

Package ‘methrix’

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Title Fast and efficient summarization of generic bedGraph files from Bisulfite sequencing

Version 1.4.0

Description Bedgraph files generated by Bisulfite pipelines often come in various flavors. Critical downstream step requires summarization of these files into methylation/coverage matrices. This step of data aggregation is done by Methrix, including many other useful downstream functions.

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LazyData false

Depends R (>= 3.6), data.table (>= 1.12.4), SummarizedExperiment

Imports rtracklayer, DelayedArray, HDF5Array, BSgenome, rjson, DelayedMatrixStats, parallel, methods, ggplot2, matrixStats, graphics, stats, utils, GenomicRanges, IRanges

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combine_methrix	<i>Combine methrix objects</i>
-----------------	--------------------------------

Description

Combine methrix objects

Usage

```
combine_methrix(m1, m2, by = c("row", "col"))
```

Arguments

m1	Frist methrix object
m2	Second methrix object
by	The direction of combine. 'column' (cbind) combines samples with same regions, 'row' combines different regions, e.g. different chromosomes.

Details

Takes a `methrix` object and returns with the same object with delayed array assay slots with HDF5 backend. Might take long time!

Value

An object of class `methrix`, HDF5 format

Examples

```
data(methrix_data)
m2 <- convert_methrix(m=methrix_data)
```

coverage_filter	<i>Filter matrices by coverage</i>
-----------------	------------------------------------

Description

Filter matrices by coverage

Usage

```
coverage_filter(
  m,
  cov_thr = 1,
  min_samples = 1,
  prop_samples = 0,
  group = NULL,
  n_chunks = 1,
  n_cores = 1
)
```

Arguments

<code>m</code>	<code>methrix</code> object
<code>cov_thr</code>	minimum coverage required to call a loci covered
<code>min_samples</code>	Minimum number of samples that should have a loci with coverage \geq <code>cov_thr</code> . If <code>group</code> is given, then this applies per group. Only need one of <code>prop_samples</code> or <code>min_samples</code> .
<code>prop_samples</code>	Minimum proportion of samples that should have a loci with coverage \geq <code>cov_thr</code> . If <code>group</code> is given, then this applies per group. Only need one of <code>prop_samples</code> or <code>min_samples</code> .
<code>group</code>	a column name from sample annotation that defines groups. In this case, the number of <code>min_samples</code> will be tested group-wise.
<code>n_chunks</code>	Number of chunks to split the <code>methrix</code> object in case it is very large. Default = 1.
<code>n_cores</code>	Number of parallel instances. <code>n_cores</code> should be less than or equal to <code>n_chunks</code> . If <code>n_chunks</code> is not specified, then <code>n_chunks</code> is initialized to be equal to <code>n_cores</code> . Default = 1.

Details

Takes `methrix` object and filters CpGs based on coverage statistics

Value

An object of class `methrix`

Examples

```
data('methrix_data')
#keep only CpGs which are covered by at-least 1 read across 3 samples
coverage_filter(m = methrix_data, cov_thr = 1, min_samples = 3)
```

extract_CPGs	<i>Extracts all CpGs from a genome</i>
--------------	--

Description

Extracts all CpGs from a genome

Usage

```
extract_CPGs(ref_genome = NULL)
```

Arguments

`ref_genome` BSgenome object or name of the installed BSgenome package. Example: `BSgenome.Hsapiens.UCSC`

Value

a list of data.table containing number of CpG's and contig lengths

Examples

```
## Not run:
hg19_cpgs = methrix::extract_CPGs(ref_genome = 'BSgenome.Hsapiens.UCSC.hg19')

## End(Not run)
```

get_matrix	<i>Extract methylation or coverage matrices</i>
------------	---

Description

Extract methylation or coverage matrices

Usage

```
get_matrix(m, type = "M", add_loci = FALSE, in_granges = FALSE)
```

Arguments

m	methrix object
type	can be M or C. Default 'M'
add_loci	Default FALSE. If TRUE adds CpG position info to the matrix and returns as a data.table
in_granges	Do you want the outcome in GRanges?

