provided the same set of reads aligned using STAR (allowing mismatches and indels). PSGInfer only accepted unaligned reads which were internally mapped using Bowtie (Langmead et al., 2009). We distinguish splice sites by the type of event they are part of, including exon skipping, intron retention, alternative 3' and 5' splice sites, and complex events that cannot be assigned to one of the canonical types. The events are labeled by Astalavista (Foissac and Sammeth, 2007) through a pairwise structural comparison of all transcript species expressed in our ground truth transcriptome (see Supplementary Figs S1 and S4).

The number of variable splice sites (i.e. 0 < usage < 1) in our simulated dataset, and the number of comparable splice sites among them (~36%), with corresponding event types defined by Astalavista are listed in Supplementary Table S2. It also lists the total number of (comparable) splice sites per type reported by all four methods. While McSplicer will quantify the usage of all splice sites except those missed by StringTie in step (B) in Figure 2, competing methods report only events that satisfy an adjustable confidence threshold (SplAdder) or are considered reliable according to internal filters (MAJQ). As a result, both MAJQ’s and SplAdder’s accuracy is evaluated on a smaller, presumably more confidently comparable splice sites all reads that do not overlap any of the event’s exons (pink). ES denotes exon skipping, A3SS alternative 3' splice site, A5SS alternative 5' splice site, IR intron retention and CMPLX complex events.

3.2 McSplicer leverages all reads mapped to a gene

McSplicer makes use of all reads mapped to a given gene to simultaneously infer parameters in the McSplicer model, while other methods except PSGInfer typically use only reads that directly support their parameters. To quantify the contribution of the simultaneous inference in McSplicer to improve the accuracy of estimators, we estimate one splice site usage parameter at a time using only reads directly supporting the parameter. Similar to the calculation of the traditional PSI metric, we remove for each event with comparable splice sites all reads that do not overlap any of the event’s downstream exon (Supplementary Fig. S5). Here, we compare the performance to the latter one, which we observed to be slightly more accurate. The former is reported in Supplementary Figure S6.

Additional details and figures are provided in Supporting Information.
exons, and run and evaluate McSplicer on the resulting restricted instance as described in the previous section. Figure 5 confirms that McSplicer profits enormously from transcriptional evidence that lies outside of the local splicing event. Across all types of events, McSplicer estimates splice site usage less accurately when reads that do not overlap an event are removed.

3.3 McSplicer estimates agree with spike-in RNA variants

To evaluate the performance of McSplicer under the added complexity imposed by data derived from a real RNA-seq experiment, we used spike-in controls that were previously added to human monocyte-derived macrophages from five different donors (Hoss et al., 2019). The Spike-In RNA Variants (SIRV) (Paul et al., 2016) comprise 69 synthetic RNA molecules that were added in known relative concentrations before library preparation. Mimicking the complexity of 7 human model genes, between 6 and 18 artificial transcripts per gene vary in different types of alternative splicing, transcription start- and end-sites, or are transcribed from overlapping genes, or the antisense strand. The concentration ratios between different SIRV isoforms span a range of more than two orders of magnitude. For each donor sample, including artificial SIRV isoforms, Hoss et al. (2019) sequenced 200 million paired-end reads of 2 × 125 bp length. McSplicer considers both mates independently as input reads $K_r$ (see Section 2.3).

Leveraging the artificial reference genome (SIRVome) and the known relative mixing ratios of SIRV isoforms, we derive ground truth splice site usages (see Supplementary Section S2.6.3). Again, we obtain event labels from Astalavista, which comprise 26 variable splice sites in simple events and 12 in complex events. In this experiment, we do not restrict the evaluation to comparable splice sites but include all variable sites since competing methods report too few true events, too few to allow for a meaningful quantification of agreement between estimated and true PSI and $\psi$ values. We obtain similar results on the remaining four samples (Supplementary Fig. S13).

