

# Package ‘cfdnakit’

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**Title** Fragmen-length analysis package from high-throughput sequencing of cell-free DNA (cfDNA)

**Version** 1.10.0

**Description** This package provides basic functions for analyzing shallow whole-genome sequencing (~0.3X or more) of cell-free DNA (cfDNA). The package basically extracts the length of cfDNA fragments and aids the visualization of fragment-length information. The package also extract fragment-length information per non-overlapping fixed-sized bins and used it for calculating ctDNA estimation score (CES).

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|                  |                                                                                                 |
|------------------|-------------------------------------------------------------------------------------------------|
| cfdnakit-package | <i>Fragmen-length analysis package from high-throughput sequencing of cell-free DNA (cfDNA)</i> |
|------------------|-------------------------------------------------------------------------------------------------|

---

## Description

This package provides basic functions for analyzing shallow whole-genome sequencing (~0.3X or more) of cell-free DNA (cfDNA). The package basically extracts the length of cfDNA fragments and aids the visualization of fragment-length information. The package also extract fragment-length information per non-overlapping fixed-sized bins and used it for calculating ctDNA estimation score (CES).

## Details

This package provides functions for analyzing using shallow whole-genome sequencing data (~0.3X or more) of circulating cell-free DNA (cfDNA). The aims is to estimate circulating tumor DNA using its characteristic short-fragmented cfDNA. The package extracts length of each cfDNA and assist the visualization of fragment-length distribution. A short-fragment ratio is calculated per non-overlapping fixed-sized bins. Genome-wide copy-number alteration estimated by the short-fragmented cfDNA. The ctDNA estimation score (CES) comprehensively estimate the circulating tumor DNA based on the short-fragment analysis.

## Author(s)

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## Examples

```
library(cfdnakit)
## Reading in a bamfile
sample_bamfile = system.file("extdata",
                             "ex.plasma.bam",
                             package = "cfdnakit")
plasma_SampleBam = read_bamfile(sample_bamfile,
                                apply_blacklist = FALSE)

## Plot a fragment-length distribution of a sample
plot_fragment_dist(list("Plasma.Sample"=plasma_SampleBam))

## Plot a fragment-length distribution of two samples
control_RDS_file =
  system.file("extdata", "BH01_CHR15.SampleBam.rds",
             package = "cfdnakit")
### Load example SampleBam of Healthy cfDNA
control_bins =
  readRDS(control_RDS_file)

comparing_list = list("Healthy.cfDNA"=control_bins,
                      "Patient.1"=plasma_SampleBam)
plot_fragment_dist(comparing_list)

## Derived and plot genome-wide short-fragment cfDNA
patient.SampleFragment =
  get_fragment_profile(plasma_SampleBam,
                      sample_id = "Patient.1")
plot_sl_ratio(patient.SampleFragment)

## Derived and plot normalized short-fragment cfDNA
PoN_rdsfile = system.file(
  "extdata",
  "ex.PoN.rds",
  package = "cfdnakit")
```

```

## Loading example PoN data
PoN.profiles = readRDS(PoN_rdsfile)

sample_zscore =
  get_zscore_profile(patient.SampleFragment,
                    PoN.profiles)
sample_zscore_segment = segmentByPSCB(sample_zscore)
plot_transformed_sl(sample_zscore, sample_zscore_segment)

## Estimate circulating tumor DNA
calculate_CES_score(sample_zscore_segment)

```

---

|                     |                                              |
|---------------------|----------------------------------------------|
| calculate_CES_score | <i>Calculate CES Score from Segmentation</i> |
|---------------------|----------------------------------------------|

---

## Description

Calculate CES Score from Segmentation

## Usage

```
calculate_CES_score(sample_segmentation)
```

## Arguments

```
sample_segmentation
  Segmentation Dataframe
```

## Value

Numeric; CES score

## Examples

```

### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)

sample_zscore_segment <- segmentByPSCB(sample_zscore)

calculate_CES_score(sample_zscore_segment)

```

call\_cnv

*Call Copy-number Variation from SLRatio and segmentation***Description**

Call Copy-number Variation from SLRatio and segmentation

**Usage**

```
call_cnv(
  sample_segmentation,
  sample_zscore,
  callChr = seq_len(22),
  tfs = c(0, 0.7),
  ploidies = c(1.5, 3),
  MaxCN = 4
)
```