Details

Takes methrix object and returns user specified methylation or coverage matrix

Value

Coverage or Methylation matrix

Examples

```
data('methrix_data')
#Get methylation matrix
get_matrix(m = methrix_data, type = 'M')
#Get methylation matrix along with loci
get_matrix(m = methrix_data, type = 'M', add_loci = TRUE)
#' #Get methylation data as a GRanges object
get_matrix(m = methrix_data, type = 'M', add_loci = TRUE, in_granges=TRUE)
```

get_region_summary	<i>Extract and summarize methylation or coverage info by regions of interest</i>
--------------------	--

Description

Extract and summarize methylation or coverage info by regions of interest

Usage

```

get_region_summary(
  m,
  regions = NULL,
  type = "M",
  how = "mean",
  overlap_type = "within",
  na_rm = TRUE,
  elementMetadata.col = NULL,
  verbose = TRUE,
  n_chunks = 1,
  n_cores = 1
)

```

Arguments

<code>m</code>	<code>methrix</code> object
<code>regions</code>	genomic regions to be summarized. Could be a <code>data.table</code> with 3 columns (chr, start, end) or a <code>GenomicRanges</code> object
<code>type</code>	matrix which needs to be summarized. Could be 'M', 'C'. Default 'M'
<code>how</code>	mathematical function by which regions should be summarized. Can be one of the following: mean, sum, max, min. Default 'mean'
<code>overlap_type</code>	defines the type of the overlap of the CpG sites with the target region. Default value is 'within'. For detailed description, see the <code>findOverlaps</code> function of the IRanges package.
<code>na_rm</code>	Remove NA's? Default TRUE
<code>elementMetadata.col</code>	columns in <code>methrix@elementMetadata</code> which needs to be summarised. Default = NULL.
<code>verbose</code>	Default TRUE
<code>n_chunks</code>	Number of chunks to split the <code>methrix</code> object in case it is very large. Default = 1.
<code>n_cores</code>	Number of parallel instances. <code>n_cores</code> should be less than or equal to <code>n_chunks</code> . If <code>n_chunks</code> is not specified, then <code>n_chunks</code> is initialized to be equal to <code>n_cores</code> . Default = 1.

Details

Takes `methrix` object and summarizes regions

Value

a coverage or methylation matrix

Examples

```

data('methrix_data')
get_region_summary(m = methrix_data,
  regions = data.table(chr = 'chr21', start = 27867971, end = 27868103),
  type = 'M', how = 'mean')

```

get_stats	<i>Estimate descriptive statistics</i>
-----------	--

Description

Estimate descriptive statistics

Usage

```
get_stats(m, per_chr = TRUE)
```

Arguments

m	methrix object
per_chr	Estimate stats per chromosome. Default TRUE

Details

Calculate descriptive statistics

Value

data.table of summary stats

See Also

[plot_stats](#)

Examples

```
data('methrix_data')  
get_stats(methrix_data)
```

load_HDF5_methrix	<i>Loads HDF5 methrix object</i>
-------------------	----------------------------------

Description

Loads HDF5 methrix object

Usage

```
load_HDF5_methrix(dir = NULL, ...)
```

Arguments

dir	The directory to read in from. Default NULL
...	Parameters to pass to loadHDF5SummarizedExperiment

Details

Takes directory with a previously saved HDF5Array format `methrix` object and loads it

Value

An object of class `methrix`

Examples

```
data('methrix_data')
methrix_data_h5 <- convert_methrix(m=methrix_data)
target_dir = paste0(getwd(), '/temp1/')
save_HDF5_methrix(methrix_data_h5, dir = target_dir, replace = TRUE)
load_HDF5_methrix(target_dir)
```

mask_methrix	<i>Masks too high or too low coverage</i>
--------------	---

Description

Masks too high or too low coverage

Usage

```
mask_methrix(m, low_count = NULL, high_quantile = 0.99, n_cores = 1)
```

Arguments

<code>m</code>	<code>methrix</code> object
<code>low_count</code>	The minimal coverage allowed. Everything below, will get masked. Default = NULL, nothing gets masked.
<code>high_quantile</code>	The quantile limit of coverage. Quantiles are calculated for each sample and everything that belongs to a higher quantile than the defined will be masked. Default = 0.99.
<code>n_cores</code>	Number of parallel instances. Can only be used if <code>methrix</code> is in HDF5 format. Default = 1.