3.4 Quantifying the effect of cryptic splice site mutations in patients with autism spectrum disorder

In this section, we illustrate the utility of splice site usages computed by McSplicer in interpreting the potential complex effect of genetic variants on RNA splicing. In Jaganathan et al. (2019), the authors use a deep neural network to identify non-coding genetic variants that disrupt mRNA splicing. They identified a set of high-confidence de novo mutations predicted to disrupt splicing in individuals with intellectual disability and individuals with autism spectrum disorders (ASD). To validate them, the study included RNA-seq experiments (270–388 million 150 bp reads per sample) of peripheral blood-derived lymphoblastoid cell lines from 36 individuals with ASD. Based on the presence of reads spanning the corresponding splice junction, the authors validate 21 aberrant splicing events associated with the predicted de novo mutations. Each of the splicing events was uniquely observed in one individual.

In Jaganathan et al. (2019), the authors point out that computing the effects size of splicing mutations based on a pre-selected set of incident splice junctions likely underestimates the true effect size since, among other shortcomings, not all isoform changes are taken into account. In contrast, McSplicer’s model of splice site usage does not depend on an ad hoc selection of specific junctions or AS units but naturally captures simultaneous changes in expression of multiple isoforms expressed by a gene. We therefore utilized McSplicer to quantify the effect size of the validated de novo mutations on splice sites in ASD patients. We excluded 11 aberrant splicing events where only 1 or 2 spliced reads supported the novel splice site or junction. For each de novo mutation and the corresponding aberrant splicing event, we used McSplicer to estimate splice site usage and to compute 95% bootstrapping confidence intervals for the individual harboring the variant and a control individual with similar sequenc ing depth. For all 10 aberrant splicing events, we observe significantly different splice site usages (i.e. the two confidence intervals do not overlap) between mutated and control ASD individuals (Supplementary Table S3). Figure 7 provides three illustrative examples. For gene ENOPH1, McSplicer estimates a decrease in usage of the acceptor site directly affected by the variant, consistent with the increased skipping of the corresponding exon that can be observed in the Sashimi plot. In gene CORO1B, a novel donor site is used exclusively in the individual with the variant, identified and quantified with non-zero usage by McSplicer. For gene PCSK7, McSplicer

splice site usages as estimated by McSplicer to the true usages in one of the five samples (donor 5). A Spearman’s rank correlation coefficient of $\rho = 0.798$ indicates a good agreement between estimated and true usages. We obtain similar results on the remaining four samples (Supplementary Fig. S13). SplAdder and MAJIQ only report between 6 and 12 among all 38 true events, too few to allow for a meaningful quantification of agreement between estimated and true PSI and $\psi$ values. Supplementary Figures S14 and S15 show the corresponding scatter plots for PSI and $\psi$ values estimated by SplAdder and MAJIQ, respectively. PSGInter failed to run on all five donor samples for unknown reasons.

Fig. 6. McSplicer results on spike-in RNA variants (SIRV), donor sample 5. Ground truth splice site usages computed from known mixing ratios of SIRV isoforms are compared to usages estimated by McSplicer. Out of 38 variable splice sites, 26 belong to simple events and 12 belong to complex events. ES denotes exon skipping, A3SS alternative 3’ splice site, A5SS alternative 5’ splice site, IR intron retention and CMPLX complex events

Fig. 7. McSplicer splice site usage estimates and 95% bootstrapping confidence intervals for three disrupted splicing events reported in ASD patients versus control individuals. Variant locations are indicated by black vertical lines. Each plot illustrates the gene structure around the event with the precise genomic window specified on top, the read coverage and the junction read count. The Sashimi plots shown here are created using the ggashimi tool (Garrido-Martín et al., 2018)
estimates a decrease in usage of the affected donor sites, consistent with the retention of the downstream intron.