**Arguments**

```
sample_segmentation      segmentation dataframe from segmentByPSCBS
sample_zscore            zscore dataframe
callChr                  chromosome to analysis : Default c(1:22)
tfs                      range of fitting tumor fraction : Default c(0,0.8)
ploidies                 range of fitting chromosomal ploidy : Default c(1.5,4)
MaxCN                   maximum copy-number : Default 4
```

**Value**

List of cnvcalling solutions

**Examples**

```
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)

sample_zscore_segment <- segmentByPSCB(sample_zscore)

sample_cnv <- call_cnv(sample_zscore_segment, sample_zscore, tfs=c(0.1,0.3), ploidies=c(1.5,2), MaxCN=3)
```

```
plot_cnv_solution(sample_cnv,selected_solution = 1)
```

---

|                     |                                         |
|---------------------|-----------------------------------------|
| create_blacklist_gr | Create Blacklist regions GRanges object |
|---------------------|-----------------------------------------|

---

**Description**

Create Blacklist regions GRanges object

**Usage**

```
create_blacklist_gr(blacklist_files)
```

**Arguments**

|                 |                                                          |
|-----------------|----------------------------------------------------------|
| blacklist_files | Character; Filepath to file containing blacklist regions |
|-----------------|----------------------------------------------------------|

**Value**

GRanges object of blacklist regions

---

|            |                                     |
|------------|-------------------------------------|
| create_PoN | Create Panel-of-Normal (PoN) object |
|------------|-------------------------------------|

---

**Description**

Create Panel-of-Normal (PoN) object

**Usage**

```
create_PoN(list_rdsfiles)
```

**Arguments**

|               |                                                            |
|---------------|------------------------------------------------------------|
| list_rdsfiles | Character; a file contains paths to Profile.Rdata per line |
|---------------|------------------------------------------------------------|

**Value**

Null

**Examples**

```

healthy.1 <- system.file("extdata","ex.healthy1.rds",package = "cfdnakit")
healthy.2 <- system.file("extdata","ex.healthy2.rds",package = "cfdnakit")

path_to_PoN_txt <- paste0(system.file("extdata",package = "cfdnakit"),"/temp.reference_healthy.listfile")
fileConn<-file(path_to_PoN_txt)
writeLines(c(healthy.1,healthy.2), fileConn)
close(fileConn)

PoN.profiles <- create_PoN(path_to_PoN_txt)
file.remove(path_to_PoN_txt)

```

---

|                     |                                           |
|---------------------|-------------------------------------------|
| extract_insert_size | <i>Extract Insert size from SampleBam</i> |
|---------------------|-------------------------------------------|

---

**Description**

Extract Insert size from SampleBam

**Usage**

```
extract_insert_size(readbam_bin, maximum_length = 600, minimum_length = 20)
```

**Arguments**

|                |                                                                                                            |
|----------------|------------------------------------------------------------------------------------------------------------|
| readbam_bin    | SampleBam Object                                                                                           |
| maximum_length | Int; Maximum length of fragment. cfDNA fragment longer than this value will not be considered; Default 600 |
| minimum_length | Int; Minimum length of fragment. cfDNA fragment shorter than this value will not be considered; Default 20 |

**Value**

Numeric Vector; Insert size of given sample

**Examples**

```

### Loading example SampleBam file
example_file <- system.file("extdata","example_patientcfDNA_SampleBam.RDS",package = "cfdnakit")
sample_bambin <- readRDS(example_file)
extract_insert_size(sample_bambin)
### Extract only insert size of fragment having specific size
extract_insert_size(sample_bambin,maximum_length=500, minimum_length = 50)

```

---

|                          |                                              |
|--------------------------|----------------------------------------------|
| filter_read_on_blacklist | <i>Filter out reads on blacklist regions</i> |
|--------------------------|----------------------------------------------|

---

**Description**

Filter out reads on blacklist regions

**Usage**

```
filter_read_on_blacklist(sample_bin, blacklist_files = NULL, genome = "hg19")
```

**Arguments**

- sample\_bin      SampleBam; Object from function read\_bamfile
- blacklist\_files      Character; Filepath to file containing blacklist regions
- genome      Character; Abbreviation of reference genome; Either hg19 or mm10. default:hg19

