# IPSA: Integrative Pipeline for Splicing Analysis

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# NOTE THIS DOCUMENT IS SUBSTANTIALLY OUT OF DATE

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# 1 Synopsis

This document contains the description of the integrative splicing analysis pipeline. The integrative pipeline consists of several sub-pipelines, of which we primarily discuss SJPIPE and TXPIPE.

SJPIPE implements the quantification of splicing events by directly analyzing split alignments in BAM files (Figure 1). The advantage of SJPIPE is that it quantifies annotated as well as novel splice junctions and that there is no annotation-related interference between independent splicing events. An obvious disadvantage, however, is that only a part of the sequencing information is being used, one that is related to local alignments at splice junctions, while the reads that are fully contained in exons or in introns are ignored.

SJPIPE pipeline starts with a BAM file, which is processed by *sjcount* utility and converted to a tab-delimited output of *multi-split counts with offsets* (see sjcount documentation for more detail). Additionally, *sjcount* produces counts of the continuous reads that overlap splice junctions. These outputs are aggregated, matched against genome and annotation, and passed to strand disambiguation. The resulting stranded counts are subject to irreproducibility assessment (npIDR) and subsequent filtering based on IDR, entropy, and staggered alignments. The workflow is unversal for many types of data, i.e., for stranded and unstranded data, for data with and without bioreplicates, etc.

The TXPIPE pipeline assesses the annotated splicing events by analyzing the transcript quantification data produced by some deconvolution program. The advantage of this approach is that it takes into account all sequencing data, i.e., not only local alignments at splice junctions, but also alignments of reads in exons and in introns. This results in a larger effective sample size compared to that in SJPIPE. A disadvantage, however, is that unannotated splicing events or dependent splicing events may distort the global splicing pattern significantly and, therefore, there is no guarantee that the inclusion rate of each given exon actually reflects splicing events that occur with the given exon (see examples below).

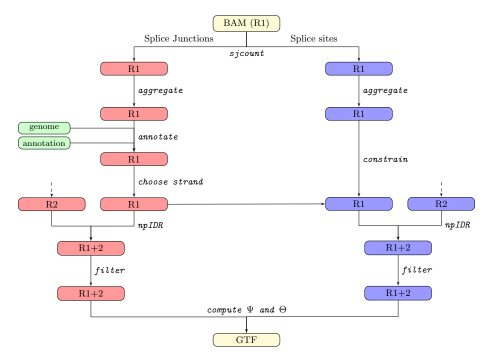


Figure 1: The SJPIPE pipeline.

# 2 Detailed description of the components

## 2.1 Perl libraries

Most of the pipeline is based standard Perl modules and a custom module called utils.pm. All Perl scripts are located in Perl/ subdirectory. Since paths are not modified, in order to call a Perl script, one should do it from the main folder as perl Perl/script.pl; otherwise utils.pm will not be found.

#### 2.2 RNA dashboard index file

An RNA dashboard index file is a two-column tab-delimited text file whose first column contains the file name or a URL, and the second column contains a list of attributes of the form attribute1=value1; attribute2=value2; etc.

## 2.3 Pipeline generator

The make.pl utility takes as an input the RNA dashboard index file and outputs to the standard output a GNU makefile that will be executed in order to to run

the pipeline.

```
Perl/make.pl -dir <dirname> -param <params> -by <attribute> -margin
          <length> -annot <gtf> -genome <name> -merge <name>
Inputs: -annot ..., the annotation (gtf), obligatory
-deltaSS ..., distance threshold for splice sites, default=0
-block ..., the blocking field for mastertable
-dir ..., the output directory, obligatory
-entropy ..., entropy lower threshold, default=1.5
-genome ..., the genome (without .dbx or .idx), obligatory
-group ..., the grouping field for IDR, obligatory
-idr ..., IDR upper threshold, default=0.1
-margin ..., margin for aggregate, default=5
-merge ..., the name of the output to merge in case if blocks are missing
-mincount ..., min number of counts for the denominator, default=10
-param ..., parameters passed to sjcount
-repository ..., the repository subdirectory for bam files
-status ..., annotation status lower threshold, default=0
        STDIN: an RNA dashboard index file
Output: STDOUT: a GNU makefile
       make -f file.mk all
```

For example, the command

Perl/make.pl -repository input/ -dir output/ -group idrGroup -param '-read1 0' -annot hg19v18.gff -genome homSap19 -merge pooled < example.dat > example.mk

creates a file called 'example.mk' based on the input from 'example.dat', where all the intermediate steps will be stored in 'output/' with the annotation file hg19v18.gtf in the current directory, and with homSap19.idx and homSap19.dbx both being readable genome files.

