

Package ‘NanoMethViz’

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Type Package

Title Visualise methylation data from Oxford Nanopore sequencing

Version 2.6.0

Description NanoMethViz is a toolkit for visualising methylation data from Oxford Nanopore sequencing. It can be used to explore methylation patterns from reads derived from Oxford Nanopore direct DNA sequencing with methylation called by callers including nanoprocess, f5c and megalodon. The plots in this package allow the visualisation of methylation profiles aggregated over experimental groups and across classes of genomic features.

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URL <https://github.com/shians/NanoMethViz>

BugReports <https://github.com/Shians/NanoMethViz/issues>

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NanoMethViz-package *NanoMethViz: Visualise methylation data from Oxford Nanopore sequencing*

Description

NanoMethViz is a toolkit for visualising methylation data from Oxford Nanopore sequencing. It can be used to explore methylation patterns from reads derived from Oxford Nanopore direct DNA sequencing with methylation called by callers including nanopolish, f5c and megalodon. The plots in this package allow the visualisation of methylation profiles aggregated over experimental groups and across classes of genomic features.

Details

The main plotting functions in this package are `plot_gene()` and `plot_region()`.

- See `vignette("ImportingData", package = "NanoMethViz")` for how to import data from Nanopolish and f5c.
- See `vignette("Introduction", package = "NanoMethViz")` for how to create visualisations using this package.

Author(s)

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See Also

Useful links:

- <https://github.com/shians/NanoMethViz>
- Report bugs at <https://github.com/Shians/NanoMethViz/issues>

bsseq_to_edger *Convert BSseq object to edgeR methylation matrix*

Description

Convert BSseq object to edgeR methylation matrix

Usage

```
bsseq_to_edger(bsseq, regions = NULL)
```

Arguments

<code>bsseq</code>	the BSseq object.
<code>regions</code>	the regions to calculate log-methylation ratios over. If left NULL, ratios will be calculated per site.

Value

a matrix compatible with the edgeR differential methylation pipeline

Examples

```
methy <- system.file("methy_subset.tsv.bgz", package = "NanoMethViz")
bsseq <- methy_to_bsseq(methy)
edger_mat <- bsseq_to_edger(bsseq)
```

bsseq_to_log_methy_ratio *Convert BSseq object to log-methylation-ratio matrix*

Description

Creates a log-methylation-ratio matrix from a BSseq object that is useful for dimensionality reduction plots.

Usage

```
bsseq_to_log_methy_ratio(bsseq, regions = NULL, prior_count = 2)
```

Arguments

<code>bsseq</code>	the BSseq object.
<code>regions</code>	the regions to calculate log-methylation ratios over. If left NULL, ratios will be calculated per site.
<code>prior_count</code>	the prior count added to avoid taking log of 0.

Value

a matrix containing log-methylation-ratios.

Examples

```
nmr <- load_example_nanomethresult()
bsseq <- methy_to_bsseq(nmr)
regions <- exons_to_genes(NanoMethViz::exons(nmr))
log_m_ratio <- bsseq_to_log_methy_ratio(bsseq, regions)
```

cluster_regions

Cluster regions by K-means

Description

Cluster regions by k-means based on their methylation profiles. In order to cluster using k-means the methylation profile of each region is interpolated and sampled at fixed points. The first 10 principal components are used for the k-means clustering. The clustering is best behaved in regions of similar width and CpG density.

Usage

```
cluster_regions(x, regions, centers = 2, grid_method = c("density", "uniform"))
```

Arguments

- | | |
|-------------|--|
| x | the NanoMethResult object. |
| regions | a table of regions containing at least columns chr, strand, start and end. |
| centers | number of centers for k-means, identical to the number of output clusters. |
| grid_method | the method for generating the sampling grid. The default option "density" attempts to create a grid with similar density as the data, "uniform" creates a grid of uniform density. |

Value

the table of regions given by the 'regions' argument with the column 'cluster' added.

