

Episo-User Guide-V1.0

1) Quick Reference

Episo needs a working version of Perl and it is run from the command line. Meanwhile, Bowtie, Tophat and Cufflinks need to be installed on your computer. First you need to download a transcript annotation file from the Ensembl or NCBI websites. Episo supports the reference transcriptome sequence files in FastA format, allowed file extensions are either .fa or .fasta. The following examples will use the paired-ends files

'example_1.fastq&example_2.fastq' (it contains 2,500,876 reads in FastQ format, 101 bp long reads, simulated by Fluxsimulator) and the transcript annotation file. They are available for download from the Episo homepage.

(1) Compiling the program

When you use the UNIX (linux, Mac OSX) you should compile some programs. You can use gcc or any ANSI C-compatible compiler. The source codes are from my Episo package. The commands are as follow.

```
gcc -o contrans contrans.c
```

```
gcc -o compare-paired compare-paired.c -lm
```

```
gcc -o anti-bisulfite anti-bisulfite.c
```

```
gcc -o selsam selsam.c
```

```
gcc -o methylation_ratio methylation_ratio.c -lm
```

```
gcc -o isoform_filter isoform_filter.c
```

(2) Generating reference transcriptome

The first step is to generate the transcript file according to annotation transcript by using the program Cufflinks. The command is as follows.

Usage: cufflinks -G annotation.gtf convert.sam

Note. The file annotation.gtf is from the Ensembl or NCBI websites and the file convert.sam is from the Episo package.

This will produce one output file: transcripts.gtf.

The second step is to generate the reference transcriptome by using the program contrans. The command is as follows.

Usage: contrans contrans.ctl

Note. The format of control file contrans.ctl is as follows. The genome file genome.fa is from the Ensembl or NCBI websites.

outfile = out	* recording the running information
gtffile = transcripts.gtf	* gtf file generated by cufflinks
fafile = genome.fa	* genome file
transfile = out_trans	* recording the transcript
seqfile = out_seq.fa	* recording the sequence of each trascript
seqlength = 50	* the length of sequence in the seqfile

This will produce two output files whose names are from the parameter “seqfile” and “transfile” in the control file contrans.ctl.

(3) Generating transcriptome indexing

Usage: bismark_genome_preparation [options] <path_to_transcriptome_folder>

Note. The output seqfile generated by the program contrans or reference transcriptome sequence file downloaded from the Ensembl or NCBI websites should be put in the <path_to_transcriptome_folder>.

A typical transcriptome indexing could be like this:

```
bismark_genome_preparation --path_to_bowtie /usr/local/bowtie --verbose
/data/transcriptome/
```

(4) Calling the methylation site

Usage: bismark-liu [options] <transcriptome_folder> -1 <mates> -2 <mates>

A typical calling example could be like this:

```
bismark-liu --path_to_bowtie /usr/local/bowtie --vanilla --sam -n 2 /data/transcriptome/ -1
example_1.fastq -2 example_2.fastq
```

This will produce three output files:

- (a) example_1.fastq_bismark_pe.txt (contains all alignments and methylation call strings)
- (b) example_1.fastq_bismark_pe_mul.txt (contains the transcript information, where the alignment belongs)
- (c) example_1.fastq_bismark_PE_report.txt (contains alignment and methylation summary)

Note. The options “vanilla” and “sam” are necessary and the bowtie version must be bowtie1. The program compare-paired and the transfile generated by contrans must be in the same directory in which bismark-liu is.

(5) Generating the anti-bisulfite RNA-Seq data

Usage: anti-bisulfite anti-bisulfite.ctl

Note. The format of control file anti-bisulfite.ctl is as follows.

outfile = out	* recording the running information
intxtfile = example_1.fastq_bismark_pe.txt	* bismark_pe.txt file generated by bismark-liu
outreadfile = anti-example	* the index of anti-bisulfite RNA-Seq fastq file
flag = p	* p means paired-ends; s means singled-end
skipped_number = 1	* the number of the rows which will be skipped
inmultxtfile = example_1.fastq_bismark_pe_mul.txt	* bismark_pe_mul.txt file generated by bismark-liu

This will produce three output files according to the control file anti-bisulfite.ctl:

(a) anti-example_1.fastq and anti-example_2.fastq (an anti-bisulfite RAN-Seq paired-end file)

(b) methylation_summary (contains the transcript information, where the methylation alignment belongs)

(6) Estimating the methylation level of each isoform

The first step is to analysis the anti-example_1.fastq and anti-example_2.fastq by using the program TopHat. The options “--bowtie1” and “--no-convert-bam” must be chosen. **The second step** is to generate the file which contains the methylation alignments only according to the sam file generated by TopHat. The command is as follows.

Usage: selsam <.sam> <skipped_number>

Note. The .sam file was generated by TopHat. The skipped_number is the number of rows which include the sign “@” in the .sam file. The output file name is accepted_hits_methylation.sam.

The third step is to use the program Cufflinks to analysis the sam file which was generated by TopHat and the file accepted_hits_methylation.sam generated by selsam respectively. The option “-G” must be chosen and the gtf file comes from the contrans.ctl file.

The last step is to estimate the methylation level of each transcript. The command is as follows.

Usage: methylation_ratio <isoform_filter> <isoform_methylation_filter> <total_number> <methylation_number>

Note. The files in <isoform_filter> and <isoform_methylation_filter> are got by using the program isoform_filter according to the files in <isoform> and <isoform_methylation>. The file in <isoform> is the output file isoforms.fpk_tracking when using the Cufflinks to analysis the sam file which was generated by TopHat. The file in <isoform_methylation> is the output file isoforms.fpk_tracking when using the Cufflinks to analysis the file accepted_hits_sam generated by selsam. The number in <total_number> is the number of rows of the file ***_bismark_pe_mul.txt generated by bismark-liu. The number in <methylation_number> is the number of rows of the file methylation_summary generated by anti-bisulfite.

