A Protocol for m6A-seq Data Analysis

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Overview

This protocol serves as a step-by-step guide to m6A-seq data analysis. We include all the major steps involved in seven modules: database checking, data preprocessing, differential expression analysis, site detection(host), site detection(virus), differential methylation analysis, and reference-based analysis. We demonstrate the procedures by analyzing two published MeRIP-seq datasets from scratch based on UNIX shell and R system. The data information are summarized in the "Dataset" subsection. The software information and workflow are summarized in the following table.

Datasets

The datasets we are using for this protocol are from two studies described in Brandon et al (2018) [1] and Martin et al (2013) [2]. These datasets can be found and accessed on Gene Expression Omnibus (GEO) database.

• Expression profile of viral and cellular m6A epitranscriptomes in KSHV life cycle in Homo sapiens.

- Accession code in GEO database: GSE93676
 - GSM2460344 iSLK-uninf-input
 - SRR5978827
 - SRR5978828
 - SRR5978829
 - <u>GSM2460345</u> iSLK-uninf-ip
 - SRR5978834
 - SRR5978835
 - <u>SRR5978836</u>
 - GSM2460350 iSLK-KSHV_BAC16-48hr-ip
 - SRR5978869
 - SRR5978870
 - <u>SRR5978871</u>
 - <u>GSM2460351</u> iSLK-KSHV_BAC16-48hr-input
 - SRR5179446
 - <u>SRR5179447</u>
 - <u>SRR5179448</u>
- Methylation profile of the Obesity-associated FTO gene in Mus musculus.
 - Accession code in GEO database: GSE47217
 - <u>GSM1147014</u> WT1_MeRIP
 - SRR866997
 - SRR866999
 - SRR867001
 - GSM1147015 WT1_No IP
 - <u>SRR866998</u>
 - <u>SRR867000</u>
 - SRR867002
 - GSM1147020 KO1_MeRIP
 - SRR866991
 - SRR866993
 - <u>SRR866995</u>
 - GSM1147021 KO1_No IP
 - SRR866992
 - <u>SRR866994</u>
 - <u>SRR866996</u>

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M0. Check Databases

By checking the following databases before analyzing the datasets, we could have a basic understanding of the data in terms of estimated quantitative profiles, conservation, biological functions, annotation, and disease association of epi-transcriptome.

m6A_Atlas

m6A-Atlas is a comprehensive knowledgebase for unraveling the m6A epitranscriptome [3].

RMDisease

<u>RMDisease</u> is a database of genetic variants that affect RNA modifications, with implications for epitranscriptome pathogenesis [4].

WHISTLE

<u>WHISTLE</u> is a high-accuracy map of the human m6A epitranscriptome predicted using a machine learning approach [5].

M1. Data Preprocessing

Download GEO Data (SRA Toolkit)

The Sequence Read Archive (SRA) is a publicly accessible archive for high throughput sequencing data. The <u>SRA Toolkit</u> from NCBI is a collection of tools for using data in the INSDC SRA. It takes the following steps to download data from SRA:

Install and Config SRA Toolkit

```
# Download and extract the latest version
$ wget --output-document sratoolkit.tar.gz https://ftp-
trace.ncbi.nlm.nih.gov/sra/sdk/2.10.9/sratoolkit.2.10.9-ubuntu64.tar.gz
$ tar -vxzf sratoolkit.2.10.9-ubuntu64.tar.gz

# Append the path to your PATH environment variable:
$ export PATH=$PATH:/path/to/sratoolkit.2.10.9-ubuntu64/bin

# Verify the installation
$ which fastq-dump
```

Download Data from SRA

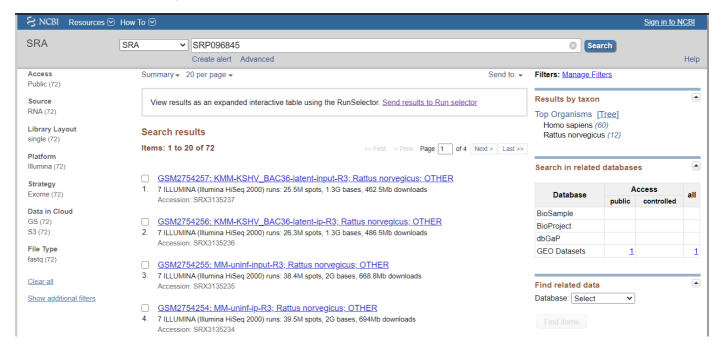
1. Access the GEO summary page by searching "GSE93676" on GEO website.



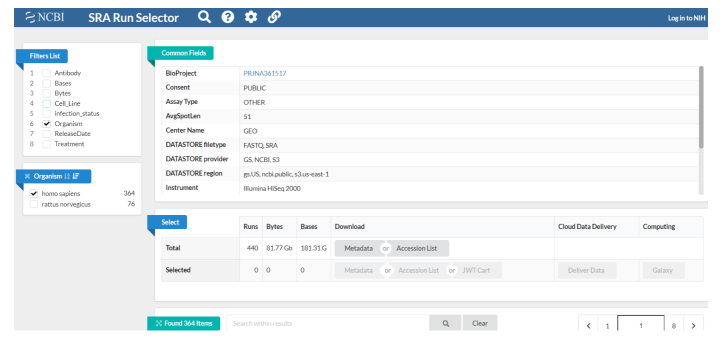
2. Find a link for "SRA" under the heading "Relations".

```
Submission date Jan 16, 2017
Last update date May 15, 2019
Contact name
                 Hui Liu
                 lhcumt@hotmail.com
E-mail(s)
Organization name China University of Mining and Technology
Street address
                 #1 Daxue Road
City
                 Xuzhou
State/province
                 Jiangsu
ZIP/Postal code
                 221116
                 China
Country
                 GPL11154 Illumina HiSeq 2000 (Homo sapiens)
Platforms (2)
                 GPL14844 Illumina HiSeq 2000 (Rattus norvegicus)
                 GSM2460344 iSLK-uninf-input
Samples (72)
■ More...
                 GSM2460345 iSLK-uninf-ip
                 GSM2460346 iSLK-KSHV BAC16-latent-input
Relations
BioProject
                 PRJNA361517
SRA
                 SRP096845
```

3. Click on the link (SRP096845) which sends you to a page of all the biological samples with specific runs and files in this study.



4. To find files of interest in one comprehensive list, navigate to the bottom of the page then click: "send to" > "Run Selector" > "go". Use "Filter List" to narrow down the choices.



5. Extract FastQ files from SRA-accession using SRA-Toolkit

```
#!/bin/bash
cd /path/to/raw_data/homo/
fetch_dump(){
prefetch $1
fastq-dump $1
}
export -f fetch_dump
for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
fetch_dump ${s}
done
```

```
#!/bin/bash
cd /path/to/raw_data/mm10/
fetch_dump(){
prefetch $1
fastq-dump $1
}
export -f fetch_dump
for s in SRR866997 SRR866998 SRR866999 SRR867000 SRR867001 SRR867002 SRR866991
SRR866992 SRR866993 SRR866994 SRR866996
do
fetch_dump ${s}
done
```

For paired-end data, you need to add --split-files option in the fastq-dump command.

```
fastq-dump --split-files SRR866997
```

Quality Assessment (FastQC)

It is always necessary to assess the quality of the sequence reads in FASTQ files from the sequencing facility. <u>FastQC</u> is a quality control application for high-throughput sequencing data. By using FastQC, we could be aware of any problems in raw sequence data before moving on to the next step.

Install FastQC

```
# FastQC requires a suitable 64-bit Java Runtime Environment (JRE) installed and in the
path. Check the version of Java:
$ java -version

# For Linux, download and extract the latest version of FastQC from [project website]
(http://www.bioinformatics.babraham.ac.uk/projects/download.html#fastqc):
$ wget http://www.bioinformatics.babraham.ac.uk/projects/fastqc/fastqc_v0.11.9.zip
$ unzip fastqc_v0.11.9.zip

# Append the path to your PATH environment variable:
$ export PATH=$PATH:/path/to/FastQC

# Verify installation:
$ fastqc -help
```

Run FastQC

```
# Examine the quality of one FastQ file:
$ fastqc -o /path/to/fastqc_result/ /path/to/raw_data/homo/SRR5978869.fastq

# or examine the quality of multiple FastQ files:
$ fastqc -o /path/to/fastqc_result/ -t 6 /path/to/raw_data/homo/*.fastq
```

Note: —o (or ——outdir) will create all output files in the specified output directory. —t specifies the number of files / threads that can be processed in parallel.

FastQC Results

FastQC produces two output files for each FastQ file: an HTML report ("SRR5978869_fastqc.html") and a packed file ("SRR5978869_fastqc.zip").

You could transfer the HTML file to local place by *FileZilla* (mac) or *WinSCP* (win), and open the file in browser. A screenshot of part of the HTML file is shown below.

№FastQC Report

Summary



Per base sequence quality

Per sequence quality scores

Per base sequence content

Per sequence GC content

Per base N content

Sequence Length Distribution

Sequence Duplication Levels

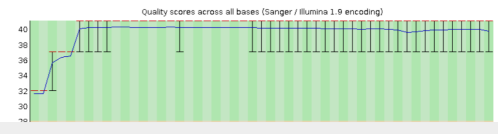
Overrepresented sequences

Adapter Content



Measure	Value		
Filename	SRR5978869.fastq		
File type	Conventional base calls		
Encoding	Sanger / Illumina 1.9		
Total Sequences	4724702		
Sequences flagged as poor quality	0		
Sequence length	51		
%GC	58		

Per base sequence quality



Produced by FastQC (version 0.11.9)

Note that two of the most important analysis modules in FastQC are "Per base sequence quality" plot and the "Overrepresented sequences" table. The "Per base sequence quality" plot provides the distribution of quality scores across all bases at each position in the reads. The "Overrepresented sequences" table displays the sequences (at least 20 bp) that occur in more than 0.1% of the total number of sequences, which aids in identifying contamination. You could also refer to Analysis Modules in FastQC documentation for the interpretation of the HTML report.

