# **TAPIS Documentation**

Release 1.2.1

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**CHAPTER** 

**ONE** 

## **GET TAPIS**

# 1.1 Required packages

The following packages are required for TAPIS core functions. Version numbers correspond to those tested during development.

- SpliceGrapher (v0.2.4)
- Pysam (v0.8.1)
- matplotlib (v1.3.1)
- bx-python (v0.5.0)
- NumPy (v1.8.2)
- GMAP (v2015-07-23)

#### 1.2 Download

current release: v1.1.2

TAPIS is hosted on bitbucket https://bitbucket.org/comp\_bio/tapis

#### 1.3 Install

```
$ tar zxvf tapis_<version>.tgz
$ cd tapis_<version>.tgz
$ python setup.py install
```

**Note:** To install in a user directory, use the option:

--home=/Path/To/Local/Library

**CHAPTER** 

**TWO** 

#### **TUTORIAL**

This tutorial is meant as a complete walk-through for identifying transcripts and poly(A) sites from PacBio reads.

- 1. Align and clean reads
- 2. Cluster reads and analyze transcripts and poly(A) sites

## 2.1 Align reads

**TAPIS** accepts any sorted, indexed BAM file for long reads but it provides a method that cleans and aligns reads with high accuracy and efficiency. To align and clean reads use the following provided script **alignPacBio.py**. Before running the script, you will need to run **gmap\_build** to make a genome reference index.

```
usage: alignPacBio.py [-h] [-v] [-i ITERATIONS] [-e EDR] [-o OUTDIR]
                      [-p PROCS] [-K MAXINTRON]
                     indexesDir indexName reference fasta
Iteratively fix aligned reads using reference genome
positional arguments:
 indexesDir
                       directory to gmap indexes
 indexName
                       name of gmap index
 reference
                      Reference sequence
                       Reads to align
 fasta
optional arguments:
 -h, --help
                      show this help message and exit
                  Verbose mode
  -v, --verbose
 -i ITERATIONS, --iterations ITERATIONS
                       Number of aligment iterations, default=3
 -e EDR, --edr EDR
                       Edit distance ratio, default=10
 -o OUTDIR, --outdir OUTDIR
                       Output directory, default=./cleanedAlignments
 -p PROCS, --procs PROCS
                       Number of processors, default=1
 -K MAXINTRON, --maxIntron MAXINTRON
                       maximum intron length for gmap, default=8000
```

## 2.2 Creating indexed, sorted BAM files

If your cleaned/aligned reads are in the form of a SAM file, you can convert it to a indexed, sorted BAM file using **convertSam.py**.

```
usage: convertSam.py [-h] [-o BAMFILE] [-p PROCS] [-m MEMORY] [-v] samfile
Generate sorted BAM and index files for given SAM file
positional arguments:
 samfile
                        Samfile to convert
optional arguments:
 -h, --help
                       show this help message and exit
 -o BAMFILE, --outfile BAMFILE
                       Name of converted BAM file [default=<sambase>.bam]
 -p PROCS, --procs PROCS
                        Number of processors to use for BAM sorting (default
 -m MEMORY, --memory MEMORY
                        Max memory (in GBs) for each processor used for BAM
                        sorting (default 2)
 -v, --verbose
                       Print verbose output
```

## 2.3 Running TAPIS

```
$ run_tapis.py --help
usage: run_tapis.py [-h] [-v] [-p] [-o OUTDIR] [-t TRIMMAX] [-w W]
                    [-m MINDIST]
                   geneModel bamfile
Assemble transcripts from PacBio alignments
positional arguments:
                     Gene models annotation file (GFF/GTF)
 geneModel
 bamfile
                      Aligned reads file (sorted and indexed)
optional arguments:
 -h, --help
                      show this help message and exit
 -v, --verbose
                      Verbose mode
 -p, --plot
                      Plot novel gene graphs and poly(A) figures, default is
                       no plotting
 -o OUTDIR, --outdir OUTDIR
                       Output directory for TAPIS results, default=tapis_out
 -t TRIMMAX, --trimMax TRIMMAX
                       Maximum length of read trimming to tolerate, default=5
 -w W, --w W
                       Width of peaks when searching for poly(A) sites,
                       default=5
 -m MINDIST, --minDist MINDIST
                       Minimum distance between any two poly(A) sites,
                       default=20
```

While **TAPIS** offers many options, default values should work for most cases.

## 2.4 Interpreting TAPIS output

**TAPIS** builds an output directory as follows:

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```
$ tree my_result
  tapis_out
  |-- polyAFigures
  |  |-- gene1.png
  |  |-- gene2.pbg
  |  |-- ...
  |  |-- geneN.png
  |-- novelGraphs
  |  |-- chrom_start_end_strand.pdf
  |  |-- ...
  |-- assembled.gtf
  |-- novelGenes.csv
  |-- novelGenes.fa
  |-- polyA_summary.csv
```

- polyAFigures contains poly(A) site depictions for genes with at least one poly(A) site supported by long reads.
- novelGraphs contains splice graph figures for transcripts not found in within any annotated gene.
- assembled.gtf gene models for transcripts detected in long reads
- novelGenes.csv tab-delimited file containing summary of novel genes detected

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#### **CHAPTER**

# **THREE**

# **CONTACT**

**TAPIS** is developed by Mike Hamilton at Colorado State University.

Bug reports and feature requests can be submitted through bitbucket.

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[SG] Rogers, MF, Thomas, J, Reddy, AS, Ben-Hur, A (2012). SpliceGrapher: detecting patterns of alternative splicing from RNA-Seq data in the context of gene models and EST data. *Genome Biol.*, 13, 1:R4.