# Package: cfTools (via r-universe)

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Type Package

Title Informatics Tools for Cell-Free DNA Study

Version 1.5.0

Description The cfTools R package provides methods for cell-free DNA (cfDNA) methylation data analysis to facilitate cfDNA-based studies. Given the methylation sequencing data of a cfDNA sample, for each cancer marker or tissue marker, we deconvolve the tumor-derived or tissue-specific reads from all reads falling in the marker region. Our read-based deconvolution algorithm exploits the pervasiveness of DNA methylation for signal enhancement, therefore can sensitively identify a trace amount of tumor-specific or tissue-specific cfDNA in plasma. cfTools provides functions for (1) cancer detection: sensitively detect tumor-derived cfDNA and estimate the tumor-derived cfDNA fraction (tumor burden); (2) tissue deconvolution: infer the tissue type composition and the cfDNA fraction of multiple tissue types for a plasma cfDNA sample. These functions can serve as foundations for more advanced cfDNA-based studies, including cancer diagnosis and disease monitoring.

License file LICENSE

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StagedInstall no

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# ${\bf URL} \ {\tt https://github.com/jasminezhoulab/cfTools}$

BugReports https://github.com/jasminezhoulab/cfTools/issues

**Repository** https://bioc.r-universe.dev **RemoteUrl** https://github.com/bioc/cfTools

RemoteRef HEAD

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beta\_matrix

Beta value matrix

# Description

**Index** 

A list of methylation levels (e.g., beta values), where each row is a sample and each column is a marker

```
data("beta_matrix")
```

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## **Format**

A tibble with 20 rows and 3 variables

marker1 Beta values of marker1 for all samplesmarker2 Beta values of marker2 for all samplesmarker3 Beta values of marker3 for all samples

#### Value

A tibble with 20 rows and 3 variables

#### Author(s)

Ran Hu <huran@ucla.edu>

CancerDetector

Cancer Detector

#### **Description**

Detect tumor-derived cfDNA and estimate the tumor burden.

## Usage

```
CancerDetector(
  readsBinningFile,
  tissueMarkersFile,
  lambda = 0.5,
  id = "sample"
)
```

## **Arguments**

readsBinningFile

a file of the fragment-level methylation states of reads that mapped to the markers.

tissueMarkersFile

a file of paired shape parameters of beta distributions for markers.

lambda a number controlling "confounding" markers' distance from average markers.

id the sample ID.

#### Value

a list containing the cfDNA tumor burden and the normal cfDNA fraction.

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## **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
readsBinningFile <- file.path(demo.dir, "CancerDetector.reads.txt.gz")
tissueMarkersFile <- file.path(demo.dir, "CancerDetector.markers.txt.gz")
lambda <- 0.5
id <- "test"

CancerDetector(readsBinningFile, tissueMarkersFile, lambda, id)</pre>
```

CancerDetector.markers

Cancer-specific marker parameter

# Description

The paired shape parameters of beta distributions for cancer-specific markers

## Usage

```
data("CancerDetector.markers")
```

## **Format**

A tibble with 1266 rows and 3 variables

markerName Name of the marker

tumor Paired beta distribution shape parameters for tumor samples

normalPlasma Paired beta distribution shape parameters for normal plasma samples

#### Value

A tibble with 1266 rows and 3 variables

#### Author(s)

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CancerDetector.reads Fragment-level methylation state for cancer detection

## **Description**

The fragment-level methylation states of reads that mapped to the cancer-specific markers

## Usage

```
data("CancerDetector.reads")
```

#### **Format**

A tibble with 9991 rows and 2 variables

markerName Name of the marker

**methState** Fragment-level methylation states, which are represented by a sequence of binary values (0 represents unmethylated CpG and 1 represents methylated CpG on the same fragment)

#### Value

A tibble with 9991 rows and 2 variables

#### Author(s)

Ran Hu <huran@ucla.edu>

cfDeconvolve

cfDNA methylation read deconvolution

## **Description**

Infer the tissue-type composition of plasma cfDNA.