Details

Takes `methrix` object and masks sites with too high or too low coverage by putting NA for coverage and beta value. The sites will remain in the object.

Value

An object of class `methrix`

Examples

```
data('methrix_data')
mask_methrix(m = methrix_data, low_count = 5, high_quantile = 0.99 )
```

methrix-class	<i>Class methrix</i>
---------------	----------------------

Description

S4 class Methrix

Slots

assays A list of two matrices containing 'Methylation' and 'Coverage' information

elementMetadata A DataFrame describing rows in corresponding assay matrices.

colData genome: the name of the BSgenome that was used to extract CpGs, isHDF5: is it stored in HDF5 Array format

metadata a list of meta data associated with the assays

NAMES NULL

methrix2bsseq	<i>Convert methrix to bsseq object</i>
---------------	--

Description

Convert [methrix](#) to bsseq object

Usage

```
methrix2bsseq(m)
```

Arguments

m [methrix](#) object

Details

Takes [methrix](#) object and returns a bsseq object

Value

An object of class bsseq

Examples

```
## Not run:
data('methrix_data')
methrix2bsseq(m = methrix_data)

## End(Not run)
```

`methrix_data`*WGBS for colon cancer, chr21 and chr22*

Description

This is a subset of original 'bsseqData' converted to 'methrix' containing Whole-genome bisulfite sequencing data (WGBS) for colon cancer on chromosome 21 and 22.

Usage

```
data('methrix_data')
```

Format

An object of class 'methrix'

References

Hansen, K. D. et al. (2011) Increased methylation variation in epigenetic domains across cancer types. *Nature Genetics* 43, 768-775.

Examples

```
data('methrix_data')
methrix_data
```

`methrix_pca`*Principal Component Analysis*

Description

Principal Component Analysis

Usage

```
methrix_pca(  
  m,  
  var = "top",  
  top_var = 1000,  
  ranges = NULL,  
  pheno = NULL,  
  do_plot = TRUE,  
  n_pc = 2  
)
```

Arguments

m	Input <code>methrix</code> object
var	Choose between random CpG sites ('rand') or most variable CpGs ('top').
top_var	Number of variable CpGs to use. Default 1000 Set it to NULL to use all CpGs (which is not recommended due to memory requirements). This option is mutually exclusive with ranges.
ranges	genomic regions to be summarized. Could be a data.table with 3 columns (chr, start, end) or a GenomicRanges object
pheno	Column name of colData(m). Default NULL. Will be used as a factor to color different groups
do_plot	Should a plot be generated?
n_pc	Default 2.

Value

PCA results

Examples

```
data('methrix_data')
methrix_pca(methrix_data, do_plot = FALSE)
```

methrix_report	<i>Creates a detailed interactive html summary report from Methrix object</i>
----------------	---

Description

Creates a detailed interactive html summary report from Methrix object. If the directory contains required files (from previous run), it directly proceeds to generate html report.

Usage

```
methrix_report(
  meth,
  output_dir = NULL,
  recal_stats = FALSE,
  plot_beta_dist = TRUE,
  beta_nCpG = 10000,
  prefix = NULL,
  n_thr = 4
)
```

Arguments

meth	<code>methrix</code> object
output_dir	Output directory name where the files should be saved. If NULL creates a tempdir
recal_stats	Whether summary statistics should be recalculated? If you are using subsetted methrix object set this to TRUE.

plot_beta_dist	Default TRUE. Can be time consuming.
beta_nCpG	Number of CpGs rto use for estimating beta value distribution. Default 10000
prefix	If provided, the name of the report and the intermediate files will start with the prefix.
n_thr	Default 4. Only used if plot_beta_dist is TRUE

