# Package 'CircSeqAlignTk'

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```
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Description CircSeqAlignTk is designed for end-to-end RNA-Seq data analysis
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     It mainly targets viroids which are composed of 246-401 nt circular RNAs.
     In addition, CircSeqAlignTk implements a tidy interface
     to generate synthetic sequencing data that mimic real RNA-Seq data,
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```
Author Jianqiang Sun [aut, cre] (<a href="https://orcid.org/0000-0002-3438-3199">https://orcid.org/0000-0002-3438-3199</a>), Xi Fu [aut], Wei Cao [aut]
```

Maintainer Jianqiang Sun <sun@biunit.dev>

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CircSeqAlignTk-package

CircSeqAlignTk: A toolkit for end-to-end analysis of RNA-seq data for circular genomes

# Description

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CircSeqAlignTk is designed for end-to-end RNA-Seq data analysis of circular genome sequences, from alignment to visualization. It mainly targets viroids which are composed of 246-401 nt circular RNAs. In addition, CircSeqAlignTk implements a tidy interface to generate synthetic sequencing data that mimic real RNA-Seq data, allowing developers to evaluate the performance of alignment tools and workflows.

# **Details**

Refer to the vignette for an overview of the package, quick start, and detailed usages.

# Author(s)

Maintainer: Jianqiang Sun <sun@biunit.dev> (ORCID)

Authors:

- Xi Fu
- Wei Cao

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#### See Also

Useful links:

- https://github.com/jsun/CircSeqAlignTk
- Report bugs at https://github.com/jsun/CircSeqAlignTk/issues

#### **Examples**

```
browseVignettes("CircSeqAlignTk")
```

align\_reads

Align sequence reads to a genome sequence

#### **Description**

This function aligns sequence reads in a FASTQ file to the reference sequences of a genome.

# Usage

```
align_reads(
  input,
  index,
  output,
  n_threads = 1,
  n_mismatch = 1,
  overwrite = TRUE,
  aligner = c("bowtie2", "hisat2"),
  add_args = NULL
)
```

# **Arguments**

input A path to a FASTQ format file for alignment. index A CircSeqAlignTkRefIndex-class object generated by the build\_index funcoutput A path to a directory for saving the intermediate and final results of alignment. Number of threads to use for aligning reads. n\_threads n\_mismatch Number of allowed mismatches in alignment. overwrite Overwrite the existing files if TRUE. aligner A string to specify the alignment is for alignment. A string of additional arguments to be passed on to the alignment tool directly. add\_args For example, -N 0 -L 22, --no-spliced-alignment -k 10, etc.

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#### **Details**

This function aligns sequence reads in a FASTQ format file in two stages: (i) aligning reads to the type 1 reference sequence (i.e., refseq.t1.fa) and (ii) collecting the unaligned reads and aligning them with the type 2 reference (i.e., refseq.t2.fa). The alignment results are saved as BAM format files in the specified directory with the suffixes \*.t1.bam and \*.t2.bam. The original alignment results may contain mismatches. Hence, filtering is performed to remove the alignment with mismatches over the specified value from the BAM format file. The filtered results of the \*.t1.bam and \*.t2.bam are saved as \*.clean.t1.bam and \*.clean.t2.bam, respectively.

Two alignment tools (Bowtie2 and HISAT2) can be specified for building indexes through the aligner argument. This function first attempts to call the specified alignment tool installed on the operation system directly; however, if the tool is not installed, then the function attempts to call bowtie2\_build or hisat2\_build functions implemented in the Rbowtie2 or Rhisat2 packages for alignment.

#### Value

A CircSeqAlignTkAlign-class object.

#### See Also

CircSeqAlignTkAlign-class

# **Examples**

build\_index

Build indexes of reference sequences for alignment

#### **Description**

This function internally calls Bowtie2 or HISAT2 to build indexes of reference sequences for alignment preparation.

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#### Usage