## 2.4 Pre-processing

## 2.4.1 Annotation pre-processing

Genomic annotation files often come in GTF format and contain many feature types simultaneously. Often these are large files and it takes a lot of time just to read them. Sometimes they may be incomplete (e.g. contain exons, but not introns) and not fully correspond to GFF3 standarts.

The *transcript\_elements.pl* utility reads a gtf, extracts only exons, and outputs for each exon the list of transcripts to which this exon belongs. It

outputs also a gtf which contains (1) exons and (2) introns inferred from the input exon information, but the output file is shorter. For example, in the input gtf were

```
chr2L FlyBase exon 100 200 . + . gene_id "8"; transcript_id "1"; chr2L FlyBase exon 300 400 . + . gene_id "8"; transcript_id "1"; chr2L FlyBase exon 500 600 . + . gene_id "8"; transcript_id "1"; chr2L FlyBase exon 100 200 . + . gene_id "8"; transcript_id "2"; chr2L FlyBase exon 500 600 . + . gene_id "8"; transcript_id "2"; ... ... ... ... ... ... ...
```

then the processed form would be

```
chr2L SJPIPE exon
                      100
                            200
                                            transcript_id "1,2";
                                            transcript_id "1";
chr2L SJPIPE intron
                      200
                            300
chr2L SJPIPE intron
                      200
                            500
                                            transcript_id "2";
chr2L SJPIPE exon
                      300
                            400
                                            transcript_id "1";
chr2L SJPIPE intron
                      400
                            500
                                            transcript_id "1";
                                            transcript_id "1,2";
chr2L SJPIPE exon
                      500
                            600
```

Formally speaking, the script groups transcript id for each exon and applies the function join(",",uniq(sort(...))) to the sorted array of transcript ids. Similarly, the script can group any other feature and apply any other function to the array, as well as it can put any value for the source field, e.g.

```
Perl/transcript_elements.pl -features transcript_id,source,position:avg
```

will return uniq(transcript\_id), uniq(source), and avg(position) in the column 9 of GTF. Note that transcripts and introns are not shown in the input gtf. The information contained in the 'transcript\_id' field will be used only in TX-PIPE pipeline to compute splicing indices from transcript quantification (see section 2.6).

**NB** In what follows, the processed annotation has to be sorted. Therefore it is necessary to pipe it with the sort step as follows.

```
Perl/transcript_elements.pl - < input.gtf | sort -k1,1 -k4,5n > output.gff
```

#### 2.4.2 Genome pre-processing

The following two utilities, *transf* and *getsegm*, which belong to the *maptools* package, are used to pre-process genomes in a more compact and readable form.

maptools can be obtained from github. The scripts in maptools/bin shall be made accessible by declaring the path to that directory.

The use of transf utility is as follows

```
transf -dir genome_directory/any.fa -dbx output.dbx -idx output.idx
```

It takes all the files in <code>genome\_directory/</code> and creates two output files, <code>output.dbx</code> and <code>output.idx</code>; the former storing the data and the latter storing the index table tot hat data. The format is similar to 2bit.

The getsegm doesn't have to run on its own. Instead, it is used in annotate.pl to get genomic nucleotides.

## 2.5 SJPIPE pipeline

#### 2.5.1 Counting splice junctions and reads that overlap splice sites

The *sjcount* utility takes a BAM file and counts the number of split reads supporting splice junctions (SJ) and continuous reads that overlap splice sites (SS). Splice sites are defined by the splice junctions that are present in the alignments. Version v3.1 also counts the abundance of split reads that span multiple junctions.