Examples

```
nmr <- load_example_nanomethresult()
gene_anno <- exons_to_genes(NanoMethViz::exons(nmr))
# uniform grid due to low number of input features
gene_anno_clustered <- cluster_regions(nmr, gene_anno, centers = 2, grid_method = "uniform")
plot_agg_regions(nmr, gene_anno_clustered, group_col = "cluster")
```

`convert_methy_format` *Convert methylation calls to NanoMethViz format*

Description

Convert methylation calls to NanoMethViz format

Usage

```
convert_methy_format(
  input_files,
  output_file,
  samples = fs::path_ext_remove(fs::path_file(input_files)),
  verbose = TRUE
)
```

Arguments

<code>input_files</code>	the files to convert
<code>output_file</code>	the output file to write results to (must end in .bgz)
<code>samples</code>	the names of samples corresponding to each file
<code>verbose</code>	TRUE if progress messages are to be printed

Value

invisibly returns the output file path, creates a tabix file (.bgz) and its index (.bgz.tbi)

`create_tabix_file` *Create a tabix file using methylation calls*

Description

Create a tabix file using methylation calls

Usage

```
create_tabix_file(
  input_files,
  output_file,
  samples = extract_file_names(input_files),
  verbose = TRUE
)
```

Arguments

input_files	the files to convert
output_file	the output file to write results to (must end in .bgz)
samples	the names of samples corresponding to each file
verbose	TRUE if progress messages are to be printed

Value

invisibly returns the output file path, creates a tabix file (.bgz) and its index (.bgz.tbi)

Examples

```
methyl_calls <- system.file(package = "NanoMethViz",
  c("sample1_nanopolish.tsv.gz", "sample2_nanopolish.tsv.gz"))
temp_file <- paste0(tempfile(), ".tsv.bgz")

create_tabix_file(methyl_calls, temp_file)
```

exons

Get exon annotation

Description

Get exon annotation

Usage

```
exons(object)
```

exons<-

Set exon annotation

Description

Set exon annotation

Usage

```
exons(object) <- value
```

`exons_to_genes` *Convert exon annotation to genes*

Description

Convert exon annotation to genes

Usage

```
exons_to_genes(x)
```

Arguments

- x the exon level annotation containing columns "gene_id", "chr", "strand" and "symbol".

Value

the gene level annotation where each gene is taken to span the earliest start position and latest end position of its exons.

Examples

```
nmr <- load_example_nanomethresult()
exons_to_genes(NanoMethViz::exons(nmr))
```

`filter_methy` *Create filtered methylation file*

Description

Create a filtered methylation file from an existing one.

Usage

```
filter_methy(x, output_file, ...)
```

Arguments

- x the path to the methylation file or a NanoMethResult object.
- output_file the output file to write results to (must end in .bgz).
- ... filtering criteria given in dplyr syntax. Use methy_col_names() to get available column names.

Value

invisibly returns 'output_file' if x is a file path, otherwise returns NanoMethResult object with methy(x) replaced with filtered value.

Examples

```
nmr <- load_example_nanomethresult()
output_file <- paste0(tempfile(), ".tsv.bgz")
filter_methy(nmr, output_file = output_file, chr == "chrX")
filter_methy(methy(nmr), output_file = output_file, chr == "chrX")
```

get_example_exons_mus_musculus

Get example exon annotations for mus musculus (mm10)

Description

This is a small subset of the exons returned by `get_exons_mus_musculus()` for demonstrative purposes. It contains the exons for the genes Brca1, Brca2, Impact, Meg3, Peg3 and Xist.

Usage

```
get_example_exons_mus_musculus()
```

Value

data.frame containing exons

Examples

```
example_exons <- get_example_exons_mus_musculus()
```

get_exons

Get exon annotations

Description

Helper functions are provided for obtaining exon annotations from relevant TxDb packages on Bioconductor for the construction of NanoMethResults objects.

Usage

```
get_exons_mm10()
get_exons_hg19()
get_exons_hg38()
```

Value

tibble (data.frame) object containing exon annotation.