**Value**

SampleBam after filtering out read on balck list regions

---

|               |                                           |
|---------------|-------------------------------------------|
| fragment_dist | <i>Get insert-size distribution table</i> |
|---------------|-------------------------------------------|

---

**Description**

Get insert-size distribution table

**Usage**

```
fragment_dist(readbam_bin, maximum_length = 600, minimum_length = 20)
```

**Arguments**

- readbam\_bin      SampleBam Object from function read\_bamfile
- maximum\_length      Int; Maximum length of fragment. cfDNA fragment longer than this value will not be considered; Default 600
- minimum\_length      Int; Minimum length of fragment. cfDNA fragment shorter than this value will not be considered; Default 20

**Value**

Distribution table of fragment length

---

get\_fragment\_profile    *Getting fragment-length information*


---

## Description

Getting fragment-length information

## Usage

```
get_fragment_profile(
  readbam_bin,
  sample_id,
  genome = "hg19",
  short_range = c(100, 150),
  long_range = c(151, 250),
  maximum_length = 600,
  minimum_length = 20
)
```

## Arguments

|                |                                                                                                            |
|----------------|------------------------------------------------------------------------------------------------------------|
| readbam_bin    | SampleBam Object                                                                                           |
| sample_id      | Character; Given sample ID                                                                                 |
| genome         | abbreviation of reference genome; namely hg19, mm10. default:hg19                                          |
| short_range    | Vector of 2 Int; Range of fragment length to be defined as short fragment; Default c(100,150)              |
| long_range     | Vector of 2 Int; Range of fragment length to be defined as long fragment; Default c(151,250)               |
| maximum_length | Int; Maximum length of fragment. cfDNA fragment longer than this value will not be considered; Default 600 |
| minimum_length | Int; Minimum length of fragment. cfDNA fragment shorter than this value will not be considered; Default 20 |

## Value

SampleFragment Object; Fragment length information for quality check and downstream analysis per bin and summary of sample

## Examples

```
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bam_bin <- readRDS(example_file)
sample_profile <- get_fragment_profile(sample_bam_bin, sample_id = "Patient1")
```

---

|                        |                                                                    |
|------------------------|--------------------------------------------------------------------|
| get_segment_bysolution | <i>Return CNV segmentation result from given all CNV solutions</i> |
|------------------------|--------------------------------------------------------------------|

---

**Description**

Return CNV segmentation result from given all CNV solutions

**Usage**

get\_segment\_bysolution(solution, sample\_segmentation, SL\_distance\_df)

**Arguments**

- solution            solution dataframe
- sample\_segmentation            Segmentation dataframe
- SL\_distance\_df    Distance matrix

**Value**

list of segmentation per solution

---

|                    |                                              |
|--------------------|----------------------------------------------|
| get_solution_table | <i>Get summarised table of cnv solutions</i> |
|--------------------|----------------------------------------------|

---

**Description**

Get summarised table of cnv solutions

**Usage**

get\_solution\_table(cnv\_solutions)

**Arguments**

- cnv\_solutions    cnvcalling result from function call\_cnv.R

**Value**

Dataframe of solution table

**Examples**

```

#'
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)

sample_zscore_segment <- segmentByPSCB(sample_zscore)

sample_cnv <- call_cnv(sample_zscore_segment, sample_zscore, tfs=c(0.1, 0.3), ploidy=c(1.5, 2), MaxCN=3)
get_solution_table(sample_cnv)

```

---

|                    |                                                    |
|--------------------|----------------------------------------------------|
| get_zscore_profile | <i>Transform SLRatio with PoN Fragment profile</i> |
|--------------------|----------------------------------------------------|

---

**Description**

Transform SLRatio with PoN Fragment profile

**Usage**

```
get_zscore_profile(fragment_profile, pon_profile)
```

**Arguments**

|                  |                |
|------------------|----------------|
| fragment_profile |                |
|                  | Sample Profile |
| pon_profile      | PoN Profiles   |

**Value**

Dataframe of robust transformed SLratio

**Examples**

```

### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)

### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

```

```
sample_zscore <- get_zscore_profile(sample_profile,pon_profiles)

sample_zscore_segment <- segmentByPSCB(sample_zscore)
```