Value

an interactive html report

Examples

```
## Not run:  
data('methrix_data')  
methrix::methrix_report(meth = methrix_data)  
  
## End(Not run)
```

order_by_sd	<i>Order mathrix object by SD</i>
-------------	-----------------------------------

Description

Order mathrix object by SD

Usage

```
order_by_sd(m)
```

Arguments

m [methrix](#) object

Details

Takes [methrix](#) object and reorganizes the data by standard deviation

Value

An object of class [methrix](#)

Examples

```
data('methrix_data')  
order_by_sd(m = methrix_data)
```

plot_coverage	Coverage QC Plots
---------------	-------------------

Description

Coverage QC Plots

Usage

```
plot_coverage(  
  m,  
  type = c("hist", "dens"),  
  pheno = NULL,  
  perGroup = FALSE,  
  lim = 100,  
  size.lim = 1e+06,  
  col_palette = "RdYlGn"  
)
```

Arguments

m	Input <i>methrix</i> object
type	Choose between 'hist' (histogram) or 'dens' (density plot).
pheno	Column name of colData(m). Will be used as a factor to color different groups in the plot.
perGroup	Color the plots in a sample-wise manner?
lim	Maximum coverage value to be plotted.
size.lim	The maximum number of observations (sites*samples) to use. If the dataset is larger than this, random sites will be selected from the genome.
col_palette	Name of the RColorBrewer palette to use for plotting.

Value

ggplot2 object

Examples

```
data('methrix_data')  
plot_coverage(m = methrix_data)
```

plot_density	<i>Density Plot of β-Values</i>
--------------	--

Description

Density Plot of β -Values

Usage

```
plot_density(  
  m,  
  ranges = NULL,  
  n_cpgs = 25000,  
  pheno = NULL,  
  col_palette = "RdYlGn"  
)
```

Arguments

m	Input <code>methrix</code> object
ranges	genomic regions to be summarized. Could be a <code>data.table</code> with 3 columns (chr, start, end) or a <code>GenomicRanges</code> object
n_cpgs	Use these many random CpGs for plotting. Default 25000. Set it to NULL to use all - which can be memory expensive.
pheno	Column name of <code>colData(m)</code> . Will be used as a factor to color different groups in the violin plot.
col_palette	Name of the <code>RColorBrewer</code> palette to use for plotting.

Value

ggplot2 object

Examples

```
data('methrix_data')  
plot_density(m = methrix_data)
```

plot_pca	<i>Plot PCA results</i>
----------	-------------------------

Description

Plot PCA results

Usage

```
plot_pca(
  pca_res,
  m = NULL,
  col_anno = NULL,
  shape_anno = NULL,
  pc_x = "PC1",
  pc_y = "PC2",
  show_labels = FALSE
)
```

Arguments

pca_res	Results from methrix_pca
m	optimal methrix object. Default NULL
col_anno	Column name of colData(m). Default NULL. Will be used as a factor to color different groups. Required methrix object
shape_anno	Column name of colData(m). Default NULL. Will be used as a factor to shape different groups. Required methrix object
pc_x	Default 'PC1'
pc_y	Default 'PC2'
show_labels	Default FLASE

Value

ggplot2 object

Examples

```
data('methrix_data')
mpc = methrix_pca(methrix_data, do_plot = FALSE)
plot_pca(mpc)
```

plot_stats

Plot descriptive statistics

Description

Plot descriptive statistics

Usage

```
plot_stats(
  plot_dat,
  what = "M",
  stat = "mean",
  ignore_chr = NULL,
  samples = NULL,
  n_col = NULL,
  n_row = NULL
)
```


Arguments

plot_dat	results from get_stats
what	Can be M or C. Default M
stat	Can be mean or median. Default mean
ignore_chr	Chromosomes to ignore. Default NULL
samples	Use only these samples. Default NULL
n_col	number of columns. Passed to 'facet_wrap'
n_row	number of rows. Passed to 'facet_wrap'