The other output is a zip file for each sample. You could unpack the zip files and have a look at the summary.

```
# Unpack a .zip file in the result directory:
$ unzip SRR5978869_fastqc.zip

# or unpack .zip files in the result directory:
$ for zip in *.zip
do
unzip $zip
done

# To see the content of a single summary file:
$ cat SRR5978827_fastqc/summary.txt

# or cat all summary files into one text file and have a look at it:
$ cat */summary.txt > ~/all/fastqc_summaries.txt
$ cat ~/all/fastqc_summaries.txt
```

For paired-end data, since the two reads of the pair are generated separately, trying to get statistics like per base sequence quality on the combined forward and reverse reads would make no sense. In this case, you may check quality by inputting BAM files.

Reads Trimming (Trim Galore)

<u>Trim Galore</u> is a Perl wrapper around Cutadapt and FastQC to consistently apply adapter and quality trimming to FastQ files. We will use this tool for quality trimming, adapter trimming, and removing short sequences.

Install Trim Galore

Before installation, ensure that <u>Cutadapt</u> and <u>FastQC</u> are already installed.

```
# Check the version of cutadapt
$ cutadapt --version

# Check the version of FastQC
$ fastqc -v
```

Install the latest version of Trim Galore from Github or project website:

```
# Install Trim Galore
$ curl -fsSL https://github.com/FelixKrueger/TrimGalore/archive/0.6.6.tar.gz -o
trim_galore.tar.gz
$ tar xvzf trim_galore.tar.gz

# Verify installation
$ trim_galore -v
```

Adaptive Quality and Adapter Trimming

In this procedure, first, low-quality base calls are trimmed off from the 3' end of the reads before adapter removal. Next, adapter sequences from the 3' end of reads are detected and removed by cutadapt. Lastly, trimmed short sequences (default: < 20bp) are filtered.

```
#!/bin/bash
trimGalore(){
    trim_galore -o /path/to/trim_galore_result/ /path/to/raw_data/homo/$1.fastq
}
export -f trimGalore
for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
trimGalore ${s}
done
```

```
#!/bin/bash
trimGalore(){
trim_galore -o /path/to/trim_galore_result/ /path/to/raw_data/mm10/$1.fastq
}
export -f trimGalore
for s in SRR866997 SRR866998 SRR866999 SRR867000 SRR867001 SRR867002 SRR866991
SRR866992 SRR866993 SRR866994 SRR866995 SRR866996
do
trimGalore ${s}
done
```

For trimming paired-end data, you need to add a _-paired option in the trim_galore command.

```
trim_galore --paired -o /path/to/trim_galore_result/ *_1.fastq *_2.fastq
```

Outputs

Trim Galore produced two output files for each FastQ file: one text file ("SRR5978869.fastq_trimming_report.txt") and a trimmed FastQ file ("SRR5978869_trimmed.fq").

1. The text file

The text file provides a summary of running parameters.

```
# To see the first few lines of the text file
$ head SRR5978869.fastq_trimming_report.txt
```

```
SUMMARISING RUN PARAMETERS

Input filename: SRR5978869.fastq

Trimming mode: single-endw

Trim Galore version: 0.6.4_dev

Cutadapt version: 3.2

Number of cores used for trimming: 1

Quality Phred score cutoff: 20

Quality encoding type selected: ASCII+33
```

2. The trimmed FastQ

The trimmed FastQ file can be used for further analysis.

Reads Alignment (HISAT2)

<u>HISAT2</u> is a fast and sensitive alignment program for mapping next-generation sequencing reads to reference genome(s) [6]. We are going to use this tool to align the reads to hg19 genome, HHV8 genome, and mm10 genome, respectively.

Install HISAT2

```
# Download and extract the latest version
$ wget ftp://ftp.ccb.jhu.edu/pub/infphilo/hisat2/downloads/hisat2-2.0.4-
Linux_x86_64.zip
$ unzip hisat2-2.0.4-Linux_x86_64.zip

# Append to PATH environment variable
$ export PATH=$PATH:/path/to/hisat2-2.0.4

# Verify installation
$ hisat2 --help
$ hisat2 --version
```

Install Samtools

<u>Samtools</u> is a suite of programs for interacting with high-throughput sequencing data [7]. SAM files produced by HISAT2 must be sorted and converted to BAM using samtools before running StringTie.

```
# Download and extract samtools
$ wget https://github.com/samtools/samtools/releases/download/1.11/samtools-
1.11.tar.bz2 -O samtools-1.11.tar.bz2
$ tar -xjvf samtools-1.11.tar.bz2

# Append to PATH environment variable
$ export PATH=$PATH:/path/to/samtools-1.11

# Verify installation
$ samtools --version
```

Read Alignment

1. Build indexes

You can either download HISAT2 indexes from its website

```
$ wget https://genome-idx.s3.amazonaws.com/hisat/hg19_genome.tar.gz
$ tar -zxvf hg19_genome.tar.gz
```

or download reference sequence and gene annotation from <u>Illumina iGenome</u> before building index by hisat2-build command.

```
$ cd /path/to/homo/
$ hisat2-build -p 20 hg19_genome.fa genome

$ cd /path/to/HHV8/
$ hisat2-build -p 20 hhv8_sequence.fasta genome

$ cd /path/to/mm10/
$ hisat2-build -p 20 mm10_genome.fasta genome
```

hisat2-build generates eight .ht2 files, from genome.1.ht2 to genome.8.ht2, which we will use for alignment in the next step.

2. Run HISAT2 with Samtools

Getting sorted BAM and index:

```
#!/bin/bash
cd /path/to/homo/ # where storing index
hisat_samtool(){
hisat2 -x genome --summary-file "$1".m6A.align_summary -p 5 -U
/path/to/trim_galore_result/"$1"_trimmed.fq | samtools view -Su |samtools sort -o
/path/to/homo_result/"$1"_sorted.bam
samtools index /path/to/homo_result/"$1"_sorted.bam
}
export -f hisat_samtool
```

```
for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
hisat_samtool ${s}
done
mkdir alignment_summary
mv *.align_summary alignment_summary/
```

```
#!/bin/bash
cd /path/to/mm10/ # where storing index
hisat_samtool(){
hisat2 -x genome --summary-file "$1".m6A.align_summary -p 5 -U
/path/to/trim_galore_result/"$1"_trimmed.fq | samtools view -Su |samtools sort -o
/path/to/mm10_result/"$1"_sorted.bam
samtools index /path/to/mm10_result/"$1"_sorted.bam
}
export -f hisat_samtool

for s in SRR866997 SRR866998 SRR866999 SRR867000 SRR867001 SRR867002 SRR866991
SRR866992 SRR866993 SRR866994 SRR866995 SRR866996
do
hisat_samtool ${s}
done
mkdir alignment_summary
mv *.align_summary alignment_summary/
```

```
#!/bin/bash
cd /path/to/hhv8/ # where storing index
hisat samtool(){
hisat2 -x genome --summary-file "$1".m6A.align_summary -p 5 -U
/path/to/trim_galore_result/"$1"_trimmed.fq | samtools view -Su |samtools sort -o
/path/to/hhv8 result/"$1" sorted.bam
samtools index /path/to/hhv8 result/"$1" sorted.bam
}
export -f hisat_samtool
for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
hisat samtool ${s}
done
mkdir alignment summary
mv *.align_summary alignment_summary/
```

Note that if the aligned results are going to assemble transcript with StringTie, a --dta option is
necessary to include the tag xs to indicate the genomic strand that produced the RNA from which the
read was sequenced. This is required by StringTie. The hisat2-samtools command should be
modified as

```
hisat2 -x genome --summary-file SRR5978827.m6A.align_summary -p 5 -U /path/to/trim_galore_result/SRR5978827_trimmed.fq --dta | samtools view -Su |samtools sort -o /path/to/homo_result/SRR5978827_sorted.bam
```

```
--dta/--downstream-transcriptome-assembly
```

Report alignments tailored for transcript assemblers including StringTie. With this option, HISAT2 requires longer anchor lengths for de novo discovery of splice sites. This leads to fewer alignments with short-anchors, which helps transcript assemblers improve significantly in computation and memory usage.

• Also note that for paired end data, you need to modify the hisat2 command as follows to get SAM file

```
hisat2 -x genome --summary-file $s.m6A.align_summary -p 5 -1
/path/to/trim_galore_result/SRR5978827_trimmed_1.fq -2
/path/to/trim_galore_result/SRR5978827_trimmed_2.fq -S
/path/to/homo_result/SRR5978827.sam
```

or modify the hisat2-samtools combined command to directly get sorted BAM file

```
hisat2 -x genome --summary-file SRR5978827.m6A.align_summary -p 5 -1 /path/to/trim_galore_result/SRR5978827_trimmed_1.fq -2 /path/to/trim_galore_result/SRR5978827_trimmed_2.fq | samtools view -Su |samtools sort -o /path/to/homo_result/SRR5978827_sorted.bam
```

M2. Differential Expression

Transcript Assembly and Quantification (StringTie)

StringTie is a highly efficient assembler for RNA-Seq alignments using a novel network flow algorithm [8]. It can simultaneously assemble and quantify expression levels for the features of the transcriptome in a Ballgown readable format. StringTie's output can be processed by specialized software like Ballgown (Alyssa et al. (2014)), Cuffdiff (Cole et al. (2010)) or other programs (DESeq2 (Anders & Huber (2010)), edgeR (Robinson et al. (2010)), etc).