```
cfDeconvolve(
  readsBinningFile,
  tissueMarkersFile,
  numTissues,
  emAlgorithmType = "em.global.unknown",
  likelihoodRatioThreshold = 2,
  emMaxIterations = 100,
  id = "sample"
)
```

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#### **Arguments**

```
readsBinningFile
                  a file of the fragment-level methylation states of reads that mapped to the mark-
                  ers. Either in plain text or compressed form.
tissueMarkersFile
                  a file of paired shape parameters of beta distributions for markers.
                  a number of tissue types.
numTissues
emAlgorithmType
                  a read-based tissue deconvolution EM algorithm type: em.global.unknown (de-
                  fault), em.global.known, em.local.unknown, em.local.known.
likelihoodRatioThreshold
                  a positive float number. Default is 2.
emMaxIterations
                  a number of EM algorithm maximum iteration. Default is 100.
id
                  the sample ID.
```

#### Value

a list containing the cfDNA fractions of different tissue types and an unknown class.

#### **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
readsBinningFile <- file.path(demo.dir, "cfDeconvolve.reads.txt.gz")
tissueMarkersFile <- file.path(demo.dir, "cfDeconvolve.markers.txt.gz")
numTissues <- 7
emAlgorithmType <- "em.global.unknown"
likelihoodRatioThreshold <- 2
emMaxIterations <- 100
id <- "test"

cfDeconvolve(readsBinningFile, tissueMarkersFile, numTissues, emAlgorithmType, likelihoodRatioThreshold, emMaxIterations, id)</pre>
```

cfDeconvolve.markers Tissue-specific marker parameter

## **Description**

The paired shape parameters of beta distributions for tissue-specific markers

```
data("cfDeconvolve.markers")
```

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#### **Format**

A tibble with 10 rows and 8 variables

markerName Name of the marker

tissue1 Paired beta distribution shape parameters for tissue1 samples

tissue2 Paired beta distribution shape parameters for tissue2 samples

tissue3 Paired beta distribution shape parameters for tissue3 samples

tissue4 Paired beta distribution shape parameters for tissue4 samples

tissue5 Paired beta distribution shape parameters for tissue5 samples

tissue6 Paired beta distribution shape parameters for tissue6 samples

tissue7 Paired beta distribution shape parameters for tissue7 samples

#### Value

A tibble with 10 rows and 8 variables

#### Author(s)

Ran Hu <huran@ucla.edu>

cfDeconvolve.reads

Fragment-level methylation state for tissue deconvolution

# Description

The fragment-level methylation states of reads that mapped to the tissue-specific markers

#### Usage

```
data("cfDeconvolve.reads")
```

## **Format**

A tibble with 942 rows and 2 variables

markerName Name of the marker

**methState** Fragment-level methylation states, which are represented by a sequence of binary values (0 represents unmethylated CpG and 1 represents methylated CpG on the same fragment)

#### Value

A tibble with 942 rows and 2 variables

#### Author(s)

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cfSort

cfSort: tissue deconvolution

# **Description**

Tissue deconvolution in cfDNA using DNN models.

## Usage

```
cfSort(readsBinningFile, id = "sample")
```

## **Arguments**

readsBinningFile

a file of the fragment-level methylation states of reads that mapped to the cfSort markers. In compressed form.

id

the sample ID.

## Value

the tissue composition of the cfDNA sample.

# **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
readsBinningFile <- file.path(demo.dir, "cfSort.reads.txt.gz")
id <- "test"

cfSort(readsBinningFile, id)</pre>
```

cfSort.reads

Fragment-level methylation state for cfSort tissue deconvolution

## **Description**

The fragment-level methylation states of reads that mapped to the cfSort markers

```
data("cfSort.reads")
```

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#### **Format**

A tibble with 99999 rows and 6 variables

markerName Name of the cfSort marker

cpgPosition Postions of CpG sites on the fragment

**methState** Fragment-level methylation states, which are represented by a sequence of binary values (0 represents unmethylated CpG and 1 represents methylated CpG on the same fragment)

methCount Number of methylated CpG sites on the fragment

unmethCount Number of unmethylated CpG sites on the fragment

strand Strand

## Value

A tibble with 99999 rows and 6 variables

## Author(s)

Ran Hu <huran@ucla.edu>

cfsort\_markers

cfSort markers

#### **Description**

Marker information for the cfSort function, where each row is the information about a marker

## Usage