```
build_index(
   input,
   output = NULL,
   n_threads = 1,
   overwrite = TRUE,
   aligner = c("bowtie2", "hisat2"),
   add_args = NULL
)
```

# **Arguments**

input A path to a FASTA format file containing a reference sequence of a genome for

indexing.

output A path to a directory for saving the reference sequences and indexes.

n\_threads Number of threads to use for aligning reads.

overwrite Overwrite the existing files if TRUE.

aligner A string to specify the alignment for indexing.

add\_args A string of additional arguments to be passed on to the alignment tool directly

(e.g., --quiet).

# Details

This function generates two types of reference sequences from a genome and indexes them in preparation for alignment. The type 1 reference sequence is identical to the sequence provided by the input argument. The type 2 reference sequence is generated by restoring the type 1 reference sequence to a circular RNA and opening the circle at the position opposite to that of type 1. The type 1 and 2 reference sequences are then saved as FASTA format files, refseq.t1.fa and refseq.t2.fa, respectively, under the directory specified by the output argument. Next, the function builds indexes for refseq.t1.fa and refseq.t2.fa.

Two alignment tools (Bowtie2 and HISAT2) can be specified for building indexes through the aligner argument. This function first attempts to call the specified alignment tool installed on the operation system directly; however, if the tool is not installed, then the function attempts to call bowtie2\_build or hisat2\_build functions implemented in the Rbowtie2 or Rhisat2 packages for indexing.

#### Value

A CircSeqAlignTkRefIndex-class object.

#### See Also

CircSeqAlignTkRefIndex-class

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#### **Examples**

calc\_coverage

Calculate alignment coverage

### **Description**

This function calculates alignment coverage according to the read strand and length from alignment results.

# Usage

```
calc_coverage(x)
```

### **Arguments**

Х

A CircSeqAlignTkAlign-class object generated by the align\_reads function.

#### Details

This function calculates alignment coverage from the two BAM files, \*.clean.t1.bam and \*.clean.t2.bam, generated by the align\_reads function. The coverage is then sorted by the strand and length of the aligned reads and summarized into data frames.

# Value

A CircSeqAlignTkCoverage-class object.

#### See Also

```
CircSeqAlignTkAlign-class, CircSeqAlignTkCoverage-class, align_reads
```

CircSeqAlignTkAlign-class

Class to store alignment results

# **Description**

A class to store alignment results, including the paths to FASTQ and BAM format files and the alignment summary. The object belongs to this class is generated by align\_reads function.

#### **Slots**

input\_fastq A path to the query FASTQ format file.

fastq A vector containing the paths to the two FASTQ format files used for alignment to the type 1 and type 2 references, respectively. See align\_reads for how the FASTQ format files are generated.

bam A vector containing the paths to the two BAM format files corresponding to the alignment results of the two FASTQ files shown in the fastq slot, respectively.

clean\_bam A vector containing the paths to the two BAM format files after filtering by number of mismatch from BAM format files shown in bam slot.

stats A data frame containing alignment summary, e.g., number of query reads, aligned reads, and unaligned reads.

reference A CircSeqAlignTkRefIndex-class storing the information of reference for alignment.

#### See Also

CircSeqAlignTkRefIndex-class, align\_reads

 ${\tt CircSeqAlignTkCoverage-class}$ 

Class to save alignment coverage

# **Description**

A class to store the alignment coverage generated by calc\_coverage function.

#### **Slots**

forward A matrix containing the alignment coverage of the forward strand reads. reversed A matrix containing the alignment coverage of the reversed strand reads. figdata A string of adapter sequence.

#### See Also

calc\_coverage

CircSeqAlignTkRefIndex-class

Class to store reference information

#### **Description**

A class to store reference information for alignment. The object belongs to this class is generated by build\_index function.

#### **Slots**

name Reference name. The sequence name written the header of FASTA format file.

seq Reference sequence.

length Length of the reference sequence.

fasta A vector containing the paths to the two FASTA format files of the type 1 and type 2 reference sequences, respectively. See build\_index for how FASTA format files are generated.

index A vector containing the paths to the two reference indexes corresponding to the two FASTA format files stored in the fasta slot, respectively.

cut\_loc The position on the user-given sequence (i.e., the type 1 sequence) to cut for generating the type 2 reference sequence.

#### See Also

build\_index

CircSeqAlignTkSim-class

Class to save information of synthetic reads

# **Description**

A class to store parameters for generating synthetic sequence reads. The object belongs to this class is generated by generate\_reads function.

#### **Slots**

seq A string of a genome sequence, which is used for sampling synthetic sequence reads. adapter A string of adapter sequence.

read\_info A data frame storing the summary information of read generation. It contains the start and end positions of sampling, strand, and nucleotide sequence of each synthetic read.

peak A data frame storing the peaks information of alignment coverage.

coverage A CircSeqAlignTkCoverage-class storing the information of alignment coverage.

fastq A path to FASTQ format file saving the synthetic reads.

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#### See Also

generate\_reads, CircSeqAlignTkCoverage-class

filter\_reads Filter sequence reads in a FASTQ file by length

# **Description**

This function removes sequence reads with lengths outside the specified range from the FASTQ file.

# Usage