The input SAM(BAM) file must be sorted by reference position. Every spliced read alignment in the input must contain the tag xs to indicate the genomic strand that produced the RNA from which the read was sequenced. These requirements are met by running HISAT2 with ——dta option and samtools.

Install StringTie

```
# Download and extract StringTie
$ wget http://ccb.jhu.edu/software/stringtie/dl/stringtie-2.1.4.Linux_x86_64.tar.gz
$ tar xvfz stringtie-2.1.4.Linux_x86_64.tar.gz

# Append to PATH environment variable
$ export PATH=$PATH:/path/to/stringtie-2.1.4.Linux_x86_64

# Verify installation
$ stringtie --version
```

Run StringTie

Run with the downloaded gene annotation:

```
#!/bin/bash
stringTie1(){
    stringtie /path/to/homo_result/"$1"_sorted.bam -p 20 -o
    /path/to/stringtie_homo/"$1".gtf -G /path/to/hg19_annotation.gff
}
export -f stringTie1

for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
stringTie1 ${s}
done
```

```
#!/bin/bash
stringTiel(){
    stringtie /path/to/mm10_result/"$1"_sorted.bam -p 20 -o
    /path/to/stringtie_mm10/"$1".gtf -G /path/to/mm10_annotation.gff
}
export -f stringTiel

for s in SRR866997 SRR866998 SRR866999 SRR867000 SRR867001 SRR867002 SRR866991
SRR866992 SRR866993 SRR866994 SRR866995 SRR866996
do
stringTiel ${s}
done
```

```
# Generate a non-redundant set of transcripts
$ cd /path/to/stringtie_homo/
$ stringtie --merge -G /path/to/hg19_annotation.gff -p 20 -o homo_stringtie_merged.gtf
homo_stringtie_list.txt

$ cd /path/to/stringtie_mm10/
$ stringtie --merge -G /path/to/mm10_annotation.gff -p 20 -o mm10_stringtie_merged.gtf
mm10_stringtie_list.txt
```

The text file contains all GTF files generated when assembling the read alignments.

```
SRR5978827.gtf
SRR5978828.gtf
```

Estimate transcript abundances and generate read coverage tables for Ballgown. Note that this is the only case where the -g option is not used with a reference annotation

```
#!/bin/bash
stringTie2(){
stringtie /path/to/homo_result/"$1"_sorted.bam -eB -p 20 -G
/path/to/stringtie_homo/homo_stringtie_merged.gtf -o /path/to/stringtie_homo/"$1".gtf
}
export -f stringTie2

for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
mkdir $s
cd $s
stringTie2 ${s}
done
```

```
#!/bin/bash
stringTie2(){
stringtie /path/to/mm10_result/"$1"_sorted.bam -eB -p 20 -G
/path/to/stringtie_mm10/mm10_stringtie_merged.gtf -o /path/to/stringtie_mm10/"$1".gtf
}
export -f stringTie2

for s in SRR866997 SRR866998 SRR866999 SRR867000 SRR867001 SRR867002 SRR866991
SRR866992 SRR866993 SRR866994 SRR866995 SRR866996
do
mkdir $s
cd $s
stringTie2 ${s}
done
```

Note:

Arguments and Options	Description
-G	Use the reference annotation file (in GTF or GFF3 format) to guide the assembly process
-e	Limits the processing of read alignments to only estimate and output the assembled transcripts matching the reference transcripts given with the $-g$ option (requires $-g$, recommended for $-B/-b$)
-B	enables the output of <i>Ballgown</i> input table files (*.ctab) containing coverage data for the reference transcripts given with theg option
-b <path></path>	Same as -B option, but these files will be created in the provided directory <path> instead of the directory specified by the _o option</path>
-p <int></int>	Specify the number of processing threads (CPUs) to use for transcript assembly. The default is 1

Outputs

- 1. StringTie's primary GTF output ("SRR5978869.gtf") contains details of the transcripts that StringTie assembles from RNA-Seq data.
- 2. Ballgown input table files ((1) e2t.ctab, (2) e_data.ctab, (3) i2t.ctab, (4) i_data.ctab, and (5) t_data.ctab) contain coverage data for all transcripts.
- 3. Merged GTF ("homo_stringtie_merged.gtf") is a uniform set of transcripts for all samples.

Differential Expression Analysis (Ballgown)

<u>Ballgown</u> is an R/Bioconductor package for flexible, isoform-level differential expression analysis of RNA-seq data [9]. Ballgown's data structures make it easy to use table-based packages like limma (<u>Smyth (2005)</u>), limma Voom (<u>Law et al. (2014)</u>), DESeq (<u>Anders & Huber (2010)</u>), DEXSeq (<u>Anders et al. (2012)</u>), or edgeR (<u>Robinson et al. (2010)</u>) for differential expression analysis.

Ballgown requires three pre-processing steps:

- 1. MeRIP-Seq reads should be aligned to a reference genome. (HISAT2)
- 2. A transcriptome should be assembled, or a reference transcriptome should be downloaded. (**StringTie**)
- 3. Expression for the features (transcript, exon, and intron junctions) in the transcriptome should be estimated in a Ballgown readable format. (**StringTie**)

Differential Expression Analysis

An example of the working directory:

```
stringtie_homo/
   SRR5978827/
        e2t.ctab
        e_data.ctab
        i2t.ctab
        i_data.ctab
        t_data.ctab
        SRR5978828/
        e2t.ctab
        e_data.ctab
        i2t.ctab
        i2t.ctab
        i2t.ctab
        i2t.ctab
        idata.ctab
        i2t.ctab
        idata.ctab
        i.data.ctab
        i.data.ctab
        i.data.ctab
```

Create an R script load_bg.R for loading ballgown object:

```
library(ballgown)
bg <- ballgown(dataDir="/path/to/stringtie_homo", samplePattern='SRR', meas='all')
save(bg, file='bg.rda')</pre>
```

Then run this script in terminal

```
$ R CMD BATCH load_bg.R
```

Differential expression analysis with Ballgown:

```
# Set directory
setwd("/path/to/ballgown/")

# Load R packages
library(ballgown)
library(genefilter)

# Load data
bg = ballgown(dataDir="/path/to/stringtie_homo", samplePattern='SRR', meas='all')

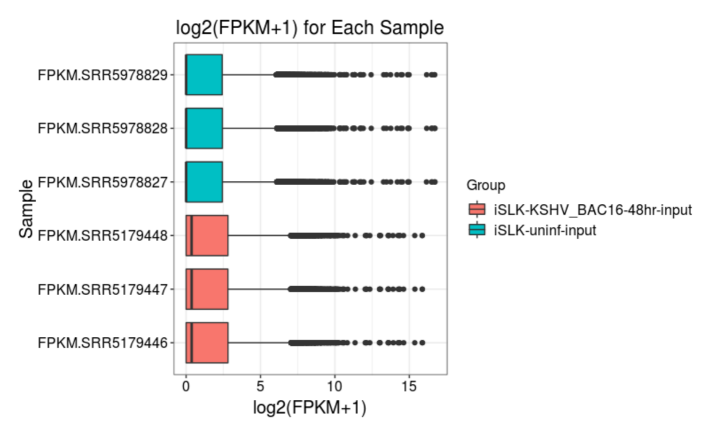
# Save data for backup
save(bg, file='bg.rda')

# Load all attributes and gene names
bg_table = texpr(bg, 'all')
bg_gene_names = unique(bg_table[, 9:10])
```