The utility returns the number of counts for each combination of chromosome, begin, end, strand, and offset, where offset is defined to be the position (within the short read sequence) of the last nucleotide preceding the splice junction. For the exact definitions of SJ, SS, offset, and examples see the help page of *sjcount* at github.

```
sjcount -bam <file> -ssj <file> -ssc <file> ...

Input: a sorted BAM file with a header

Options:
-read1 0/1, reverse complement read1 no/yes (default=1)
-read2 0/1, reverse complement read2 no/yes (default=0)
-nbins number of overhang bins, (default=1)
-maxnh, the max value of the NH tag, (default=none)
-lim nreads stop after nreads, (default=no limit)
-unstranded, force strand to be '.'
-continuous, no mismatches when overlapping splice boundaries
-gz, gzip output ('.gz' extension will *NOT* be added to output file name)
-quiet, suppress verbose output
```

Output: -ssj: Splice Junction counts, tab-delimited (default=stdout)

```
Columns are: path, number_of_splits, offset, count
-ssc: Splice boundary counts, tab-delimited (default=none)
Columns are: path, number_of_splits, offset, count
```

NB: the coordinates are 1-based

#### 2.5.2 Aggregating SJ counts over offsets

The aggregate.pl utility takes the output of sjcount on STDIN and performs aggregation by the 5th column (offset) using three different aggregation functions (see examples below). It outputs a TSV file with three extra columns being (5) total count, (6) staggered read count, (7) entropy. The output is sent to stdout.

```
Perl/aggregate.pl
Input: TSV (ssj or ssc) on STDIN
Output: TSV on STDOUT
Options:

-margin ..., the margin for offset, default=0
-maxintron ..., max intron length, default=0
-minintron ..., min intron length, default=0
-readLength ..., the read length, default=0
Columns in the output are: chr, begin, end, strand, count, staggered, entropy
```

It is possible to exclude short reads with small overhangs on either side by using -margin and -readLength parameters. This is particularly important when such a margin was imposed during the mapping step, but in order to be comparable when counting reads that overlap splice sites one should use the same restriction. Also, it is possible to exclude SJs that are too long or too short (-minintron/-maxintron).

The aggregation functions are applied to the sample  $\{x_k\}$  of counts for each combination of chromosome, begin, end, and strand vs. the offset value k. The aggregation function therefore has the general form  $f(x_1, \ldots, x_n)$ .

When  $f(x_1, ..., x_n) = x_1 + \cdots + x_n$ , the result coincides with the collapsed (total) number of counts, i.e., as if offsets were ignored. For for  $f(x_1, ..., x_n) = \theta(x_1) + \cdots + \theta(x_n)$ , where  $\theta(x) = 1$  for x > 0 and  $\theta(x) = 0$  for  $x \le 0$ , the result

is the number of staggered read counts. The function

$$f(x_1, \dots, x_n) = \log_2(\sum_{i=1}^n x_i) - \frac{\sum_{i=1}^n x_i \log_2(x_i)}{\sum_{i=1}^n x_i}$$

gives the entropy of the distribution, which can be used later to filter out non-uniform distibution of read counts. For example, if the input were

chr1	50	90	+	20	5
chr1	100	200	-	10	25
chr1	100	200	-	11	12
chr1	100	200	-	15	4
chr1	100	200	+	10	1
chr1	100	300	+	11	12

the output would have been

					• • •	
chr1	100	200	-	41	3	1.28

where 1.28 is the entropy of the distribution.

# 2.5.3 Checking the annotation status of a SJ and retrieving splice site nucleotides

The annotate.pl takes an aggregated TSV file (the output of aggregate.pl), the genomic annotation, and the genome, and outputs a TSV with two more columns: (8) the annotation status and (9) splice sites. The output is sent to stdout.

#### Perl/annotate.pl

The annotation file is a simplified, processed form of the standard annotation gtf. It can be obtained by the *transcript\_elements.pl* utility (see section 2.3.1).

The genome consists of two compressed files, \*.dbx and \*.idx, which can be obtained from the genomic fasta sequence by using transf utility of the maptools package.

In previous versions the annotation status was defined numerically as follows:

- (0) None of the splice sites of the given SJ is annotated;
- (1) One of the splice sites of the given SJ is annotated, and the other is not;
- (2) Both splice sites of the given SJ are annotated but the intron between them is not;
- (3) Both splice sites of the given SJ are annotated, and so is the intron between them.

In the current version the annotation status is defined for each splice site as

- (0) not within deltaSS nucleotides from an annotated SS;
- (1) within deltaSS nucleotides from an annotated SS;
- (2) annotated.