4 Conclusion

We have introduced McSplicer, a novel method that estimates the usage of exon start and end sites, and in particular the usage of splice sites across expressed transcripts. Rather than attempting to reconstruct expressed transcripts, McSplicer is based on a simplified probabilistic splicing model that has generated the set of expressed transcripts. It is not restricted to a pre-defined class of alternative splicing events or units but our probabilistic model is able to describe arbitrarily complex types of splicing patterns based on few, easy to interpret, parameters. We estimate these parameters, i.e. splice site usages, using all read data at once and demonstrate in simulation experiments that this yields more accurate estimates compared to other methods that use only reads directly supporting their parameters. Through its integration with transcript assembly methods such as StringTie, McSplicer quantifies the usage of annotated as well as novel splice sites.

Our model for relative transcript abundance assumes the Markovian property across indicators (Z) for whether a segment is transcribed. This assumption allows for an efficient algorithm to estimate parameters of the model, but it potentially limits the ability of our model to model longer range dependencies such as between the recognition of 5’ and 3’ splice sites or between the removal of introns within transcripts. If true dependencies are longer than our model can describe, the individual estimators for splice site usages may still be accurate, but we expect transcript frequencies implied by our model to be less accurate (LeGault and Dewey, 2013). One way to model longer range dependencies is to use higher order Markov chains as long as the data provide sufficient information to estimate these dependencies.

The splice site usages computed by McSplicer can be leveraged in various types of downstream analyses, such as the statistical comparison of splice site usage between different conditions (Li et al., 2018), the quantification of various types of splice events, the identification of subgroups of samples that show similar splicing patterns [i.e. unsupervised clustering (Ntranos et al., 2016)], or the discrimination between alternatively spliced and constitutive exons (Patrick et al., 2013).

We have used McSplicer to quantify the effect size of splicing mutations in ASD patients. In this context, splice site usage as computed by McSplicer can be considered analogous to the ‘strength’ of a splice site predicted by methods such as SplicePort (Dogan et al., 2007) from sequence-based features. Point mutations in the consensus splice site sequence can affect the strength of a splice site and result in the skipping of the exon or the activation of cryptic splice sites. In fact, a single nucleotide substitution might produce multiple (erroneous) splicing isoforms at the same time, as has been observed, for example, for specific mutations in patients with cystic fibrosis (3 isoforms) (Ramalho et al., 2003) and X-linked spondyloepiphyseal dysplasia tarda (7 isoforms) (Xiong et al., 2009). McSplicer does not attempt to reconstruct every single aberrant isoform, but similar to a weakening (strengthening) of a splice site as predicted from sequence alterations by, e.g. the Shapiro splice site probability score (Shapiro and Senapathy, 1987), the effect of a mutation will be reflected in a reduced or increased usage of the corresponding splice site estimated from RNA-seq reads.

The procedure we applied to compute the effect size of splicing mutations in our analysis of ASD patients data does not use the full data from multiple individuals and fails to consider variability among individuals, possibly leading to an increased number of false positives. Methods that model differences in splice site usages between individuals from multiple groups and exploit the variability among them should perform better in estimating effect size and quantifying their uncertainty.

Acknowledgements

The authors thank Matthew Stephens for invaluable discussions on the proposed model and Zhen Zuo for help with testing the method on other datasets. They thank the members of H. Shim, S. Canzar and T. Speed groups for helpful comments.

Funding

This work was supported by the Purdue startup fund. L.A. was supported by a Deutsche Forschungsgemeinschaft fellowship through the Graduate School of Quantitative Biosciences Munich.

Conflict of Interest: none declared.

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**Title:**
McSplicer: a probabilistic model for estimating splice site usage from RNA-seq data

**Date:**
2021-01-30

**Citation:**
Alqassem, I., Sonthalia, Y., Klitzke-Feser, E., Shim, H. & Canzar, S. (2021). McSplicer: a probabilistic model for estimating splice site usage from RNA-seq data. BIOINFORMATICS, 37 (14), pp.2004-2011. https://doi.org/10.1093/bioinformatics/btab050.

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