---

|                  |                                                     |
|------------------|-----------------------------------------------------|
| GRCh2UCSCGRanges | <i>Convert GRCh chromosome format to UCSC style</i> |
|------------------|-----------------------------------------------------|

---

**Description**

Convert GRCh chromosome format to UCSC style

**Usage**

```
GRCh2UCSCGRanges(which)
```

**Arguments**

which                    GRanges object;

**Value**

GRanges; GRanges after chromosome format conversion

---

|                         |                                               |
|-------------------------|-----------------------------------------------|
| <i>if_exist_baifile</i> | <i>Check if bai file exist from given bam</i> |
|-------------------------|-----------------------------------------------|

---

**Description**

Check if bai file exist from given bam

**Usage**

```
if_exist_baifile(bamfile)
```

**Arguments**

bamfile                    Character; Path to sample bamfile

**Value**

Boolean if the bai file exist

---

|                   |                                                         |
|-------------------|---------------------------------------------------------|
| if_ucsc_chrformat | <i>Check UCSC chromosomes format for input bam file</i> |
|-------------------|---------------------------------------------------------|

---

**Description**

Check UCSC chromosomes format for input bam file

**Usage**

```
if_ucsc_chrformat(bamfile_path)
```

**Arguments**

|              |                                   |
|--------------|-----------------------------------|
| bamfile_path | Character; Path to sample bamfile |
|--------------|-----------------------------------|

**Value**

Boolean; if the input bam file is UCSC format, chr prefix

---

|                    |                                           |
|--------------------|-------------------------------------------|
| make_density_table | <i>Make Fragment-length density table</i> |
|--------------------|-------------------------------------------|

---

**Description**

Make Fragment-length density table

**Usage**

```
make_density_table(readbam_bin, minimum_length, maximum_length)
```

**Arguments**

|                |                                                                                 |
|----------------|---------------------------------------------------------------------------------|
| readbam_bin    | List; A list containing SampleBam object/objects from the read_bamfile function |
| minimum_length | numeric;                                                                        |
| maximum_length | numeric                                                                         |

**Value**

data.frame

---

```
overlap_bin_with_segment
```

*Overlap and merge bin data frame with segmentation dataframe*

---

### Description

Overlap and merge bin data frame with segmentation dataframe

### Usage

```
overlap_bin_with_segment(per_bin_profile, sample_segmentation)
```

### Arguments

```
per_bin_profile      bin dataframe
sample_segmentation  segmentation dataframe
```

### Value

dataframe of overlapping bin and segmentation

---

```
plot_cnv_solution
```

*Plot Fragment-length profile with CNV calling result*

---

### Description

Plot Fragment-length profile with CNV calling result

### Usage

```
plot_cnv_solution(
  cnvcall,
  selected_solution = 1,
  genome = "hg19",
  ylim = c(-30, 30)
)
```

### Arguments

```
cnvcall      solution results from call_cnv function
selected_solution  solution rank to plot

genome      Character; version of reference genome (default hg19)
ylim        Vector of 2 Int; ylim of plot (default c(-20,20))
```

**Value**

ggplot object plot Genomics CNV profile of selected solution

**Examples**

```
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)
sample_zscore_segment <- segmentByPSCB(sample_zscore)

sample_cnv <- call_cnv(sample_zscore_segment, sample_zscore, tfs=c(0.1, 0.3), ploidy=c(1.5, 2), MaxCN=3)
plot_cnv_solution(sample_cnv, selected_solution = 1)
```

---

plot\_distance\_matrix    *Plot Distance Matrix from CNVCalling*

---

**Description**

Plot Distance Matrix from CNVCalling

**Usage**

```
plot_distance_matrix(cnvcall)
```

**Arguments**

cnvcall                      cnvcalling result from function call\_cnv.R

**Value**

ggplot object ; distance matrix per cnvcalling solution

**Examples**

```
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")
```

```

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)
sample_zscore_segment <- segmentByPSCB(sample_zscore)

sample_cnv <- call_cnv(sample_zscore_segment, sample_zscore, tfs=c(0.1, 0.3), ploidies=c(1.5, 2), MaxCN=3)
plot_distance_matrix(sample_cnv)