The status of a splice junction is defined according to the table (rows – donor site; columns – acceptor site)

	0	0 1			
	not annot	within deltaSS	annotated		
0	0	0.5	1		
1	0.5	0.75	1.5		
2	1	1.5	2 or 3		

The splice site nucleotides are the four intronic nucleotides, two flanking ones from each end, such as GTAG or ATAC. Since for this field and for the annotation status strand has to be defined, two lines are produced in the case of unstranded data (one for each strand). For instance, if the input were

```
chr1 100 200 + 41 3 1.28
chr1 100 200 - 41 3 1.28
```

and there were, indeed, an annotated junction at (chr1, 100, 200, -) with GTAG, then the output would have been

```
. . .
                200
                               41
                                      3
                                                      0
                                                             CTAC
chr1
        100
                                              1.28
chr1
        100
                200
                               41
                                      3
                                              1.28
                                                      3
                                                              GTAG
```

Note that sequence retriever uses the <code>getsegm</code> program of the <code>maptools</code> package, so maptools has to be installed and path has to be added.

#### 2.5.4 Strand choice

At this step, a unique value of strand is chosen for each SJ. This is done by *choose\_strand.pl* utility.

For each combination of chromosome, begin, and end, the strand with greater annotation status (see section 2.4.3) is chosen. In case of a tie (usually 0 on both strands), the strand is chosen based on the "largest" splice site nucleotides in terms of lexicographic order (TTTT>GTAG>...). There will be an option to choose a custom order of trustable splice site sequences (e.g., GTAG>ATAC>others).

For instance, if the input were

• • •			• • •			• • •		
chr1	100	200	+	41	3	1.28	0	CTAC
chr1	100	200	-	41	3	1.28	3	GTAG
chr1	150	200	+	21	2	1.01	0	GTAG
chr1	150	200	-	21	2	1.01	0	CTAC

the output would have been

```
1.28
chr1
        100
               200
                             41
                                    3
                                                   3
                                                          GTAG
chr1
        150
               200
                             21
                                    2
                                           1.01
                                                   0
                                                          GTAG
```

#### 2.5.5 Constraining splice site counts

Since now a unique value of strand is chosen for each SJ, the counts of reads overlapping splice sites have to be constrained to a smaller set of splice sites. This is done by <code>constrain\_ssc.pl</code> utility.

Perl/constrain\_ssc.pl

 ${\tt Input:} \quad {\tt TSV} \ {\tt ssc} \ ({\tt reads} \ {\tt overlapping} \ {\tt splice} \ {\tt sites}) \ {\tt on} \ {\tt STDIN}$ 

-ssj = TSV file to constrain to, obligatory

Output: TSV on STDOUT

Here -ssj is the splice junction file after strand choice was made, -ssc is the output of <code>sjcount</code>. The output of <code>constrain\_ssc.pl</code> is sent to <code>stdout</code>. If the ssc input is unstranded, then the strand of a splice site is taken from ssj, where the strand is already defined. In some cases it will lead to two lines being produced in the case of unstranded data (one for each strand). For example, if the ssj input were

chr1	100	200	536	-	41	3	1.28	3	GTAG
chr1	150	200	439	+	21	2	1.01	0	GTAG

then (chr1, 200, +) and (chr1, 200, -) are both valid splice sites and the corresponding ssc counts will be reported for each of the two strands.

### 2.6 Ascertainment of reproducibility (IDR)

In this step the number of counts from (generally, as many as possible) bioreplicates are assessed for irreproducibility. This step is done by *idr4sj.r* 

```
Rscript R/idr4sj.r inp1.tsv [inp2.tsv] ... [inpN.tsv] output.tsv
```

where inp1,2,...N the bioreplicates and output is the file is the last in the command line. The output contains one extra column (10) equal to IDR score. Columns 7, 8, and 9 are summed (not averaged!) between bioreplicates. In case if only one input file, the IDR score is set to 0. Currently, in case of more than two bioreplicates, only the first two files will be considered (others ignored).

### 2.6.1 Filtering

There is no specific routine for filtering because it can be done by *awk* by requiring column 7 (entropy) to be greater than threshold (usually, 3 bits) and the column 10 (IDR) be not greater than 0.1.

#### 2.6.2 Calculation of splicing indices

As soon as the SJ (and splice site) counts were assessed for reproducibility and filtered, the next step is to compute the inclusion and processing rates by zeta.pl utility. The inclusion and processing rates can be defined for exons and for introns and exist under different names [1]. Since, by definition, splice junctions know nothing about the set of exons that one might want to assess, the global exon inclusion and processing rates are computed for for a given set of annotated exons, as specified in the annotation file. In contrast, the inclusion and processing rates of SJ are computed for all splice junctions that remain intact after filtering, but also the annotated SJ are also assessed and reported.

