SpliceLauncher

SpliceLauncher is a pipeline tool to study the alternative splicing. The pipeline works in three steps: * Get a read count matrix from fastq files, by a dedicated RNAseq pipeline (A step in diagram below). * Generate data files used hereafter (B step in diagram below) * Run SpliceLauncher from a read count matrix (C step and furthermore in diagram below).

SpliceLauncher

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Repository contents

- dataTest: example of input files
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Prerequisites to install SpliceLauncher

The SpliceLauncher pipeline needs to install the following tools and R librairies:

- STAR (v2.6 or later)
- samtools (v1.3 or later)
- BEDtools (v2.17 or later)
- R with WriteXLS and Cairo packages
- Perl

STAR

Following instruction were from the STAR manual

Get the g++ compiler for linux

```
sudo apt-get update
sudo apt-get install g++
sudo apt-get install make
```

Download the latest release and uncompress it

```
# Get latest STAR source
wget https://github.com/alexdobin/STAR/archive/2.7.0c.tar.gz
tar -xzf 2.7.0c.tar.gz
cd STAR-2.7.0c

# Alternatively, get STAR source using git
git clone https://github.com/alexdobin/STAR.git
```

Compile under Linux

```
# Compile

cd STAR/source

make STAR
```

Samtools

Download the samtools package at: https://github.com/samtools/samtools/releases/latest

Configure samtools for linux:

```
cd samtools-1.x
./configure --prefix=/where/to/install
make
make install
```

For more information, please see the samtools manual

BEDtools

Installation of BEDtools for linux:

```
wget https://github.com/arq5x/bedtools2/releases/download/v2.25.0/bedtools-2.25.0.tar.gz
tar -zxvf bedtools-2.25.0.tar.gz
cd bedtools2
make
```

For more information, please see the BED tools tutorial

Install R libraries

Open the R console:

```
install.packages("WriteXLS")
install.packages("Cairo")
```

Installing SpliceLauncher

Download the latest release from of SpliceLauncher source using git

```
git clone https://github.com/raphaelleman/SpliceLauncher
cd ./SpliceLauncher
```

Download the reference files

The reference files are the genome (Fasta) and the corresponding annotation file (GFF3):

- 1. Reference genome in fasta format
- 2. The annotation file in GFF v3 format

Steps: 1. Download Fasta genome: from RefSeq FTP server or from Gencode.

For example, human hg19 genome file from RefSeq: Bash #the ftp URL depends on your assembly genome choice wget ftp://ftp.ncbi.nlm.nih.gov/refseq/H_sapiens/annotation/GRCh37_latest/refseq_identifiers/GRCh37_latest_genomic.fna.gz gunzip ./GRCh37_latest_genomic.fna.gz

2. Download the GFF annotation file, either from Ref Seq FTP server or from Gencode.

For example, human hg19 annotation file from RefSeq:

```
wget ftp://ftp.ncbi.nlm.nih.gov/refseq/H_sapiens/annotation/GRCh37_latest/refseq_identifiers/GRCh37_latest_genomic
gunzip ./GRCh37_latest_genomic.gff.gz
head ./GRCh37 latest genomic.gff
##gff-version 3
#!gff-spec-version 1.21
#!processor NCBI annotwriter
#!genome-build GRCh37.p13
#!genome-build-accession NCBI_Assembly:GCF_000001405.25
#!annotation-date
#!annotation-source
##sequence-region NC 000001.10 1 249250621
##species https://www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id=9606
                                                         ID=id0;Dbxref=taxon:9606;Name=1;chromosome=1;gbkey=Src
NC_000001.10 RefSeq region 1 249250621 . + .
NC_000001.10 BestRefSeq gene
                                  11874 14409
                                                             ID=gene0;Dbxref=GeneID:100287102,HGNC:HGNC:37102;N
NC_000001.10
              BestRefSeq transcript 11874 14409 . +
                                                                 ID=rna0;Parent=gene0;Dbxref=GeneID:100287102,6
NC_000001.10
                                  11874
                                                             ID=id1;Parent=rna0;Dbxref=GeneID:100287102,Genbank
               BestRefSeq exon
                                          12227
                                                                                                            Þ.
```

Configure SpliceLauncher with INSTALL mode

SpliceLauncher is provide with a config.cfg file. This last contains the path for softwares and files used by SpliceLauncher. The mode INSTALL of SpliceLauncher updates this config.cfg file. The INSTALL mode uses the GFF (v3) file and the FASTA genome to extract all necessary information and to generate the STAR genome indexes. These information are storage in a BED file that contains the exon coordinates, in a sjdb file that contains the intron coordinates and a text file that contains the details of transcript structures. You need to define where these files will be saved by the -0, --output argument

Use INSTALL mode of SpliceLauncher:

```
cd /path/to/SpliceLauncher/
mkdir ./refSpliceLauncher # Here this folder will contain the reference files used by SpliceLauncher
bash ./SpliceLauncher.sh --runMode INSTALL \
        -0 ./refSpliceLauncher \
        --STAR /path/to/STAR \
        --samtools /path/to/samtools \
        --bedtools /path/to/bedtools \
        --gff /path/to/gff \
        --fasta /path/to/fasta
```

Running the SpliceLauncher tests

The example files are provided indataTest, with the example data provided in single end RNAseq (1x75pb) on BRCA1 and BRCA2 transcripts:

```
Bash cd /path/to/SpliceLauncher bash ./SpliceLauncher.sh --runMode Align,Count,SpliceLauncher -F ./dataTest/fastq/ -0 ./testSpliceLauncher/
```

After running, the BAM files from alignment are in a Bam folder, the count files are in getClosestExons and the results of SpliceLauncher analysis are in testSpliceLauncher_result.