Examples

```
mm10_exons <- get_exons_mm10()  
  
hg19_exons <- get_exons_hg19()  
  
hg38_exons <- get_exons_hg38()
```

get_exons_homo_sapiens

Get exon annotations for Homo sapiens (hg19)

Description

Get exon annotations for Homo sapiens (hg19)

Usage

```
get_exons_homo_sapiens()
```

Value

data.frame containing exons

Examples

```
h_sapiens_exons <- get_exons_homo_sapiens()
```

get_exons_mus_musculus

Get exon annotations for Mus musculus (mm10)

Description

Get exon annotations for Mus musculus (mm10)

Usage

```
get_exons_mus_musculus()
```

Value

data.frame containing exons

Examples

```
m_musculus_exons <- get_exons_mus_musculus()
```

```
load_example_nanomethresult
```

Load an example NanoMethResult object

Description

Load an example NanoMethResult object for demonstration of plotting functions. Run `load_example_nanomethresults` without the function call to see how the object is constructed.

Usage

```
load_example_nanomethresult()
```

Value

a NanoMethResults object

Examples

```
nmr <- load_example_nanomethresult()
```

```
methyl
```

Get methylation data

Description

Get methylation data

Usage

```
methyl(object)
```

Arguments

object the object.

Value

the path to the methylation data.

Examples

```
showMethods("methy")
```

```
methy<-
```

Set methylation data

Description

Set methylation data

Usage

```
methy(object) <- value
```

```
methy_col_names
```

Column names for methylation data

Description

Column names for methylation data

Usage

```
methy_col_names()
```

Value

column names for methylation data

Examples

```
methy_col_names()
```

methy_to_bsseq	<i>Create BSSeq object from methylation tabix file</i>
----------------	--

Description

Create BSSeq object from methylation tabix file

Usage

```
methy_to_bsseq(methy, out_folder = tempdir(), verbose = TRUE)
```

Arguments

methy	the path to the methylation tabix file.
out_folder	the folder to store intermediate files. One file is created for each sample and contains columns "chr", "pos", "total" and "methylated".
verbose	TRUE if progress messages are to be printed

Value

a BSSeq object.

Examples

```
nmr <- load_example_nanomethresult()
bsseq <- methy_to_bsseq(nmr)
```

methy_to_edger	<i>Convert NanoMethResult object to edgeR methylation matrix</i>
----------------	--

Description

Convert NanoMethResult object to edgeR methylation matrix

Usage

```
methy_to_edger(methy, regions = NULL, out_folder = tempdir(), verbose = TRUE)
```

Arguments

methy	the path to the methylation tabix file.
regions	the regions to calculate log-methylation ratios over. If left NULL, ratios will be calculated per site.
out_folder	the folder to store intermediate files. One file is created for each sample and contains columns "chr", "pos", "total" and "methylated".
verbose	TRUE if progress messages are to be printed

Value

a matrix compatible with the edgeR differential methylation pipeline

Examples

```
nmr <- load_example_nanomethresult()
edger_mat <- methy_to_edger(nmr)
```

ModBamFiles

*Constructor for a ModBamFiles object***Description**

This function creates a ModBamFiles object containing information about the samples and file paths.

Usage

```
ModBamFiles(samples, paths)

## S4 method for signature 'ModBamFiles'
show(object)
```

Arguments

- | | |
|---------|---|
| samples | a character vector with the names of the samples. |
| paths | a character vector with the file paths for the BAM files. |
| object | a ModBamFiles object. |

Value

A ModBamFiles object with the sample and path information.

ModBamFiles-class

*ModBamFiles class***Description**

This is a class for holding information about modbam files. It is a data.frame containing information about samples and paths to modbam files.

ModBamResult-class *Modbam methylation results*

Description

(Experimental) A ModBamResult object stores modbam data used for NanoMethViz visualisation.

Usage

```
## S4 method for signature 'ModBamResult'
methy(object)

## S4 replacement method for signature 'ModBamResult,ModBamFiles'
methy(object) <- value

## S4 method for signature 'ModBamResult'
samples(object)

## S4 replacement method for signature 'ModBamResult,data.frame'
samples(object) <- value

## S4 method for signature 'ModBamResult'
exons(object)

## S4 replacement method for signature 'ModBamResult,data.frame'
exons(object) <- value

ModBamResult(methy, samples, exons = NULL)
```

Arguments

<code>object</code>	the ModBamResult object.
<code>value</code>	the exon annotation.
<code>methy</code>	a ModBamFiles object.
<code>samples</code>	the data.frame of sample annotation containing at least columns sample and group.
<code>exons</code>	(optional) the data.frame of exon information containing at least columns gene_id, chr, strand, start, end, transcript_id and symbol.