Details

plot descriptive statistics results from [get_stats](#)

Value

ggplot2 object

See Also

[get_stats](#)

Examples

```
data('methrix_data')
gs = get_stats(methrix_data)
plot_stats(gs)
```

plot_violin

Violin Plot for β -Values

Description

Violin Plot for β -Values

Usage

```
plot_violin(
  m,
  ranges = NULL,
  n_cpgs = 25000,
  pheno = NULL,
  col_palette = "RdYlGn"
)
```

Arguments

m	Input <code>methrix</code> object
ranges	genomic regions to be summarized. Could be a <code>data.table</code> with 3 columns (chr, start, end) or a <code>GenomicRanges</code> object
n_cpgs	Use these many random CpGs for plotting. Default 25000. Set it to NULL to use all - which can be memory expensive.
pheno	Column name of <code>colData(m)</code> . Will be used as a factor to color different groups in the violin plot.
col_palette	Name of the <code>RColorBrewer</code> palette to use for plotting.

Value

ggplot2 object

Examples

```
data('methrix_data')
plot_violin(m = methrix_data)
```

read_bedgraphs	<i>Versatile BedGraph reader.</i>
----------------	-----------------------------------

Description

Versatile BedGraph reader.

Usage

```
read_bedgraphs(
  files = NULL,
  pipeline = NULL,
  zero_based = TRUE,
  stranded = FALSE,
  collapse_strands = FALSE,
  ref_cpgs = NULL,
  ref_build = NULL,
  contigs = NULL,
  vect = FALSE,
  vect_batch_size = NULL,
  coldata = NULL,
  chr_idx = NULL,
  start_idx = NULL,
  end_idx = NULL,
  beta_idx = NULL,
  M_idx = NULL,
  U_idx = NULL,
  strand_idx = NULL,
  cov_idx = NULL,
  synced_coordinates = FALSE,
  n_threads = 1,
```

```

    h5 = FALSE,
    h5_dir = NULL,
    h5temp = NULL,
    verbose = TRUE
)

```

Arguments

files	bedgraph files.
pipeline	Default NULL. Currently supports "Bismark_cov", "MethylDackel", "MethylTools", "BisSNP", "BSseeker2_CGmap" If not known use idx arguments for manual column assignments.
zero_based	Are bedgraph regions zero based ? Default TRUE
stranded	Default FALSE
collapse_strands	If TRUE collapses CpGs on different crick strand into watson. Deafult FALSE
ref_cpgs	BSgenome object, or name of the installed BSgenome package, or an output from extract_CPGs . Example: BSgenome.Hsapiens.UCSC.hg19
ref_build	reference genome for bedgraphs. Default NULL. Only used for additional details. Doesnt affect in any way.
contigs	contigs to restrict genomic CpGs to. Default all autosomes and allosomes - ignoring extra contigs.
vect	To use vectorized code. Default FALSE. Set to TRUE if you don't have large number of BedGraph files.
vect_batch_size	Default NULL. Process samples in batches. Applicable only when vect = TRUE
coldata	An optional DataFrame describing the samples. Row names, if present, become the column names of the matrix. If NULL, then a DataFrame will be created with basename of files used as the row names.
chr_idx	column index for chromosome in bedgraph files
start_idx	column index for start position in bedgraph files
end_idx	column index for end position in bedgraph files
beta_idx	column index for beta values in bedgraph files
M_idx	column index for read counts supporting Methylation in bedgraph files
U_idx	column index for read counts supporting Un-methylation in bedgraph files
strand_idx	column index for strand information in bedgraph files
cov_idx	column index for total-coverage in bedgraph files
synced_coordinates	Are the start and end coordinates of a stranded bedgraph are synchronized between + and - strands? Possible values: FALSE (default), TRUE if the start coordinates are the start coordinates of the C on the plus strand.
n_threads	number of threads to use. Default 1. Be-careful - there is a linear increase in memory usage with number of threads. This option is does not work with Windows OS.
h5	Should the coverage and methylation matrices be stored as 'HDF5Array'
h5_dir	directory to store H5 based object
h5temp	temporary directory to store hdf5
verbose	Be little chatty ? Default TRUE.