Here, whenever a ratio is calculated, we usually relate inclusion quantity to the sum of inclusion and exclusion quantities. The latter, however, can be a small integer number and, therefore, a threshold is needed to cut off estimates with large standard errors. This threshold is -mincount. There is also an option to enforce strandless computation, but it will be deprecated in future versions.

The procedure of zeta.pl is to read and to index all SJs and then for each splice site to create a list of exons which start or end at the given splice site. Then, reading sequentially the count file, the program increments the counters for exon inclusion, exon exclusion, and also for SJ usage. The output is a GFF with the corresponding features, e.g.

where psi and cosi are exon percent-spliced-in and completeness of splicing rates; psi5, psi3, cosi5, cosi3 are the respective percent-spliced-in and completeness of splicing indices of an intron, measured from the 5'-end and from the 3'-end. The rest of the parameters are counts.

## 2.7 TXPIPE pipeline

# 2.7.1 Computation of splicing indices from transcript quantification data

The tx.pl utility takes the transcript quantification data (gtf) and the preprocessed genomic annotation (the output of  $transcript\_elements.pl$ ). For each exon in the annotation it returns the sum of abundances of transcripts that contain the given exon (as an exon, i.e., there is a line in the gtf file saying that the exon belongs to the transcript) as a fraction of the sum of abundances of transcripts which cover the given exon (i.e., the transcript starts upstream and ends downstream of the exon). Note that this definition doesn't require gene id at all.

The transcript abundance is defined in the gtf field 9 as 'RPKM' attribute or as the mean of the two bioreplicates defined by 'RPKM1' and 'RPKM2' attributes.

#### 2.8 Master tables and endpoints

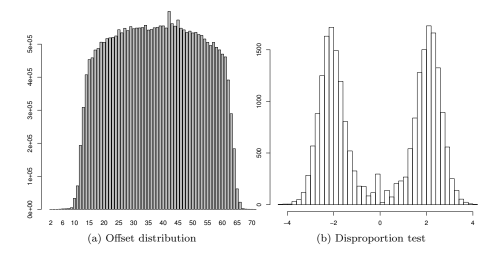
The  $merge\_gff.pl$  utility is formally not a part of any pipeline but it can be used to merge the content of a number of gff/gtf files into a square (R-readable) matrix.

The program reads input files specified in the '-i' parameter (could be many) one by one and selects features specified in '-o' parameter. For instance,

```
Perl/merge_gff.pl -i cell_line1.gtf HELAS3 -i cell_line2.gtf NHEK -o psi result_psi.tsv -o cosi result_cosi.tsv
```

will generate two files, result\_psi.tsv and result\_cosi.tsv, each containing a square table, e.g.,

```
HELAS3 NHEK
chr1_100_200_+ 0.52 0.75
chr1_300_400_+ 0.00 1.00
... ...
```



It can be applied to any feature that was specified in the gtf input in cell\_line1.gtf and cell\_line2.gtf.

# 3 Quality control functions

## 3.1 Distribution of offsets

The following utility plot the distribution of offset frequencies, i.e., how frequently each offset value was seen. This distribution may be helpful in guessing the correct margin value because some mappers have intrinsic thresholds that may be different for reads overlapping SJ and SB.

Rscript offset.r <file.tsv> <file.pdf>

An example such diagram is shown in Figure 2a.

## 3.2 Strand disproportion

Another useful test is the distribution of  $\log(c_+) - \log(c_-)$ , where  $c_+$  and  $c_-$  are the number of counts on the plus and the minus strand, respectively. SJ with  $c_+ = 0$  or  $c_- = 0$  are excluded. If the data is stranded and read1/read2 flags were set up correctly, then the distribution of  $\log(c_+) - \log(c_-)$  shall be bimodal as shown in Figure 2b, reflecting the fact that one strand has much more split reads than the other.

Rscript disproportion.r <file.tsv> <file.pdf>

## 3.3 Summary stats on annotation status and splice sites

As soon as splice junction counts are computed, it makes sense to ask what proportion of splice junctions is annotated and what is the distribution of frequencies of splice site nucleotides. This is done by sjstat.r utility.

R/sjstat.r <file.tsv> > <file.log>

# References

[1] D. D. Pervouchine, D. G. Knowles, and R. Guigo. Intron-centric estimation of alternative splicing from RNA-seq data. *Bioinformatics*, 29(2):273–274, Jan 2013.