```
filter_reads(input, output, read_lengths = seq(21, 24), overwrite = TRUE)
```

# **Arguments**

input A path to a FASTQ file targeted for filtering.

output A path to save the filtered reads in FASTQ format.

read\_lengths A series of integers to specify read length. Reads other than the length specified

will be excluded during alignment.

overwrite Overwrite the existing files if TRUE.

## **Details**

Studies on small RNA-seq data from viroid-infected plants have mostly focused on reads with lengths ranging from 21 nt to 24 nt. This function is intended to be used to remove sequence reads with lengths outside the specified range. The default range is 21-24 nt, which can be changed through the read\_lengths argument.

Note that, if filtering by read length has already been performed during the quality control process, there is no need to use this function.

# Value

A path to the filtered FASTQ file.

```
output_dpath <- tempdir()

fq <- system.file(package="CircSeqAlignTk", "extdata", "srna.fq.gz")
output_fq <- file.path(output_dpath, "sran.filtered.fq.gz")
filter_reads(fq, output_fq, seq(21, 24))</pre>
```

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generate\_reads

Generate synthetic sequence reads

#### Description

This function generates synthetic sequence reads to mimic RNA-seq reads sequenced from organelles or organisms with circular genome sequences in FASTQ format file.

# Usage

```
generate_reads(
  n = 10000,
  seq = NULL,
  output = NULL,
  adapter = NULL,
  srna_length = NULL,
  read_length = 150,
  mismatch_prob = 0,
  peaks = NULL,
  read_name_prefix = NULL)
```

#### **Arguments**

n	Number of reads	should be generated.
[1]	Number of reads	should be generated.

seq A file path to a genome sequence in FASTA format file or a string of genome

sequence.

output A file path to store the synthetic reads in FASTQ format file. The extension

should be one of .fq, .fastq. Note that to compress the FASTQ format file,

add .gz or .gzip to the extension (e.g., .fq.gz, .fq.gzip).

adapter A path to a FASTA format file containing a string of adapter sequence. If NULL

is specified, the sequence "AGATCGGAAGAGCACACGTCTGAACTCCAGT-CAC" is used as the adapter sequence. If NA is specified, the adapter sequence is

not included in the synthetic reads.

srna\_length A data frame to specify the lengths of sequence reads sampled from the genome

sequence. The data frame should contain two columns named as length and prob. The values in the length column is used to specify the lengths of sequence reads; the values in the prob column is used to specify the probability that reads with specified length among all reads. If the argument is not given (i.e., srna\_length = NULL), a data frame is randomly generated before sampling

the reads.

read\_length The length of synthetic reads. If adapter is specified, the reads are generated by

concatenating sequence reads and adapter sequences until the specified length.

If adapter = None, ignore this argument.

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mismatch\_prob

A vector to specify probabilities of mismatches occurring in the reads. In order not to allow any mismatches in the reads, set the argument to  $\emptyset$ . To allow multiple mismatches in the reads, set multiple probabilities (e.g.,  $c(\emptyset.05, 0.01)$ ).

peaks

A data frame to specify the peaks of the alignment coverage. The data frame should contain four columns named as mean, std, strand, and prob. The values in the mean and std columns are used to sample the start position of sequence reads from the genome sequence given by seq. The values in the strand column should be + or - to specify which read strand generates the peak. The values in the prob column should be probabilities to use the mean, std, and strand of the same row for read generation. If the argument is not given (i.e., peaks = NULL), a data frame is randomly generated before sampling the reads.

read\_name\_prefix

The prefix of read name in FASTQ format file. If NULL, generate the prefix randomly.

#### Value

A CircSeqAlignTkSim-class object containing parameters for read generation.

#### See Also

CircSeqAlignTkSim-class