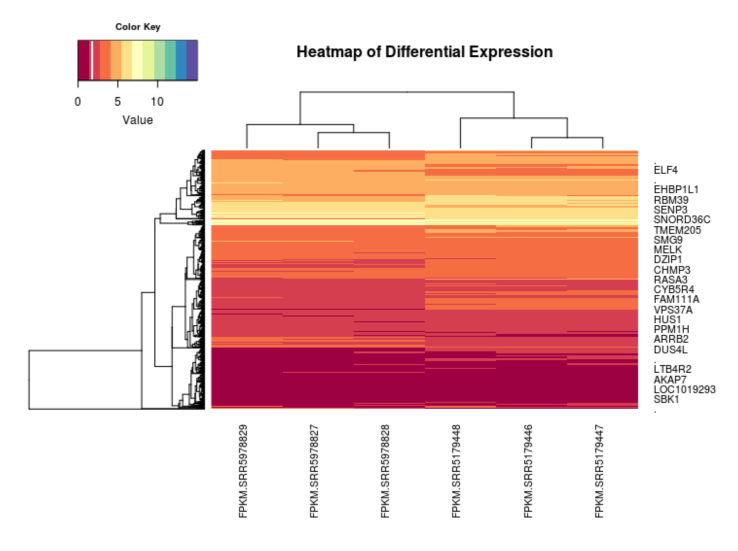
```
# Get gene expression data frame
gene expression = as.data.frame(gexpr(bg))
# Add pData specifying groups
group = c(rep("iSLK-KSHV_BAC16-48hr-input", 3), rep("iSLK-uninf-input", 3))
pData(bg) = data.frame(id=sampleNames(bg), group=group)
# Perform differential expression (DE) analysis with no filtering
results transcripts = stattest(bg, feature="transcript", covariate="group", getFC=TRUE,
meas="FPKM")
results_genes = stattest(bg, feature="gene", covariate="group", getFC=TRUE,
meas="FPKM")
results_genes = merge(results_genes, bg_gene_names, by.x=c("id"), by.y=c("gene_id"))
# Filter low-abundance genes.
bg_filt = subset(bg, "rowVars(texpr(bg)) > 1", genomesubset=TRUE)
# Load all attributes including gene name
bg_filt_table = texpr(bg_filt , 'all')
bg_filt_gene_names = unique(bg_filt_table[, 9:10])
# Perform DE analysis now using the filtered data
results transcripts = stattest(bg filt, feature="transcript", covariate="group",
getFC=TRUE, meas="FPKM")
results_genes = stattest(bg_filt, feature="gene", covariate="group", getFC=TRUE,
meas="FPKM")
results_genes = merge(results_genes, bg_filt_gene_names, by.x=c("id"),
by.y=c("gene_id"))
# Identify the significant genes with p-value < 0.05</pre>
sig transcripts = subset(results transcripts, results transcripts$pval < 0.05)</pre>
sig_genes = subset(results_genes, results_genes$pval < 0.05)</pre>
# Visualization1: view the range of values and general distribution of FPKM values
FPKM_dist = function(gene_expression) {
  log_fpkm = log2(gene_expression+1)
 fpkm_long = gather(data.frame(log_fpkm), "Sample", "log2(FPKM+1)")
  index table = pData(bg)
  index_table$id = paste("FPKM.", index_table$id, sep='')
 colnames(index table) = c("Sample", "Group")
 fpkm_long = left_join(fpkm_long, index_table, by = "Sample")
 p = ggplot(fpkm_long, aes(x=Sample, y=`log2(FPKM+1)`, fill = Group)) +
    geom boxplot() + coord flip() +
    labs(title = "log2(FPKM+1) for Each Sample") +
   theme bw() +
    theme(plot.title = element text( size=18, vjust=0.5, hjust=0.5),
          axis.title.x = element_text(size=15, vjust=0.5),
          axis.title.y = element_text( size=15, vjust=0.5),
          axis.text.x = element_text(size=12, colour="black"),
```

```
axis.text.y = element text(size=12, colour="black"),
          legend.text = element text(size=12, colour="black"),
          legend.title = element text(size=12, colour="black"),
          plot.caption = element_text(size=12, colour="gray75", face="italic", hjust =
1, vjust = 1))
 return(p)
p1 = FPKM_dist(gene_expression)
# Visualization2: heat map of differential expression
heatmap = function(gene expression, results genes){
  library(RColorBrewer)
 Colors=brewer.pal(11, "Spectral")
 results_genes[,"de"] = log2(results_genes[,"fc"])
  sigpi = which(results genes[,"pval"]<0.05)</pre>
  topn = order(abs(results_genes[sigpi,"fc"]), decreasing=TRUE)[1:25]
  topn = order(results genes[sigpi, "qval"])[1:25]
  sigp = results_genes[sigpi,]
  sigde = which(abs(sigp[, "de"]) >= 2)
  sig tn de = sigp[sigde,]
 mydist=function(c) {dist(c,method="euclidian")}
 myclust=function(c) {hclust(c,method="average")}
 main title="Heatmap of Differential Expression"
 par(cex.main=0.8)
  sig_genes_de=sig_tn_de[,"id"]
 sig_gene_names_de=sig_tn_de[,"gene_name"]
 data=log2(as.matrix(gene_expression[as.vector(sig_genes_de),])+1)
  p = heatmap.2(data,
                 hclustfun=myclust,
                 distfun=mydist,
                 na.rm = TRUE,
                 scale="none",
                 dendrogram="both",
                 margins=c(7,5),
                 ROWV=TRUE, Colv=TRUE,
                 symbreaks=FALSE, key=TRUE, symkey=FALSE,
                 density.info="none", trace="none",
                 main=main title,
                 cexRow=0.8, cexCol=0.8,
                 labRow=sig gene names de,
                 # col=rev(heat.colors(75)),
                 col=Colors)
  return(p)
p2 = heatmap(gene expression, results genes)
```

A barplot shows the range and general distribution of FPKM values:



A heat map shows the significant differential expressed genes among all samples:



Note that pData should hold a data frame of phenotype information for the samples in the experiment, and be added during ballgown object construction. It can also be added later.

Also note that you can remove all transcripts with a low variance across the samples before differential analysis.

Results

DE results for dataset GSE93676:

```
# To see the first few lines of significant genes and transcripts:
head(sig transcripts)
       feature id
                           fc
                                    pval
                                               qval
# 4 transcript 40 1.237362e-02 0.047948681 0.7342077
# 13 transcript 68 3.643648e-13 0.041800485 0.7342077
# 35 transcript 133 2.283918e+17 0.009168513 0.7342077
# 51 transcript 167 6.634653e-10 0.034473953 0.7342077
# 70 transcript 244 2.241706e-16 0.048610761 0.7342077
# 76 transcript 276 2.637682e-18 0.002536443 0.7342077
head(sig genes)
             id feature
                                fc
                                         pval
                                                    qval gene_name
# 1
       100128071 gene 1.445238e-59 0.04327419 0.7012609
                                                          FAM229A
# 16
          119710 gene 1.378979e-51 0.01122029 0.6171722 C11orf74
            6289 gene 1.718444e-71 0.01229523 0.6171722
                                                             SAA2
# 57
# 74 MSTRG.10000 gene 1.942633e+58 0.01965612 0.6250552
                                                          NGLY1
# 97 MSTRG.10028 gene 5.764237e+05 0.04138674 0.6962953 PDCD6IP
# 141 MSTRG.10096 gene 1.998718e-66 0.02531592 0.6422143
                                                           PTH1R
```

DE results for dataset GSE47217:

```
head(sig_transcripts)
        feature id
                           fc
                                     pval
# 57 transcript 479 0.9339480 0.036290263 0.7220780
# 66 transcript 522 0.4784699 0.002714966 0.5205885
# 86 transcript 685 0.8718279 0.008432731 0.5794108
# 107 transcript 847 0.8360476 0.031839607 0.7176861
# 117 transcript 908 1.0784123 0.004445226 0.5428939
# 137 transcript 1051 0.4439950 0.036100181 0.7220780
head(sig_genes)
             id feature
                              fc
                                       pval
                                                qval gene name
# 1 MSTRG.1000
                 gene 1.0657124 0.04960470 0.5469039 Ppp1r15b
# 36 MSTRG.10096 gene 0.9003980 0.03641091 0.5417405
                                                         Lpin2
# 55 MSTRG.10153 gene 0.9036117 0.02278910 0.5227679
                                                         Birc6
                 gene 0.9036117 0.02278910 0.5227679
# 56 MSTRG.10153
# 85 MSTRG.1024 gene 1.2618979 0.03277910 0.5417405 Tmem183a
```

M3. Site Detection - Host

Peak Calling (exomePeak2)

<u>exomePeak2</u> is an R/Bioconductor package which provides bias-aware quantification and peak detection for Methylated RNA immunoprecipitation sequencing data (MeRIP-Seq) [10]. We are going to use this package for peak calling to predict significant methylation sites.

Installation

```
if(!requireNamespace("BiocManager", quietly = TRUE))
  install.packages("BiocManager")
BiocManager::install("exomePeak2")
```

Peak Calling

```
library(exomePeak2)
set.seed(1)
root = "/path/to/homo result"
setwd(root)
f1 = file.path(root, "SRR5978834_sorted.bam")
f2 = file.path(root, "SRR5978835_sorted.bam")
f3 = file.path(root, "SRR5978836 sorted.bam")
IP\_BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR5978827_sorted.bam")
f2 = file.path(root, "SRR5978828_sorted.bam")
f3 = file.path(root, "SRR5978829_sorted.bam")
INPUT_BAM = c(f1, f2, f3)
exomePeak2(bam ip = IP BAM,
           bam input = INPUT BAM,
           genome = "hg19",
           library_type = "1st_strand",
           paired_end = FALSE)
```

```
library(exomePeak2)
set.seed(1)
```

An output folder named <code>exomePeak2_output</code> will be created in the working directory containing. The most important two files "Mod.bed" and "Mod.csv" will be used in further analysis.

```
- exomePeak2_output
- LfcGC.pdf
- RunInfo.txt
- Mod.bed
- Mod.csv
- Mod.rds
- ADDInfo
- ADDInfo_SizeFactors.csv
- ADDInfo_GLM_allDesigns.csv
- ADDInfo_ReadsCount.csv
- ADDInfo_RPKM.csv
```

Motif Discovery (Homer)

<u>Homer</u> is a software for motif discovery and next-gen sequencing analysis [11]. We are going to use this tool to find enriched m6A motifs within peaks that has been found.

Install Homer

```
# Download and install following the instruction:
http://homer.ucsd.edu/homer/introduction/install.html
$ mkdir homer
$ cd homer
$ wget http://homer.ucsd.edu/homer/configureHomer.pl
$ perl configureHomer.pl -install

# Append to PATH environment variable
$ export PATH=$PATH:/path/to/homer/.//bin/

# Verify installation
$ findMotifs.pl
```

Find Motifs

```
$ findMotifsGenome.pl Mod.bed /path/to/hg19_genome.fa /path/to/MotifOutput -rna -p 10 -
len 5,6
```

Note that Homer can analyze strand-specific genomic regions for motifs by running findMotifsGenome.pl with an rna option. Also note that filtering out the peaks longer than 1000bp can improve the reliability of motifs found by Homer.

The figure below shows the enriched motifs in peaks on hg19 transcripts.

Rank	Motif	P-value	log P-pvalue	% of Targets	% of Background
1	UCGAC	1e-314	-7.239e+02	66.02%	44.91%
2	ACUACC	1e-89	-2.069e+02	51.22%	40.05%
3	UUCGGA EXE	1e-88	-2.030e+02	60.04%	48.90%

The figure below shows the enriched motifs in peaks on mm10 transcripts.