Value

a NanoMethResult object to be used with plotting functions
 a ModBamFiles data.frame.
 the sample annotation.
 the exon annotation.
 the exon annotation.

Functions

- `methy(ModBamResult)`: modbam information getter.
- `methy(object = ModBamResult) <- value`: modbam information setter.
- `samples(ModBamResult)`: sample annotation getter.
- `samples(object = ModBamResult) <- value`: sample annotation setter.
- `exons(ModBamResult)`: exon annotation getter.
- `exons(object = ModBamResult) <- value`: exon annotation setter.
- `ModBamResult()`: Constructor

Slots

`methy` a ModBamFiles data.frame specifying the samples and paths to bam files.
`samples` the data.frame of sample annotation containing at least columns sample and group.
`exons` the data.frame of exon information containing at least columns gene_id, chr, strand, start, end, transcript_id and symbol.

NanoMethResult-class *Nanopore Methylation Result*

Description

A NanoMethResult object stores data used for NanoMethViz visualisation. It contains stores a path to the methylation data, sample information and optional exon information. The object is constructed using the NanoMethResult() constructor function described in "Usage".

Usage

```
NanoMethResult(methy, samples, exons = NULL)

## S4 method for signature 'NanoMethResult'
methy(object)

## S4 replacement method for signature 'NanoMethResult,ANY'
methy(object) <- value

## S4 method for signature 'NanoMethResult'
samples(object)

## S4 replacement method for signature 'NanoMethResult,data.frame'
samples(object) <- value

## S4 method for signature 'NanoMethResult'
exons(object)

## S4 replacement method for signature 'NanoMethResult,data.frame'
exons(object) <- value
```

Arguments

<code>methy</code>	the path to the methylation tabix file.
<code>samples</code>	the data.frame of sample annotation containing at least columns sample and group.
<code>exons</code>	(optional) the data.frame of exon information containing at least columns gene_id, chr, strand, start, end, transcript_id and symbol.
<code>object</code>	the NanoMethResult object.
<code>value</code>	the exon annotation.

Value

a NanoMethResult object to be used with plotting functions
 the path to the methylation data.
 the sample annotation.
 the exon annotation.

Functions

- `NanoMethResult()`: Constructor
- `methy(NanoMethResult)`: methylation data path getter.
- `methy(object = NanoMethResult) <- value`: methylation data path setter.
- `samples(NanoMethResult)`: sample annotation getter.
- `samples(object = NanoMethResult) <- value`: sample annotation setter.
- `exons(NanoMethResult)`: exon annotation getter.
- `exons(object = NanoMethResult) <- value`: exon annotation setter.

Slots

`methy` the path to the methylation tabix file.
`samples` the data.frame of sample annotation containing at least columns sample and group.
`exons` the data.frame of exon information containing at least columns gene_id, chr, strand, start, end, transcript_id and symbol.

Examples

```

methy <- system.file(package = "NanoMethViz", "methy_subset.tsv.bgz")
sample <- c(
  "B6Cast_Prom_1_b16",
  "B6Cast_Prom_1_cast",
  "B6Cast_Prom_2_b16",
  "B6Cast_Prom_2_cast",
  "B6Cast_Prom_3_b16",
  "B6Cast_Prom_3_cast"
)
group <- c(

```

```

"bl6",
"cast",
"bl6",
"cast",
"bl6",
"cast"
)
sample_anno <- data.frame(sample, group, stringsAsFactors = FALSE)
exon_tibble <- get_example_exons_mus_musculus()
NanoMethResult(methy, sample_anno, exon_tibble)

x <- load_example_nanomethresult()
methy(x)

```

plot_agg_genes *Plot gene aggregate plot*

Description

Plot gene aggregate plot

Usage

```
plot_agg_genes(
  x,
  genes = NULL,
  binary_threshold = 0.5,
  group_col = NULL,
  flank = 2000,
  stranded = TRUE,
  span = 0.05,
  palette = ggplot2::scale_colour_brewer(palette = "Set1")
)
```

Arguments

- x** the NanoMethResult object.
- genes** a character vector of genes to include in aggregate plot, if NULL then all genes are used.
- binary_threshold** the modification probability such that calls with modification probability above the threshold are considered methylated, and those with probability equal or below are considered unmethylated.
- group_col** the column to group aggregated trends by. This column can be in from the regions table or samples(x).
- flank** the number of flanking bases to add to each side of each region.

stranded	TRUE if negative strand features should have coordinates flipped to reflect features like transcription start sites.
span	the span for loess smoothing.
palette	the ggplot colour palette used for groups.