```
output_dpath <- tempdir()</pre>
sim <- generate_reads(output = file.path(output_dpath, 'sample1.fq.gz'))</pre>
srna_length <- data.frame(length = c(21, 22, 23, 24),
                           prob = c(0.5, 0.3, 0.1, 0.1)
sim <- generate_reads(output = file.path(output_dpath, 'sample2.fq.gz'),</pre>
                      srna_length = srna_length)
sim <- generate_reads(output = file.path(output_dpath, 'sample3.fq.gz'),</pre>
                      mismatch\_prob = c(0.1, 0.1)
peaks <- data.frame(mean =</pre>
                              c(50, 100, 150),
                              c(3, 5, 5),
                     std =
                     strand = c('+', '-', '+'),
                     prob = c(0.4, 0.4, 0.2))
sim <- generate_reads(output = file.path(output_dpath, 'sample4.fq.gz'),</pre>
                      peaks = peaks)
```

get\_slot\_contents

Get the slot contents from a formal class

# **Description**

This function returns the slot contents from a formal class. It is convenient to use @ when accessing the contents of a slot, however, using @ will generate warnings during the unit tests under software development. This function was created to avoid that warning. Users do not have to use this function.

#### Usage

```
get_slot_contents(object, name)
```

# **Arguments**

object An object from a formally defined class.

name The name of the slot.

#### Value

The contents of the specified slot from the given object.

# Examples

```
output_dpath <- tempdir()
sim <- generate_reads(output = file.path(output_dpath, 'sample1.fq.gz'))
head(get_slot_contents(sim, 'peak'))</pre>
```

```
merge.CircSeqAlignTkSim
```

Merge multiple synthetic datasets

# **Description**

Merge multiple synthetic datasets generated by generate\_reads.

# Usage

```
## S3 method for class 'CircSeqAlignTkSim'
merge(..., output = NULL, overwrite = TRUE)
```

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#### **Arguments**

... CircSeqAlignTkSim class objects.

output A file path to store the synthetic reads in FASTQ format file. The extension

should be one of .fq, .fastq. Note that to compress the FASTQ format file,

add .gz or .gzip to the extension (e.g., .fq.gz, .fq.gzip).

overwrite Overwrite the existing files if TRUE.

#### **Details**

Merge multiple synthetic datasets generated by generate\_reads into one dataset.

#### Value

A CircSeqAlignTkSim-class object.

#### See Also

```
CircSeqAlignTkSim-class, generate_reads
```

### **Examples**

plot\_coverage

Visualize alignment coverage

# Description

This function visualizes the alignment coverage using an area chart. By default, the upper and lower directions of the y-axis represent the alignment coverage of the reads aligned in the forward and reversed strands, respectively.

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#### Usage

```
plot_coverage(x, read_lengths = NULL, fill = "read_length", scale_fun = NULL)
## S3 method for class 'CircSeqAlignTkCoverage'
plot(x, ...)
```

# **Arguments**

A CircSeqAlignTkCoverage-class object generated by the calc\_coverage function.

read\_lengths

Numeric numbers to specify the lengths of reads targeted for visualization. If NULL (default), plot the alignment coverage of reads with all lengths.

fill

Specify NULL or read\_length. If read\_length is specified, then color the area chart according to the read length.

scale\_fun

Set log10 or log to plot the alignment coverage in logarithmic scale.

Other graphical parameters.

#### Value

An object of ggplot2.

#### See Also

CircSeqAlignTkCoverage-class, calc\_coverage

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