The final results are displayed in the filetestSpliceLauncher_outputR.xlsx, this last is in testSpliceLauncher_result folder. The scheme of this

file is:

Column names	Example	Description
Conca	chr13_32915333_32920963	The junction id (chr_start_end)
chr	chr13	Chromosome number
start	32915333	Genomic coordinate of start junction End if on reverse strand
end	32920963	Genomic coordinate of end junction Start if on reverse strand
strand	+	Strand of the junction ('+': forward; '-':reverse)
Strand_transcript	forward	Strand of transcript
NM	NM_000059	The transcript id according RefSeq nomenclature
Gene	BRCA2	Gene symbol
Sample	2250	Read count
P_Sample	15.25659623	% of relative expression
event_type	SkipEx	The nature of junction: Physio: Natural junction SkipEx: Exon skipping 5AS: Donor splice site shift 3AS: Acceptor splice site shift NoData: Unannotated juntion
AnnotJuncs	Δ12	The junction names
cStart	c.6841	Transcriptomic start coordinate of the junction
cEnd	c.6938	Transcriptomic end coordinate of the junction
mean_percent	12.60242	Average in % of relative expression across samples
read_mean	2683.769231	Average of read count across samples
nbSamp	11	Number of time that the junction has been seen in samples
DistribAjust	-	The Distribution of junction expression (Gamma/N.binom)
Significative	NO	If a sample shown an abnormal expression of the junction

SpliceLauncher options

-runMode INSTALL, Align, Count, SpliceLauncher * The runMode defines the steps of analysis with: * INSTALL: Updates the config.cfg file for SpliceLauncher pipeline * Align: Generates BAM files from the FASTQ files * Count: Generates the matrix read count from the BAM files * SpliceLauncher: Generates final output from the matrix read count

Option for INSTALL mode

- -C, -config /path/to/configuration file/ * Path to the config.cfg file,only if you want to use your own config file
- -O, -output /path/to/output/ * Directory to save the reference files (BED, sjdb, txt) and the indexed genome
- -STAR /path/to/STAR * Path to the STAR executable
- $\hbox{\bf --samtools /path/to/samtools * Path to the samtools executable}$
- ${\color{red}\textbf{-bedtools}\,/\text{path/to/bedtools}} \ {\color{blue}\textbf{*}\,\text{Path}\,\text{to}\,\text{the}\,\text{bedtools}\,\text{executable}}$

- -gff /path/to/gff file * Path to the GFF file (v3)
- -fasta /path/to/fasta * Path to the genome fasta file
- -t, -threads N * Nb threads used to index the STAR genome

Option for Align mode

- -F, -fastq /path/to/fastq/ * Repository of the FASTQ files
- -O, -output /path/to/output/ * Repository of the output files
- -p * Processes to paired-end analysis
- -t, -threads N * Nb threads used for the alignment
- -g, -genome /path/to/genome * Path to the genome directory, only if you to use a genome directory different of the genome defined in config.cfg file
- -STAR /path/to/STAR * Path to the STAR executable, only if you to use a STAR software different of the STAR defined in config.cfg file
- -samtools /path/to/samtools * Path to the samtools executable, only if you to use a samtools software different of the samtools defined in config.cfg file

Option for Count mode

- -B, -bam /path/to/BAM files * Repository of the BAM folder
- -O, -output /path/to/output/ * Repository of the output files
- -samtools /path/to/samtools * Path to the samtools executable, only if you to use a samtools software different of the samtools defined in config.cfg file
- -bedtools\t/path/to/bedtools * Path to the bedtools executable,only if you to use a bedtools software different of the bedtools defined in config.cfg file
- -b, -BEDannot /path/to/your_annotation_file.bed * Path to exon coordinates file (in BED format),only if you to use exon coordinates different of the coordinates defined in config.cfg file

Option for SpliceLauncher mode

- -I, -input/path/to/inputFile * Read count matrix (.txt)
- -O, -output /path/to/output/ * Directory to save the results
- -TranscriptList /path/to/transcriptList.txt * Set the list of transcripts to use as reference
- -txtOut * Print main output in text instead of xls
- -bedOut * Get the output in BED format
- -Graphics * Display graphics of alternative junctions (Warnings: increase the runtime)
- -n, -NbIntervals 10 * Nb interval of Neg Binom (Integer)
- -SampleNames name1|name2|name3 * Sample names, '|'-separated, by default use the sample file names

If list of transcripts (--TranscriptList): -removeOther * Remove the genes with unselected transcripts to improve runtime

If graphics (-g, --Graphics): --threshold 1 * Threshold to shown junctions (%)

-R, -RefSeqAnnot /path/to/RefSpliceLauncher.txt * Transcript information file,only if you to use a transcript information file different of file defined in config.cfg file

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