Rank	Motif	P-value	log P-pvalue	% of Targets	% of Background
1	EGGACE	1e-268	-6.186e+02	63.42%	45.31%
2	CAUCGE	1e-72	-1.674e+02	28.97%	21.01%
3	SGASSA	1e-70	-1.620e+02	44.26%	35.29%

Visualization of Reads (IGV)

The Integrative Genomics Viewer (IGV) is a high-performance viewer that efficiently handles large heterogeneous data sets [13]. We are going to use this tool to visualize aligned reads on genome and the differential methylation sites.

Visualization of Aligned Reads

Upload BAM and its index (.bai) file to IGV web application and zoom in to see the aligned reads.





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Visualization of Methylation Sites

1. Install IGV

```
# Java of version 11 is required.
$ java -version

# Download and install from http://software.broadinstitute.org/software/igv/download
$ wget https://data.broadinstitute.org/igv/projects/downloads/2.9/IGV_2.9.2.zip
$ unzip IGV_2.9.2.zip
$ cd IGV_2.9.2
$ nohup bash igv.sh

# Verify installation
$ igvtools version
```

2. Generate TDF

```
#!/bin/bash
Data="/path/to/homo_result"
Output="/path/to/homo_igv"

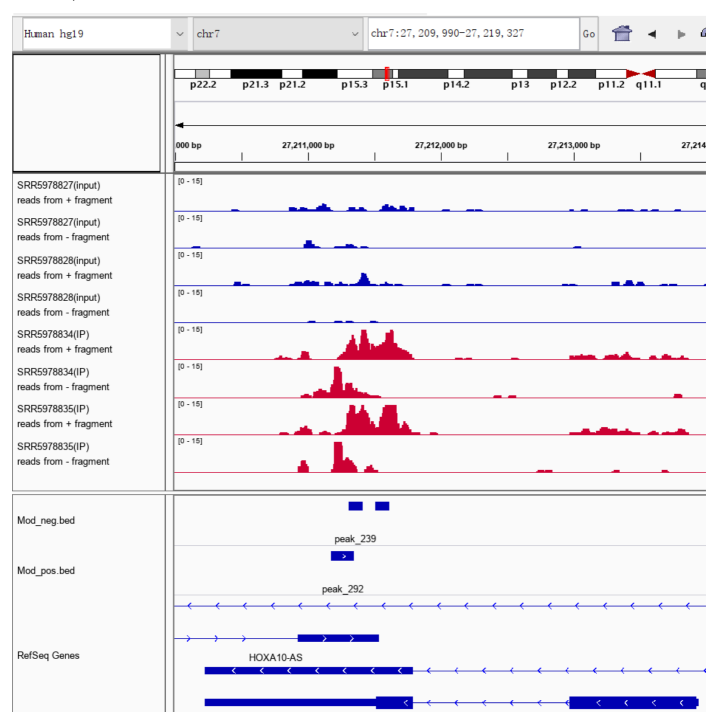
for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
igvtools count --strands read -z 5 -w 10 -e 0 $Data/${s}_sorted.bam $Output/${s}.tdf
homo_genome.fa
wait
done
```

```
#!/bin/bash
Data="/path/to/mm10_result"
Output="/path/to/mm10_igv"

for s in SRR866997 SRR866998 SRR866999 SRR867000 SRR867001 SRR867002 SRR866991
SRR866992 SRR866993 SRR866994 SRR866995 SRR866996
do
igvtools count --strands read -z 5 -w 10 -e 0 $Data/${s}_sorted.bam $Output/${s}.tdf
mm10_genome.fa
wait
done
```

3. Run IGV

The generated TDF files and BED file can then be visualized using IGV browser. As clearly shown in the figure below zooming in a region of an antisense gene, the exomePeak2 has the ability to distinguish methylation sites on specific strands.



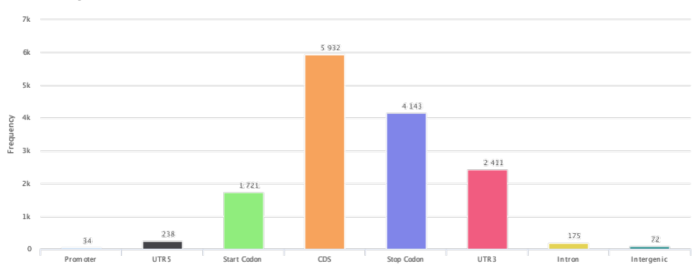
RNA Annotation (RNAmod)

<u>RNAmod</u> is a convenient web-based platform for the meta-analysis and functional annotation of modifications on mRNAs [14].

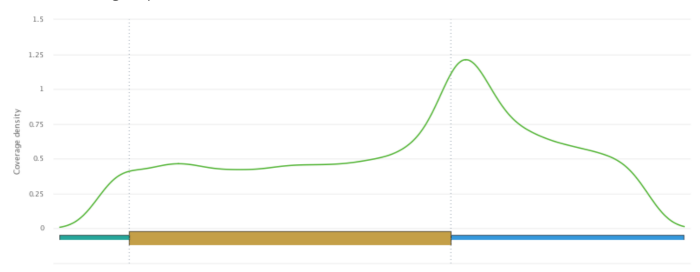
To obtain modification site information, click on "Single case" in the <u>Home page</u>, upload the BED file generated from exomePeak2 package, and submit the job. Then you will receive a job ID and will use it to query the job status and get the results in the <u>result page</u>. You can either view the results on browser or download the zip file to the local place.

Some of the figures about modification site information are displayed below:

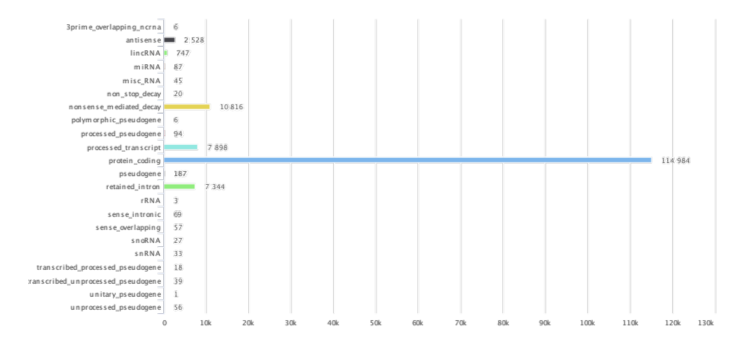
• Peaks gene features distribution:



• mRNA metagene plot



• Gene types distribution



M4. Site Detection - Virus

Peak Calling (exomePeak2)

We are going to use exomePeak2 for peak calling to find enriched m6A sites on HHV8 transcripts. The BED file as output could be used for further analysis.

```
library(exomePeak2)
set.seed(1)
root = "/path/to/hhv8_result"
setwd(root)
f1 = file.path(root, "SRR5978834 sorted.bam")
f2 = file.path(root, "SRR5978835 sorted.bam")
f3 = file.path(root, "SRR5978836_sorted.bam")
IP_BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR5978827_sorted.bam")
f2 = file.path(root, "SRR5978828 sorted.bam")
f3 = file.path(root, "SRR5978829 sorted.bam")
INPUT BAM = c(f1, f2, f3)
GENE_ANNO_GTF = file.path("/path/to/sequence.gff3")
exomePeak2(bam ip = IP BAM,
           bam input = INPUT BAM,
           gff dir = GENE ANNO GTF,
           library_type = "1st_strand",
           paired_end = FALSE)
```

An output folder named exomePeak2 output will be created in the working directory containing:

```
- exomePeak2_output
- LfcGC.pdf
- RunInfo.txt
- Mod.bed
- Mod.csv
- Mod.rds
- ADDInfo
- ADDInfo_SizeFactors.csv
- ADDInfo_GLM_allDesigns.csv
- ADDInfo_ReadsCount.csv
- ADDInfo_RPKM.csv
```

Visualization of Reads (IGV)

Visualization of Aligned Reads

Upload the following files to IGV web application.

- HHV8 virus genome (.fa) and its index (.fai) to "Genome".
- BAM and its index (.bai) file to "Track".





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Visualization of Methylation Sites

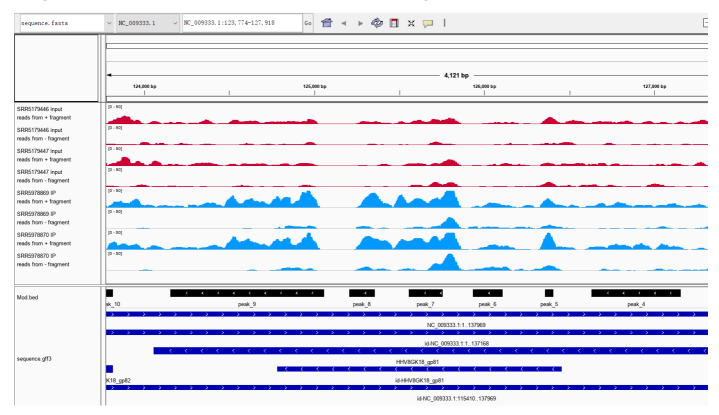
1. Generate TDF

```
#!/bin/bash
Data="/path/to/hhv8_result"
Output="/path/to/hhv8_igv"

for s in SRR5978827 SRR5978828 SRR5978829 SRR5978834 SRR5978835 SRR5978836 SRR5978869
SRR5978870 SRR5978871 SRR5179446 SRR5179447 SRR5179448
do
igvtools count -z 5 -w 10 -e 0 $Data/${s}_sorted.bam $Output/${s}.tdf hhv8_genome.fa
wait
done
```

2. Run IGV

The generated TDF files and BED file can then be visualized using IGV browser.