Details

Reads BedGraph files and generates methylation and coverage matrices. Optionally arrays can be serialized as on-disk HDFS5 arrays.

Value

An object of class `methrix`

Examples

```
## Not run:
bdg_files = list.files(path = system.file('extdata', package = 'methrix'),
  pattern = '*\\.bedGraph\\.gz$', full.names = TRUE)
hg19_cpgs = methrix::extract_CPGs(ref_genome = 'BSgenome.Hsapiens.UCSC.hg19')
meth = methrix::read_bedgraphs( files = bdg_files, ref_cpgs = hg19_cpgs,
  chr_idx = 1, start_idx = 2, M_idx = 3, U_idx = 4,
  stranded = FALSE, zero_based = FALSE, collapse_strands = FALSE)

## End(Not run)
```

region_filter	<i>Filter matrices by region</i>
---------------	----------------------------------

Description

Filter matrices by region

Usage

```
region_filter(m, regions, type = "within")
```

Arguments

m	<code>methrix</code> object
regions	genomic regions to filter-out. Could be a <code>data.table</code> with 3 columns (chr, start, end) or a <code>GenomicRanges</code> object
type	defines the type of the overlap of the CpG sites with the target regions. Default value is 'within'. For detailed description, see the <code>foverlaps</code> function of the <code>data.table</code> package.

Details

Takes `methrix` object and filters CpGs based on supplied regions in `data.table` or `GRanges` format

Value

An object of class `methrix`

Examples

```
data('methrix_data')
region_filter(m = methrix_data,
  regions = data.table(chr = 'chr21', start = 27867971, end = 27868103))
```

remove_snps	<i>Removes CpG sites from the object if they overlap with common SNPs</i>
-------------	---

Description

Removes CpG sites from the object if they overlap with common SNPs

Usage

```
remove_snps(
  m,
  populations = NULL,
  maf_threshold = 0.01,
  reduce_filtering = FALSE,
  forced = FALSE,
  keep = FALSE,
  n_chunks = 1,
  n_cores = 1
)
```

Arguments

m	<code>methrix</code> object
populations	Populations to use. Default is all.
maf_threshold	The frequency threshold, above which the SNPs will be removed. Default is 0.01
reduce_filtering	If TRUE, the SNPs with a MAF < 0.1 will be evaluated and only the highly variable ones will be removed. Default FALSE.
forced	the <code>reduce_filtering</code> is not recommended with less than 10 samples, but can be forced. Default is FALSE.
keep	Do you want to keep the sites that were filtered out? In this case, the function will return with a list of two <code>methrix</code> objects.
n_chunks	Number of chunks to split the <code>methrix</code> object in case it is very large. Can only be used if input data is in HDF5 format. Default = 1.
n_cores	Number of parallel instances. Can only be used if input data is in HDF5 format. <code>n_cores</code> should be less than or equal to <code>n_chunks</code> . If <code>n_chunks</code> is not specified, then <code>n_chunks</code> is initialized to be equal to <code>n_cores</code> . Default = 1.

Details

Takes `methrix` object and removes common SNPs. SNPs overlapping with a CpG site and have a minor allele frequency (MAF) above a threshold in any of the populations used will be selected and the corresponding CpG sites will be removed from the `methrix` object. With the `reduce_filtering` option, SNPs with MAF < 0.1 will be further evaluated. If they show low variance in the dataset, there is probably no genotype variability in the population, therefore the corresponding CpG site won't be removed. Please keep in mind that variance thresholds are

Value

methrix object or a list of methrix objects

Examples

```
data('methrix_data')
remove_snps(m = methrix_data, maf_threshold=0.01)
```

remove_uncovered	<i>Remove loci that are uncovered across all samples</i>
------------------	--