```

---

plot\_fragment\_dist      *Plot Fragment-length Distribution*

---

### Description

Plot Fragment-length Distribution

### Usage

```
plot_fragment_dist(readbam_list, maximum_length = 550, minimum_length = 20)
```

### Arguments

|                |                                                                                                            |
|----------------|------------------------------------------------------------------------------------------------------------|
| readbam_list   | List; A list containing SampleBam object/objects from the read_bamfile function                            |
| maximum_length | Int; Maximum length of fragment. cfDNA fragment longer than this value will not be considered; Default 550 |
| minimum_length | Int; Minimum length of fragment. cfDNA fragment shorter than this value will not be considered; Default 20 |

### Value

distribution plot

### Examples

```

example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)

### adding more samples to the plot
example_file2 <- system.file("extdata", "BH01_CHR15.SampleBam.rds", package = "cfdnakit")
control_bambin <- readRDS(example_file2)
readbam_list <- list(plasma1 = sample_bambin, Healthy.blood.plasma=control_bambin)
plot_fragment_dist(readbam_list)

```

---

|               |                                       |
|---------------|---------------------------------------|
| plot_sl_ratio | <i>Plot Short/Long-fragment Ratio</i> |
|---------------|---------------------------------------|

---

**Description**

Plot Short/Long-fragment Ratio

**Usage**

```
plot_sl_ratio(fragment_profile, ylim = c(0, 0.4), genome = "hg19")
```

**Arguments**

|                  |                                                       |
|------------------|-------------------------------------------------------|
| fragment_profile | list                                                  |
| ylim             | plot y-axis limit                                     |
| genome           | Character; version of reference genome (default hg19) |

**Value**

plot

**Examples**

```
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")
plot_sl_ratio(fragment_profile = sample_profile)

### change plot y-axis
plot_sl_ratio(fragment_profile = sample_profile, ylim=c(0.1,0.5))

### change reference genome
plot_sl_ratio(fragment_profile = sample_profile, genome="hg38")
```

---

|                     |                                                     |
|---------------------|-----------------------------------------------------|
| plot_transformed_sl | <i>Plot z-transformed Short/Long-fragment Ratio</i> |
|---------------------|-----------------------------------------------------|

---

**Description**

Plot z-tranformed Short/Long-fragment Ratio

Usage

```
plot_transformed_sl(
  sample_transformed_sl,
  sample_segment_df = NULL,
  ylim = c(-30, 30),
  genome = "hg19"
)
```

Arguments

- sample\_transformed\_sl  
Dataframe z-transformed SLRatio from get\_zscore\_profile
- sample\_segment\_df  
Dataframe segmenation from segmentByPSCB
- ylim  
plot y-axis limit
- genome  
Character; version of reference genome (default hg19)

Value

Genome-wide plot of z-transformed SLRatio

Examples

```
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)
sample_zscore_segment <- segmentByPSCB(sample_zscore)
plot_transformed_sl(sample_zscore, sample_zscore_segment)
## Change reference genome
plot_transformed_sl(sample_zscore, sample_zscore_segment, genome="hg38")
```

---

|              |                                                                                                                                           |
|--------------|-------------------------------------------------------------------------------------------------------------------------------------------|
| read_bamfile | <i>Read a bam file Read a bam file from give path. Alignment and sequencing read information will be binned into non-overlapping size</i> |
|--------------|-------------------------------------------------------------------------------------------------------------------------------------------|

---

Description

Read a bam file Read a bam file from give path. Alignment and sequencing read information will be binned into non-overlapping size

**Usage**

```
read_bamfile(
  bamfile_path,
  binsize = 1000,
  blacklist_files = NULL,
  genome = "hg19",
  target_bedfile = NULL,
  min_mapq = 20,
  apply_blacklist = TRUE
)
```

**Arguments**

|                 |                                                                                              |
|-----------------|----------------------------------------------------------------------------------------------|
| bamfile_path    | Character; Path to sample bamfile                                                            |
| binsize         | Int; Size of non-overlapping windows in KB. Only 100,500 and 1000 is available; Default 1000 |
| blacklist_files | Character; Filepath to file containing blacklist regions                                     |
| genome          | Character; abbreviation of reference genome; available genome: hg19,hg38, mm10. default:hg19 |
| target_bedfile  | Character; Path to exon/target bedfile; Default NULL                                         |
| min_mapq        | Int; minimum read mapping quality; Default 20                                                |
| apply_blacklist | Logical; To exclude read on the blacklist regions Default TRUE                               |

**Value**

SampleBam Object; A list object containing read information from the BAM file.