```
data("cfsort_markers")
```

## **Format**

A tibble with 51035 rows and 4 variables

marker\_index The marker index used in cfSort methodalpha\_threshold The alpha threshold for each markerpair The pair of tissues used for identifying the markergroup The group number for each marker

#### Value

A tibble with 51035 rows and 4 variables

#### Author(s)

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CpG\_OB\_demo

Methylation information for CpG on the original bottom strand (OB)

# Description

Methylation information for CpG on the original bottom strand (OB), which is one of the outputs from 'bismark methylation extractor'

## Usage

```
data("CpG_OB_demo")
```

#### **Format**

A tibble with 2224 rows and 5 variables

sequence ID ID of the sequence

methylation state Methylated or unmethylated CpG site

chromosome name Chromosome name

chromosome start Chromosome start position

methylation call Methylation call

## Value

A tibble with 2224 rows and 5 variables

## Author(s)

Ran Hu <huran@ucla.edu>

CpG\_OT\_demo

Methylation information for CpG on the original top strand (OT)

## **Description**

Methylation information for CpG on the original top strand (OT), which is one of the outputs from 'bismark methylation extractor'

```
data("CpG_OT_demo")
```

#### **Format**

A tibble with 2556 rows and 5 variables

sequence ID ID of the sequence

methylation state Methylated or unmethylated CpG site

chromosome name Chromosome name

chromosome start Chromosome start position

methylation call Methylation call

#### Value

A tibble with 2556 rows and 5 variables

## Author(s)

Ran Hu <huran@ucla.edu>

demo.fragment\_level.meth.bed

Fragment-level methylation information

# Description

A BED file of fragment-level methylation information

#### Usage

```
data("demo.fragment_level.meth.bed")
```

## **Format**

A tibble with 552 rows and 9 variables

chr Chromosome

**start** Chromosome start

end Chromosome end

name ID of the sequence

fragmentLength Fragment length

strand Strand

cpgNumber Number of CpG sites on the fragment

cpgPosition Postions of CpG sites on the fragment

methState A string of methylation states of CpG sites on the fragment

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# Value

A tibble with 552 rows and 9 variables

# Author(s)

Ran Hu <huran@ucla.edu>

demo.refo\_frag.bed

Fragment-level information

# Description

A BED file of fragment-level information

## Usage

```
data("demo.refo_frag.bed")
```

## **Format**

A tibble with 559 rows and 6 variables

chr Chromosome

start Chromosome start

end Chromosome end

fragmentLength Fragment length

strand Strand

name ID of the sequence

#### Value

A tibble with 559 rows and 6 variables

# Author(s)

demo.refo\_meth.bed

demo.refo\_meth.bed

Methylation information on fragments

# Description

A BED file of methylation information on fragments

#### Usage

```
data("demo.refo_meth.bed")
```

#### **Format**

A tibble with 552 rows and 8 variables

chr Chromosome

cpgStart Start postion of first CpG on the fragment

cpgEnd End postion of first CpG on the fragment

strand Strand

cpgNumber Number of CpG sites on the fragment

cpgPosition Postions of CpG sites on the fragment

methState A string of methylation states of CpG sites on the fragment

name ID of the sequence

## Value

A tibble with 552 rows and 8 variables

# Author(s)

Ran Hu <huran@ucla.edu>

demo.sorted.bed

Paired-end sequencing reads

# Description

Paired-end sequencing reads information

```
data("demo.sorted.bed")
```

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#### **Format**

A tibble with 1117 rows and 6 variables

chr Chromosome namestart Chromosome startend Chromosome endname Sequence ID

score Mapping quality score

strand Strand

## Value

A tibble with 1117 rows and 6 variables

## Author(s)

Ran Hu <huran@ucla.edu>

GenerateFragMeth

Generate fragment-level information about methylation states

## **Description**

Join two lists containing the fragment information and the methylation states on each fragment into one list.

## Usage

```
GenerateFragMeth(frag_bed, meth_bed, output.dir = "", id = "")
```

## Arguments

output.dir

frag_bed	a BED file containing information for every fragment, which is the output of
	MergePEReads().
meth bed	a BED file containing methylation states on every fragment, which is the output

a BED file containing methylation states on every fragment, which is the output of MergeCpGs().

a path to the output directory. Default is "", which means the output will not be

written into a file.

an ID name for the input data. Default is "" which means the output will not h

an ID name for the input data. Default is "", which means the output will not be

written into a file.

#### Value

id

a list in BED file format and/or written to an output BED file.

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## **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
frag_bed <- read.delim(file.path(demo.dir, "demo.refo_frag.bed.txt.gz"),
colClasses = "character")
meth_bed <- read.delim(file.path(demo.dir, "demo.refo_meth.bed.txt.gz"),
colClasses = "character")
output <- GenerateFragMeth(frag_bed, meth_bed)</pre>
```

GenerateMarkerParam

Generate the methylation pattern of markers

# Description

Output paired shape parameters of beta distributions for methylation markers.