Description

Remove loci that are uncovered across all samples

Usage

```
remove_uncovered(m)
```

Arguments

m [methrix](#) object

Details

Takes [methrix](#) object and removes loci that are uncovered across all samples

Value

An object of class [methrix](#)

Examples

```
data('methrix_data')
remove_uncovered(m = methrix_data)
```

save_HDF5_methrix	<i>Saves HDF5 methrix object</i>
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Description

Saves HDF5 methrix object

Usage

```
save_HDF5_methrix(m = NULL, dir = NULL, replace = FALSE, ...)
```

Arguments

m	methrix object
dir	The directory to use. Created, if not existing. Default NULL
replace	Should it overwrite the pre-existing data? FALSE by default.
...	Parameters to pass to saveHDF5SummarizedExperiment

Details

Takes [methrix](#) object and saves it

Value

Nothing

Examples

```
data('methrix_data')
methrix_data_h5 <- convert_methrix(m=methrix_data)
target_dir = paste0(getwd(), '/temp/')
save_HDF5_methrix(methrix_data_h5, dir = target_dir, replace = TRUE)
```

subset_methrix *Subsets [methrix](#) object based on given conditions.*

Description

Subsets [methrix](#) object based on given conditions.

Usage

```
subset_methrix(
  m,
  regions = NULL,
  contigs = NULL,
  samples = NULL,
  overlap_type = "within"
)
```

Arguments

m	methrix object
regions	genomic regions to subset by. Could be a data.table with 3 columns (chr, start, end) or a GenomicRanges object
contigs	chromosome names to subset by
samples	sample names to subset by
overlap_type	defines the type of the overlap of the CpG sites with the target region. Default value is 'within'. For detailed description, see the overlaps function of the data.table package.

Details

Takes `methrix` object and filters CpGs based on coverage statistics

Value

An object of class `methrix`

Examples

```
data('methrix_data')
#Subset to chromosome 1
subset_methrix(methrix_data, contigs = 'chr21')
```

<code>write_bedgraphs</code>	<i>Writes bedGraphs from methrix object</i>
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Description

Writes bedGraphs from methrix object

Usage

```
write_bedgraphs(
  m,
  output_dir = NULL,
  rm_NA = TRUE,
  force = FALSE,
  n_thr = 4,
  compress = TRUE,
  SeqStyle = "UCSC",
  multiBed = NULL,
  metilene = FALSE,
  phenoCol = NULL
)
```

Arguments

<code>m</code>	<code>methrix</code> object
<code>output_dir</code>	Output directory name where the files should be saved. If NULL creates a tempdir
<code>rm_NA</code>	remove NAs
<code>force</code>	forces to create files if they are existing
<code>n_thr</code>	Default 4.
<code>compress</code>	Whether to compress the output. Default TRUE
<code>SeqStyle</code>	Default 'UCSC' with 'chr' prefix.
<code>multiBed</code>	Default NULL. If provided a filename, a single bedGraph file with all samples included is generated.
<code>metilene</code>	Default FALSE. If TRUE outputs bedgraphs in 'metilene' format that can be directly used for DMR calling with 'metilene'. This option works only when <code>multiBed = TRUE</code> .
<code>phenoCol</code>	Default NULL. 'condition' column from colData. Only applicable if <code>metilene = TRUE</code>

Value

writes bedgraph files to output

Examples

```
data('methrix_data')
write_bedgraphs(m = methrix_data, output_dir = './temp')
#Export to metline format for DMR calling with metline
write_bedgraphs(m = methrix_data, output_dir = "./temp", rm_NA = FALSE, metilene = TRUE, multiBed = "metline_ip")
```

write_bigwigs	<i>Exports methrix object as bigWigs</i>
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Description

Exports methrix object as bigWigs

Usage

```
write_bigwigs(m, output_dir = getwd(), samp_names = NULL)
```

Arguments

m	<code>methrix</code> object
output_dir	Output directory name where the files should be saved. Default getwd()
samp_names	sample names to export

Examples

```
data('methrix_data')
write_bigwigs(m = methrix_data, output_dir = './temp')
```

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