**Examples**

```
f1 <- system.file("extdata","ex.plasma.bam",package = "cfDNAkit")
### read bam file with default params (hg19, 1000K binsize)
sample.bam <- read_bamfile(f1, apply_blacklist=FALSE)
```

---

|                |                                                      |
|----------------|------------------------------------------------------|
| read_PoN_files | <i>Read Fragment Profile from a list of rds file</i> |
|----------------|------------------------------------------------------|

---

**Description**

Read Fragment Profile from a list of rds file

**Usage**

```
read_PoN_files(list_rdsfiles)
```

**Arguments**

list\_rdsfiles    path to file containing list of rds file

**Value**

list containing content of rds file

---

|               |                                     |
|---------------|-------------------------------------|
| segmentByPSCB | <i>Segmentation data with PSCBS</i> |
|---------------|-------------------------------------|

---

**Description**

Segmentation data with PSCBS

**Usage**

segmentByPSCB(sample\_transformed\_sl)

**Arguments**

sample\_transformed\_sl  
dataframe of z-transformed SLRatio

**Value**

Dataframe of segmentation result

**Examples**

```
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
### Example PoN
PoN_rdsfile <- system.file("extdata", "ex.PoN.rds", package = "cfdnakit")
pon_profiles <- readRDS(PoN_rdsfile)
sample_profile <- get_fragment_profile(sample_bambin, sample_id = "Patient1")

sample_zscore <- get_zscore_profile(sample_profile, pon_profiles)
sample_zscore_segment <- segmentByPSCB(sample_zscore)
```

---

```
test_isize_KolmogorovSmirnov
```

*KolmogorovSmirnov test for insert size*

---

**Description**

KolmogorovSmirnov test for insert size

**Usage**

```
test_isize_KolmogorovSmirnov(control_insert_size, sample_insert_size)
```

**Arguments**

```
control_insert_size
```

Vector of insert size of a control sample

```
sample_insert_size
```

Vector of insert size of a testing sample

**Value**

KS.Test result

**Examples**

```
### Loading example SampleBam file
example_file <- system.file("extdata", "example_patientcfDNA_SampleBam.RDS", package = "cfdnakit")
sample_bambin <- readRDS(example_file)
control_rds <- "BH01_CHR15.SampleBam.rds"
control_RDS_file <- system.file("extdata", control_rds, package = "cfdnakit")
control_fragment_profile <- readRDS(control_RDS_file)
sample.isize <- extract_insert_size(sample_bambin)
healthy.isize <- extract_insert_size(control_fragment_profile)
test_isize_KolmogorovSmirnov(sample.isize, healthy.isize)
```

---

```
UCSC2GRChSampleBam
```

*Convert UCSC chromosome format to GRCh style from a list of alignment information*

---

**Description**

Convert UCSC chromosome format to GRCh style from a list of alignment information

**Usage**

```
UCSC2GRChSampleBam(sample.bam)
```

Arguments

sample.bam      list of alignment information from function read\_bamfile

Value

List; list of alignment information after conversion

---

|                   |                                  |
|-------------------|----------------------------------|
| util.bias_correct | <i>Correct GC Bias readcount</i> |
|-------------------|----------------------------------|

---

Description

Correct GC Bias readcount

Usage

util.bias\_correct(readcount, bias)

Arguments

readcount      numeric  
bias            numeric

Value

numeric

---

|                  |                                                                 |
|------------------|-----------------------------------------------------------------|
| zscore_transform | <i>zscore_transform transforms SLRatio profile into z-score</i> |
|------------------|-----------------------------------------------------------------|

---

Description

zscore\_transform transforms SLRatio profile into z-score

Usage

zscore\_transform(per\_bin\_profile)

Arguments

per\_bin\_profile  
SampleFragment from function get\_fragment\_profile

Value

dataframe of z-score per bin

---

`%>%`*Pipe operator*

---

**Description**

See `magrittr::%>%` for details.

**Arguments**

|                  |                                                            |
|------------------|------------------------------------------------------------|
| <code>lhs</code> | A value or the <code>magrittr</code> placeholder.          |
| <code>rhs</code> | A function call using the <code>magrittr</code> semantics. |

**Value**

The result of calling `rhs(lhs)`.

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