## Usage

```
GenerateMarkerParam(x, sample.types, marker.names, output.file = "")
```

## **Arguments**

x	a list of methylation levels (e.g., beta values), where each row is a sample and each column is a marker.
sample.types	a vector of sample types (e.g., tumor or normal, tissue types) corresponding to the rows of the list.
marker.names	a vector of marker names corresponding to the columns of the list.
output.file	a character string naming the output file. Default is "", which means the output will not be written into a file.

#### Value

a list containing the paired shape parameters of beta distributions for markers and/or written to an output file.

# **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
methLevel <- read.table(file.path(demo.dir, "beta_matrix.txt.gz"),
row.names=1, header = TRUE)
sampleTypes <- read.table(file.path(demo.dir, "sample_type.txt.gz"),
row.names=1, header = TRUE)$sampleType
markerNames <- read.table(file.path(demo.dir, "marker_index.txt.gz"),
row.names=1, header = TRUE)$markerIndex</pre>
```

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```
output <- GenerateMarkerParam(methLevel, sampleTypes, markerNames)</pre>
```

markers.bed

Genomic postions of markers

# Description

A BED file of genomic regions of markers

#### Usage

```
data("markers.bed")
```

#### **Format**

A tibble with 3 rows and 4 variables

chr Chromosome

start Chromosome start

end Chromosome end

markerName Marker name

## Value

A tibble with 3 rows and 4 variables

## Author(s)

Ran Hu <huran@ucla.edu>

marker\_index

Marker name

## Description

A vector of marker names corresponding to the columns of the list of methylation levels.

# Usage

```
data("marker_index")
```

#### **Format**

A tibble with 3 rows and 1 variables

markerIndex Marker name

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## Value

A tibble with 3 rows and 1 variables

## Author(s)

Ran Hu <huran@ucla.edu>

Merge(	CpGs
--------	------

Generate fragment-level methylation states of CpGs

## **Description**

Merge the methylation states of all CpGs corresponding to the same fragment onto one line in output.

## Usage

```
MergeCpGs(CpG_OT, CpG_OB, output.dir = "", id = "")
```

# Arguments

CpG_OT	a file of methylation information for CpG on the original top strand (OT), which is one of the outputs from 'bismark methylation extractor'.
CpG_OB	a file of methylation information for CpG on the original bottom strand (OB), which is one of the outputs from 'bismark methylation extractor'.
output.dir	a path to the output directory. Default is "", which means the output will not be written into a file.
id	an ID name for the input data. Default is "", which means the output will not be written into a file.

## Value

a list in BED file format and/or written to an output BED file.

# **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
CpG_OT <- file.path(demo.dir, "CpG_OT_demo.txt.gz")
CpG_OB <- file.path(demo.dir, "CpG_OB_demo.txt.gz")
output <- MergeCpGs(CpG_OT, CpG_OB)</pre>
```

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MergePEReads

Generate fragment-level information for paired-end sequencing reads

## **Description**

Merge BED file (the output of 'bedtools bamtobed') to fragment-level for paired-end sequencing reads.

## Usage

```
MergePEReads(bed_file, output.dir = "", id = "")
```

# Arguments

bed\_file a (sorted) BED file of paired-end reads.

output.dir a path to the output directory. Default is "", which means the output will not be

written into a file.

id an ID name for the input data. Default is "", which means the output will not be

written into a file.

#### Value

a list in BED file format and/or written to an output BED file.

## **Examples**

```
## input files
demo.dir <- system.file("data", package="cfTools")
PEReads <- file.path(demo.dir, "demo.sorted.bed.txt.gz")
output <- MergePEReads(PEReads)</pre>
```

sample\_type

Sample type

# **Description**

A vector of sample types (e.g., tumor or normal, tissue types) corresponding to the rows of the list of methylation levels.

```
data("sample_type")
```

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# Format

A tibble with 20 rows and 1 variables sampleType Sample type

# Value

A tibble with 20 rows and 1 variables

# Author(s)

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