M5. Differential Methylation

m6A-seq Data Analysis (exomePeak2)

Here we are going to use exomePeak2 for peak calling and differential methylation analysis. The BED file as output will be used for further analysis.

```
library(exomePeak2)
set.seed(1)
root = "/path/to/homo result"
setwd(root)
f1 = file.path(root, "SRR5978834_sorted.bam")
f2 = file.path(root, "SRR5978835_sorted.bam")
f3 = file.path(root, "SRR5978836_sorted.bam")
IP BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR5978827_sorted.bam")
f2 = file.path(root, "SRR5978828_sorted.bam")
f3 = file.path(root, "SRR5978829_sorted.bam")
INPUT_BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR5179446_sorted.bam")
f2 = file.path(root, "SRR5179447 sorted.bam")
f3 = file.path(root, "SRR5179448_sorted.bam")
TREATED_INPUT_BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR5978869_sorted.bam")
f2 = file.path(root, "SRR5978870_sorted.bam")
f3 = file.path(root, "SRR5978871 sorted.bam")
TREATED_IP_BAM = c(f1, f2, f3)
exomePeak2(bam_ip = IP_BAM,
           bam_input = INPUT_BAM,
           bam_treated_input = TREATED_INPUT_BAM,
           bam_treated_ip = TREATED_IP_BAM,
           genome = "hg19",
           library_type = "1st_strand",
           paired_end = FALSE)
```

```
library(exomePeak2)
set.seed(1)
root = "/path/to/mm10_result"
setwd(root)

f1 = file.path(root, "SRR866997_sorted.bam")
f2 = file.path(root, "SRR866999_sorted.bam")
f3 = file.path(root, "SRR867001_sorted.bam")
IP_BAM = c(f1,f2,f3)

f1 = file.path(root, "SRR866998_sorted.bam")
f2 = file.path(root, "SRR8667000_sorted.bam")
```

```
f3 = file.path(root, "SRR867002 sorted.bam")
INPUT BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR866992 sorted.bam")
f2 = file.path(root, "SRR866994_sorted.bam")
f3 = file.path(root, "SRR866996_sorted.bam")
TREATED_INPUT_BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR866991 sorted.bam")
f2 = file.path(root, "SRR866993_sorted.bam")
f3 = file.path(root, "SRR866995 sorted.bam")
TREATED IP BAM = c(f1, f2, f3)
exomePeak2(bam_ip = IP_BAM,
           bam input = INPUT BAM,
           bam_treated_input = TREATED_INPUT_BAM,
           bam treated ip = TREATED IP BAM,
           genome = "mm10",
           library_type = "1st_strand",
           paired_end = FALSE)
```

An output folder named exomePeak2_output will be created in the working directory containing:

```
- exomePeak2_output
- LfcGC.pdf
- RunInfo.txt
- DiffMod.bed
- DiffMod.csv
- DiffMod.rds
- ADDInfo
- ADDInfo
- ADDInfo_SizeFactors.csv
- ADDInfo_GLM_allDesigns.csv
- ADDInfo_ReadsCount.csv
- ADDInfo_RPKM.csv
```

Distribution of m6A sites (MetaTX)

We are going to use MetaTX to visualize the distribution of methylation sites.

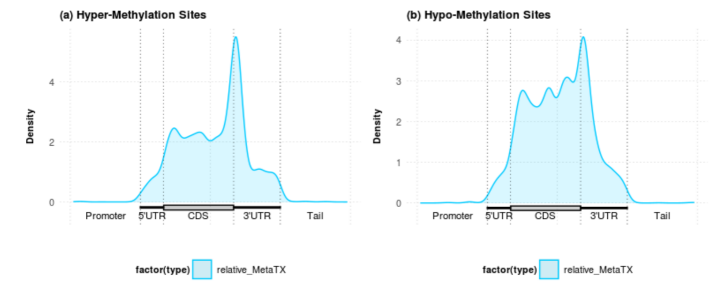
Visualization of the Distribution of Peaks

The following code produces separate figures of the distribution of hyper-methylation sites and hypomethylation sites.

```
# Load libraries
```

```
library(MetaTX)
library(rtracklayer)
library(readr)
library(bedr)
library(genomation)
library(GenomicRanges)
# Import BED file from exomePeak2
file = "/path/to/exomePeak2 output peakcalling 1strand/Mod.bed"
file1 = "/path/to/exomePeak2 output peakcalling 1strand/Mod.csv"
data csv = read csv(file1)
gr_obj = import(file)
# Separate by hyper and hypo methylation sites
data1 = gr obj[which(data csv$DiffModLog2FC > 0),]
data2 = gr_obj[which(data_csv$DiffModLog2FC < 0),]</pre>
# convert to bed files
df1 = data.frame(seqnames=seqnames(data1),
                  starts=start(data1),
                  ends=end(data1),
                  names=elementMetadata(data1)$name,
                  scores=elementMetadata(data1)$score,
                  strands=strand(data1))
df2 = data.frame(seqnames=seqnames(data2),
                  starts=start(data2),
                  ends=end(data2),
                  names=elementMetadata(data2)$name,
                  scores=elementMetadata(data2)$score,
                  strands=strand(data2))
write.table(df1, file="Mod_metaTX_diff_pos.bed", quote=F, sep="\t", row.names=F,
col.names=F)
write.table(df2, file="Mod_metaTX_diff_neg.bed", quote=F, sep="\t", row.names=F,
col.names=F)
# Import separated bed files
file_pos = "Mod_metaTX_diff_pos.bed"
file neg = "Mod metaTX diff neg.bed"
gr_obj_pos = import(file_pos)
gr obj neg = import(file neg)
gr_obj_pos = resize(gr_obj_pos, width = 1, fix = "center")
gr_obj_neg = resize(gr_obj_neg, width = 1, fix = "center")
# Download information about mRNA components
txdb = TxDb.Hsapiens.UCSC.hg19.knownGene
cds by tx0 1 = cdsBy(txdb, "tx")
fiveUTR_tx0_1 = fiveUTRsByTranscript(txdb,use.names=FALSE)
threeUTR_tx0_1 = threeUTRsByTranscript(txdb,use.names=FALSE)
```

```
# Map peaks to the RNA model
remap_results_m6A_1 = remapCoord(features = gr_obj_pos, txdb = txdb, num_bin = 10,
includeNeighborDNA = TRUE, cds by tx0 = cds by tx0 1, fiveUTR tx0 = fiveUTR tx0 1,
                                  threeUTR_tx0 = threeUTR_tx0_1)
# Plot 1
p1 = metaTXplot(remap_results_m6A_1,
                  num bin
                                       = 10,
                  includeNeighborDNA
                                       = TRUE,
                                      = c(1, 3, 2, 3),
                  relativeProportion
                  title = '(a) Hyper-Methylation Sites',
                  legend = 'relative',
                  type = 'relative'
)
# Map peaks to the RNA model
remap_results_m6A_2 = remapCoord(features = gr_obj_neg, txdb = txdb, num_bin = 10,
includeNeighborDNA = TRUE, cds by tx0 = cds by tx0 1, fiveUTR tx0 = fiveUTR tx0 1,
                                  threeUTR_tx0 = threeUTR_tx0_1)
# Plot 2
p2 = metaTXplot(remap_results_m6A_2,
                  num_bin
                                       = 10,
                  includeNeighborDNA
                                       = TRUE,
                  relativeProportion
                                      = c(1, 3, 2, 3),
                  title = '(b) Hypo-Methylation Sites',
                  legend = 'relative',
                  type = 'relative'
# Plot 1 + 2
ggdraw() +
 draw_plot(p1, 0, 0, .5, 1) +
 draw_plot(p2, .5, 0, .5, 1)
```



Report Isoform Probabilities

MetaTX also provides a function for returning the probabilities of a particular feature being located on different isoforms.

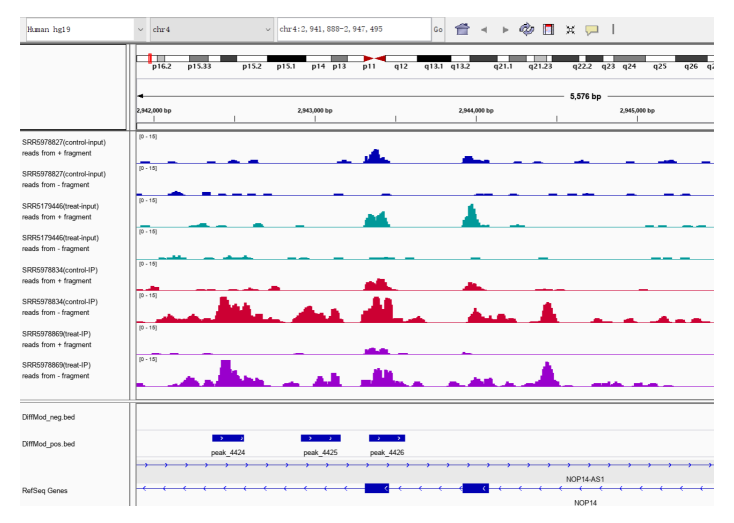
```
isoform_probs <- isoformProb(remap_results_m6A_1, num_bin = 10, includeNeighborDNA =
TRUE, lambda = 2)
write.csv(isoform_probs, "isoform_probs.csv")</pre>
```

Here are the first few rows of the outputs.

index_trans (double)	index_methyl (double)	seqnames (character)	methyl_pos (double)	strand (character)	trans_ID (double)	isoform_prob (double)		
1	1	chr19	58867266	-	70456	0.00000000		
2	1	chr19	58867266	-	70457	0.51380127		
3	1	chr19	58867266	-	70458	0.48619873		
4	2	chr20	43269048	-	72132	0.00000000		
5	3	chr18	25532067	-	65378	0.43630632		
6	3	chr18	25532067	-	65379	0.56369368		
7	4	chr3	101396111	+	14200	1.00000000		
8	5	chrX	119387453	+	76492	0.45666710		
9	5	chrX	119387453	+	76493	0.47472665		

Visualization of Reads (IGV)

Here we use the IGV tool to visualize reads and peaks comparing control and experimental groups. This task can be simply done by uploading the generated TDF files and BED file onto the IGV browser.