The filtering command is as follows.

Usage: isoform_filter <isoform> <isoform_methylation> <filter_number>

Note. The number in <filter_number> is FPKM value under which the records in the files <isoform> and <isoform_methylation> are deleted.

2) Estimating the methylation level of each transcript at a single site

After generating the anti-bisulfite RNA-Seq data, we can estimate the methylation level of each transcript at a single site. In order to estimate the methylation level of each transcript at a single site, we need the output files generated by TopHat&selsam according to the output files generated by anti-bisulfite. The pipeline is as follows.

(1) Compiling the program

```
gcc -o selreads selreads.c -lm
```

```
gcc -o selsam-single-parallel selsam-single-parallel.c
```

(2) Outputting the alignments which include the assigned single site

Usage: selreads selreads.ctl

Note. The format of control file selreads.ctl is as follows. The value of parameter “length” is the length of read which in the fastq file generated by anti-bisulfite.

outfile = out	* recording the running information
intxtfile = example_1.fastq_bismark_pe.txt	* bismark_pe.txt file generated by bismark-liu
intransfile = out_trans	* trans file generated by contrans
outreadfile = sel_reads	* the index of reads file in which the reads include assigned methylated site
location = 59	* the methylated site location
flag = p	* p means paired-ends; s means singled-end
length = 75	* the length of read
chrom_name = test_chromosome	* the name for chromosome
skipped_number = 1	* the number of the rows which will be skipped

This will produce one output file: methylation_summary_sam (contains the names of alignments which include the assigned single site in the control file selreads.ctl)

(3) Generating the sam files which are used to be analysed by Cufflinks

Usage: selsam-single-parallel <sam file> <methylation_summary_sam> <skipped_number> <tag> <the number of rows in methylation_summary_sam> 1 <output file name>

Note. The value in <tag> is 1 or 0. When the value in <tag> is 1, the file in <sam file> is the output file generated by TopHat according to the output files generated by anti-bisulfite. When the value in <tag> is 0, the file in <sam file> is the output file generated by selsam. The value in <skipped_number> is the number of rows which include the sign “@” in the .sam file.

(4) Using the program Cufflinks to analysis the output files generated by selsam-single-parallel when the value in <tag> is 1 and 0 respectively. The option “-G” must be chosen and the gtf file comes from the contrans.ctl file.

(5) Estimating the methylation level of each isoform at a single site

Usage: methylation_ratio <isoform_filter> <isoform_methylation_filter> <total_number>
<methylation_number>

Note. The files in <isoform_filter> and <isoform_methylation_filter> are got by using the program isoform_filter according to the files in <isoform> and <isoform_methylation>. The file in <isoform> is the output file isoforms.fpk_tracking when using the Cufflinks to analysis the output file generated by selsam-single-parallel when the value in <tag> is 1. The file in <isoform_methylation> is the output file isoforms.fpk_tracking when using the Cufflinks to analysis the output file generated by selsam-single-parallel when the value in <tag> is 0. The number in <total_number> is the number of rows of the file methylation_summary_sam generated by selreads. The number in <methylation_number> is the number of rows which include the character “methylated_liu” in the file methylation_summary_sam generated by selreads.

3) Computing the methylation rate of each site

The methylation rate of each site is the ratio of methylated reads in all reads which include the site. The methylated read is the read in which the assigned site is methylated. The pipeline is as follows.

(1) Compiling the program

```
gcc -o trans2genom-bismark trans2genom-bismark.c -lm
```

```
gcc -o trans2genom-bismark-methy trans2genom-bismark-methy.c -lm
```

(2) Extracting the methylation call for every single C analysed

Usage: bismark_methylation_extractor-liu [options] <filenames>

A typical command to extract context-dependent (CpG/CHG/CHH) methylation could like this:

```
Bismark_methylation_extractor-liu -p --vanilla --no_overlap --comprehensive  
example_1.fastq_bismark_pe.txt
```

Note. The file in <filenames> is generated by bismark-liu.

This will produce six output files:

(a) CHG_context_example_1.fastq_bismark_pe.txt

(b) CHH_context_example_1.fastq_bismark_pe.txt

(c) CpG_context_example_1.fastq_bismark_pe.txt

(3) Computing the methylation rate of each site

In order to compute the methylation rate of each site, a series of shell script commands should be executed. These commands are as follows.

```
#!/bin/sh

sort -k 2 <out_trans> >> out_trans-sort

sed '1d' <CHG_context_*_pe.txt> | sort -k 3 >> CHG_CTOB-sort.txt

nice ./trans2genom-bismark CHG_CTOB-sort.txt out_trans-sort

cat methylation_genom >> methylation_genom-all

rm methylation_genom -f

sed '1d' <CHH_context_*_pe.txt> | sort -k 3 >> CHH_CTOB-sort.txt

nice ./trans2genom-bismark CHH_CTOB-sort.txt out_trans-sort

cat methylation_genom >> methylation_genom-all

rm methylation_genom -f

sed '1d' <CpG_context_*_pe.txt> | sort -k 3 >> CpG_CTOB-sort.txt

nice ./trans2genom-bismark CpG_CTOB-sort.txt out_trans-sort

cat methylation_genom >> methylation_genom-all

rm methylation_genom -f

sort -k 4 methylation_genom-all >> methylation_genom-all-sort

nice ./trans2genom-bismark-methy methylation_genom-all-sort
```

Note. The file in <out_trans> is generated by contrans. The files in <CHG_context_*_pe.txt>, <CHH_context_*_pe.txt>, and <CpG_context_*_pe.txt> are the output files generated by bismark_methylation_extractor-liu.