GO Enrichment Analysis (DAVID)

The **D**atabase for **A**nnotation, **V**isualization and **I**ntegrated **D**iscovery (**DAVID**) is a website providing a comprehensive set of functional annotation tools for investigators to understand biological meaning behind large list of genes [15]. We are going to use this tool to discover enriched gene functions.

Prepare and Upload Gene Lists

From the output file ("DiffMod.csv") generated from exomePeak2, we remove duplicated values in "geneID" column and copy all the unique IDs to a txt file ("geneID_diff.txt").

Upload txt files to DAVID website with Identifier as "ENTREZ_GENE_ID", species as "Homo sapiens", and "Gene List" selected. Submit all lists and wait for results.

Analyze Results

Open "Functional Annotation Chart" and click on "Download File" to download the txt file containing results.



DAVID Bioinformatics Resources 6.8

Laboratory of Human Retrovirology and Immunoinformatics (LHRI)

Functional Annotation Chart

Current Gene List: geneID_diffmod Current Background: Homo sapiens 330 DAVID IDs

■ Options

130 chart records

Rerun Using Options | Create Sublist

Material Problems Download File

Help and Manual

Sublist	<u>Category</u>	<u>Ierm</u>	¢RT	Genes	Count	<u>%</u> ♦ PV	alue	<u>Benjamini</u>	
	UP_KEYWORDS	Alternative splicing	RT		163	49.4 7	.4E-	1.2E-4	
	UP_KEYWORDS	Phosphoprotein	<u>RT</u>		126	38.2 4	.3E-	1.8E-2	
	UP_KEYWORDS	Cytoplasm	<u>RT</u>		81	24.5 3	.2E-	3.0E-2	
	GOTERM_BP_DIRECT	interstrand cross-link repair	<u>RT</u>	=	6	1.8 3	.6E-	3.8E-1	
	UP_SEQ_FEATURE	splice variant	<u>RT</u>		121	36.7 ⁴ / ₄	.1E-	3.6E-1	
	GOTERM_CC_DIRECT	nucleoplasm	<u>RT</u>		53	16.1 6	.2E-	9.8E-2	
	GOTERM_CC_DIRECT	Fanconi anaemia nuclear complex	<u>RT</u>	i	4	1.2 7	.0E-	9.8E-2	
	GOTERM_BP_DIRECT	negative regulation of protein complex assembly	<u>RT</u>	Ē	4	1.2 3	.2E-	6.1E-1	
	UP_KEYWORDS	Nucleus	<u>RT</u>		83	25.2 3	.8E-	1.3E-1	
	UP_KEYWORDS	Protein transport	<u>RT</u>	=	17	5.2 2	.3E-	1.3E-1	
	GOTERM_CC_DIRECT GOTERM_BP_DIRECT UP_KEYWORDS	Fanconi anaemia nuclear complex negative regulation of protein complex assembly Nucleus	RT RT		4 4 83	1.2 7 4 1.2 1 3 25.2 1	.0E- .2E- .8E-	•	

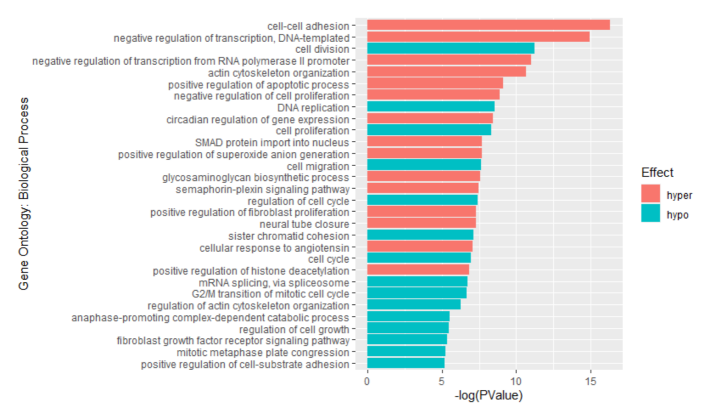
Import txt file into R, analyze results and display in figures.

```
library(readr)
library(dplyr)
chart = read_tsv("chart_diff.txt")
# generate a figure for differential expressed genes
generateFigure = function(chart, num, term = "GOTERM_BP_DIRECT"){
  p = selectPvalue(chart)
  frame = as.data.frame(chart %>%
    filter(Category == term) %>%
   select(c("Term", "%", "PValue")) %>%
   rename("Ratio"=`%`) %>%
   mutate(Term = as.factor(gsub("^.*?~", "",Term)),
           Ratio = Ratio / 100))[1:30,] # make sure no less than 30 terms in total
  fig2 = frame %>%
    ggplot(aes(x=reorder(Term, -PValue),y=-log(PValue),fill = Ratio)) +
    geom_bar(stat="identity") +
   coord_flip() +
   xlab("Gene Ontology: Biological Process")
  return(fig2)
}
generateFigure(chart, 30, term)
```

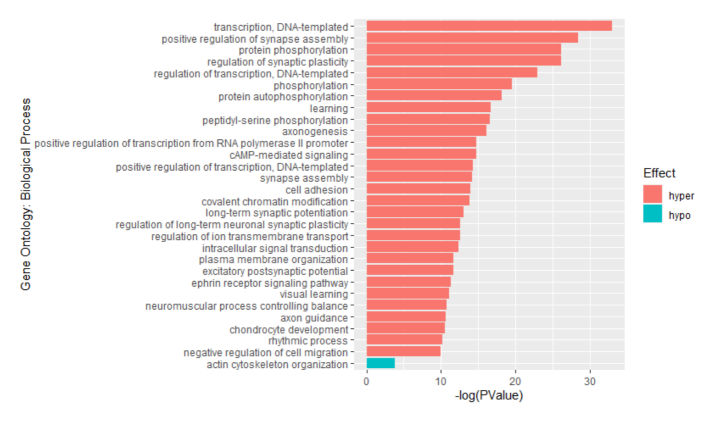
```
# generate a figure for hyper/hypo genes
chartp = read tsv("pos david mm.txt")
chartn = read_tsv("neg_david_mm.txt")
chartp = chartp %>% mutate(Effect = "hyper")
chartn = chartn %>% mutate(Effect = "hypo")
charts = rbind(chartp, chartn)
generateFigure hyper hype = function(charts, term = "GOTERM BP DIRECT"){
 print(charts %>%
          filter(Category == term & Effect == 'hyper') %>%
          select(c("Term", "%", "PValue", "Effect")) %>%
          rename("Ratio"=`%`) %>%
         mutate(Term = as.factor(gsub("^.*?~", "",Term)),
                 Ratio = Ratio / 100) %>% nrow())
 frame_p = as.data.frame(charts %>%
   filter(Category == term & Effect == 'hyper') %>%
   select(c("Term", "%", "PValue", "Effect")) %>%
   rename("Ratio"=`%`) %>%
   mutate(Term = as.factor(gsub("^.*?~", "",Term)),
           Ratio = Ratio / 100))[1:29,]
 print(charts %>%
          filter(Category == term & Effect == 'hypo') %>%
          select(c("Term", "%", "PValue", "Effect")) %>%
          rename("Ratio"=`%`) %>%
         mutate(Term = as.factor(gsub("^.*?~", "",Term)),
                 Ratio = Ratio / 100) %>% nrow())
  frame n = as.data.frame(charts %>%
     filter(Category == term & Effect == 'hypo') %>%
     select(c("Term", "%", "PValue", "Effect")) %>%
    rename("Ratio"=`%`) %>%
    mutate(Term = as.factor(gsub("^.*?~", "",Term)),
           Ratio = Ratio / 100))[1,]
  frame = rbind(frame p, frame n)
  fig2 = frame %>%
   ggplot(aes(x=reorder(Term, -PValue), y=-log(PValue), fill = Effect)) +
   geom bar(stat="identity") +
   coord flip() +
   xlab("Gene Ontology: Biological Process")
 return(fig2)
}
```

generateFigure hyper hypo(charts)

Enriched biological processes regulated by differential methylated genes on hg19 genome.



Enriched biological processes regulated by differential methylated genes on mm10 genome.

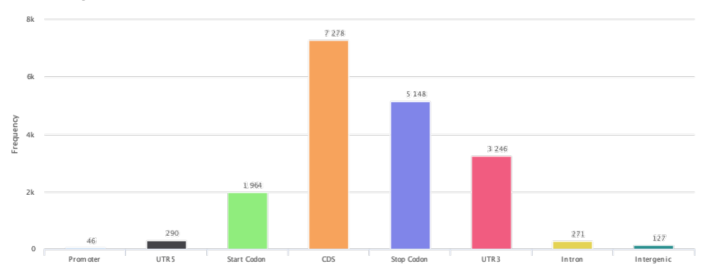


RNA Annotation (RNAmod)

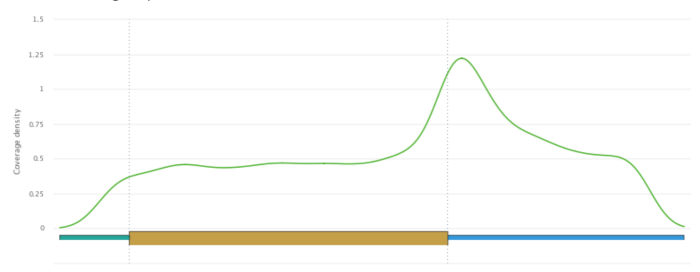
To obtain modification site information, click on "Single case" in the RNAmod <u>Home page</u>, upload the BED file generated from exomePeak2 package, and submit the job. Then you will receive a job ID and will use it to query the job status and get the results in the <u>result page</u>. You can either view the results on browser or download the zip file to the local place.

Some of the figures about modification site information are displayed below:

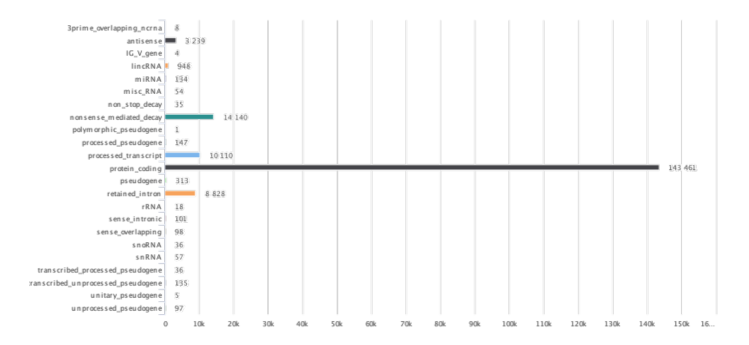
• Peaks gene features distribution:



• mRNA metagene plot



• Gene types distribution



M6. Reference based analysis

exomePeak2

Here we use exomePeak2 to conduct reference based quantification and differential analysis.

Download and Convert Basic Site Information

Download single base RNA modification annotation of m6A on human genome from m6A-Atlas database, which should be tabular data in a txt file. Convert tabular data to genomic ranges by:

```
library(readr)
library(GenomicRanges)
# Downloaded from m6A-Atlas database
my.file="/path/to/home/tangyujiao/big/download/m6A H.sapiens basical information.txt"
# Load txt file
m6A basic info <- read table2(my.file, col names = FALSE)
m6A_basic_info <- m6A_basic_info[, c(1:11)]</pre>
colNames <- c("ID", "chr", "start", "end", "num1", "strand", "LOC", "ENS", "RNA", "Gene",
colnames(m6A_basic_info) <- colNames</pre>
# Concert to Grange object
mod_annot <- makeGRangesFromDataFrame(m6A_basic_info,</pre>
                          keep.extra.columns=FALSE,
                          ignore.strand=FALSE,
                          seginfo=NULL,
                          segnames.field="chr",
                          start.field="start",
                          end.field="end",
                          strand.field="strand",
```

```
starts.in.df.are.Obased=FALSE)

# Save Grange to rds
saveRDS(mod_annot, "/path/to/mod_annot.rds")
```

Reference-based Analysis

```
library(exomePeak2)
set.seed(1)
root = "/path/to/homo_result"
setwd(root)
f1 = file.path(root, "SRR5978834_sorted.bam")
f2 = file.path(root, "SRR5978835_sorted.bam")
f3 = file.path(root, "SRR5978836 sorted.bam")
IP BAM = c(f1, f2, f3)
f1 = file.path(root, "SRR5978827_sorted.bam")
f2 = file.path(root, "SRR5978828_sorted.bam")
f3 = file.path(root, "SRR5978829_sorted.bam")
INPUT_BAM = c(f1, f2, f3)
f2 = "/path/to/mod annot.rds"
MOD ANNO GRANGE <- readRDS(f2)</pre>
exomePeak2(bam_ip = IP_BAM,
           bam_input = INPUT_BAM,
           genome = "hg19",
           library_type = "1st_strand",
           paired end = FALSE,
           mod_annot = MOD_ANNO_GRANGE)
```

An output folder named exomePeak2_output will be created in the working directory containing:

```
- exomePeak2_output
- LfcGC.pdf
- RunInfo.txt
- Mod.bed
- Mod.csv
- Mod.rds
- ADDInfo
- ADDInfo_SizeFactors.csv
- ADDInfo_GLM_allDesigns.csv
- ADDInfo_ReadsCount.csv
- ADDInfo_RPKM.csv
```

Appendix

Construct BSgenome Object

exomePeak2 requires BSgenome object of matched species. However, when there is no available BSgenome object, we have to manually construct one. Therefore, this page provides instructions for constructing BSgenome object from scratch by using *Zea mays* (Corn) genome as an example.

1. Download Genome FASTA

Download *Zea mays* (Corn) reference genome "AGPv4" from <u>Illumina iGenome</u>, and find genome fasta file.

```
$ wget http://igenomes.illumina.com.s3-website-us-east-
1.amazonaws.com/Zea_mays/Ensembl/AGPv4/Zea_mays_Ensembl_AGPv4.tar.gz
$ tar -xvzf Zea_mays_Ensembl_AGPv4.tar.gz
```

2. Convert fasta to 2bit file format

Download "faToTwoBit" tool by

```
$ rsync -aP \
    rsync://hgdownload.soe.ucsc.edu/genome/admin/exe/linux.x86_64/faToTwoBit ./
```

Convert file format

```
$ mkdir
/home/zhen.di/corn/Zea_mays/Ensembl/AGPv4/Sequence/WholeGenomeFasta/preparation/
$ cd
/home/zhen.di/corn/Zea_mays/Ensembl/AGPv4/Sequence/WholeGenomeFasta/preparation/
$ cp ../genome.fa zmAGPv4.fa
$ faToTwoBit zmAGPv4.fa zmAGPv4.2bit
```

3. Extract chromosome names

```
$ less -S zmAGPv4.fa | grep ">" |awk '{print $1}' | sed 's/^>//g' >
zmAGPv4.chromName.txt
```

Find chromosome name by

```
$ cat zmAGPv4.chromName.txt
```

4. Prepare seed file "BSgenome.Zmays.Ensemble.zmAGPv4-seed" in the same directory

```
Package: BSgenome.Zmays.Ensemble.zmAGPv4

Title: Genome sequences for Zea mays (Ensemble AGPv4)

Description: A BSgenome package containing the full genome sequences for Zea mays (Maize) as provided by Ensemble (B73 AGPv4, Sept. 2020) and stored in Biostrings objects.

Version: 1.0
```

```
organism: Zea mays
common_name: Maize
provider: Ensemble
provider_version: zmAGPv4
release_date: Sept. 2020
release_name: Maize Genome Sequencing B73 RefGen_v4.0
#source_url: ftp://ftp.ensemblgenomes.org/pub/plants/release-49/fasta/zea_mays/dna/
circ_seqs: c("Mt","Pt")
organism_biocview: Zea_mays
BSgenomeObjname: Zmays
seqs_srcdir:
/home/zhen.di/corn/Zea_mays/Ensembl/AGPv4/Sequence/WholeGenomeFasta/preparation/
seqfiles_suffix: .fa
seqnames: c("10", "1", "2", "3", "4", "5", "6", "7", "8", "9", "MT", "Pt")
seqfile_name: zmAGPv4.2bit
```

Please make sure the information included in the seed file is correct by referring to <u>BSgenomeForge</u> <u>doc</u> and <u>description</u>.

5. Construct BSgenome data package

```
# Construct BSgenome data package in R
library(rtracklayer)
library(BSgenome)
setwd("/home/zhen.di/corn/Zea_mays/Ensembl/AGPv4/Sequence/WholeGenomeFasta/preparat
ion")
forgeBSgenomeDataPkg("BSgenome.Zmays.Ensemble.zmAGPv4-seed")

## The construction completes successfully with messages:

# Creating package in ./BSgenome.Zmays.Ensemble.zmAGPv4

# Loading
'/home/zhen.di/corn/Zea_mays/Ensembl/AGPv4/Sequence/WholeGenomeFasta/preparation//z
mAGPv4.2bit' ... DONE

# Writing sequences to
'./BSgenome.Zmays.Ensemble.zmAGPv4/inst/extdata/single_sequences.2bit' ... DONE
```

6. Construct R package

```
$ R CMD build BSgenome.Zmays.Ensemble.zmAGPv4
```

7. Check whether the tar.gz file has been successfully created by

```
$ R CMD check BSgenome.Zmays.Ensemble.zmAGPv4_1.0.tar.gz
```

You should receive messages showing everything is OK.

8. Install package into R

```
$ R CMD INSTALL BSgenome.Zmays.Ensemble.zmAGPv4_1.0.tar.gz
```

9. Construct TxDb object

```
library(AnnotationHub) # help to build the annotation object
library(biomaRt)
library(GenomicFeatures)
# library(exomePeak2)
library(rtracklayer)
library(BSgenome)
library(BSgenome.Zmays.Ensemble.zmAGPv4)

maize_txdb<- makeTxDbFromBiomart(biomart = "plants_mart",dataset = "zmays_eg_gene",host = "http://plants.ensembl.org")
saveDb(maize_txdb, file="maize_v4.sqlite")
maize_txdb <- loadDb("maize_v4.sqlite")</pre>
```

10. Use TxDb object in exomePeak2 function

```
exomePeak2(bam_ip = c("IP_1.bam","IP_2.bam"),
    bam_input = c("Input_1.bam","Input_2.bam"),
    txdb = maize_txdb,
    bsgenome = BSgenome.Zmays.Ensemble.zmAGPv4,
    paired_end = TRUE)
```

Reference

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- [10] Zhen Wei (2020). exomePeak2: Bias Awared Peak Calling and Quantification for MeRIP-Seq. R package version 1.0.0. [bioc]
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