Value

a ggplot object containing the aggregate methylation trend of genes.

Examples

```
nmr <- load_example_nanomethresult()
plot_agg_genes(nmr)
```

plot_agg_regions	<i>Plot aggregate regions</i>
------------------	-------------------------------

Description

Plot aggregate regions

Usage

```
plot_agg_regions(
  x,
  regions,
  binary_threshold = 0.5,
  group_col = NULL,
  flank = 2000,
  stranded = TRUE,
  span = 0.05,
  palette = ggplot2::scale_colour_brewer(palette = "Set1")
)
```

Arguments

x	the NanoMethResult object.
regions	a table of regions containing at least columns chr, strand, start and end. Any additional columns can be used for grouping.
binary_threshold	the modification probability such that calls with modification probability above the threshold are considered methylated, and those with probability equal or below are considered unmethylated.
group_col	the column to group aggregated trends by. This column can be in from the regions table or samples(x).

<code>flank</code>	the number of flanking bases to add to each side of each region.
<code>stranded</code>	TRUE if negative strand features should have coordinates flipped to reflect features like transcription start sites.
<code>span</code>	the span for loess smoothing.
<code>palette</code>	the ggplot colour palette used for groups.

Value

a ggplot object containing the aggregate methylation trend.

Examples

```
nmr <- load_example_nanomethresult()
gene_anno <- exons_to_genes(NanoMethViz::exons(nmr))
plot_agg_regions(nmr, gene_anno)
plot_agg_regions(nmr, gene_anno, group_col = "sample")
plot_agg_regions(nmr, gene_anno, group_col = "group")
```

`plot_gene`

Plot gene methylation

Description

Plot the methylation of a gene symbol specified within the exon(x) slot.

Usage

```
plot_gene(x, gene, ...)

## S4 method for signature 'NanoMethResult,character'
plot_gene(
  x,
  gene,
  window_prop = 0.3,
  anno_regions = NULL,
  binary_threshold = NULL,
  avg_method = c("mean", "median"),
  spaghetti = FALSE,
  heatmap = FALSE,
  span = NULL,
  gene_anno = TRUE,
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),
  line_size = 2
)
## S4 method for signature 'ModBamResult,character'
```

```
plot_gene(
  x,
  gene,
  window_prop = 0.3,
  anno_regions = NULL,
  binary_threshold = NULL,
  avg_method = c("mean", "median"),
  spaghetti = FALSE,
  heatmap = FALSE,
  span = NULL,
  gene_anno = TRUE,
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),
  line_size = 2
)
```

Arguments

<code>x</code>	the NanoMethResult or ModBamResult object.
<code>gene</code>	the gene symbol for the gene to plot.
<code>...</code>	additional arguments
<code>window_prop</code>	the size of flanking region to plot. Can be a vector of two values for left and right window size. Values indicate proportion of gene length.
<code>anno_regions</code>	the data.frame of regions to annotate.
<code>binary_threshold</code>	the modification probability such that calls with modification probability above the threshold are set to 1 and probabilities equal to or below the threshold are set to 0.
<code>avg_method</code>	the average method for pre-smoothing at each genomic position. Data is pre-smoothed at each genomic position before the smoothed aggregate line is generated for performance reasons. The default is "mean" which corresponds to the average methylation fraction. The alternative "median" option is closer to an average within the more common methylation state.
<code>spaghetti</code>	whether or not individual reads should be shown.
<code>heatmap</code>	whether or not read-methylation heatmap should be shown.
<code>span</code>	the span for loess smoothing.
<code>gene_anno</code>	whether or not gene annotation tracks are plotted.
<code>palette</code>	the ggplot colour palette used for groups.
<code>line_size</code>	the size of the lines.

Value

a patchwork plot containing the methylation profile in the specified region.

Functions

- `plot_gene(x = ModBamResult, gene = character)`: S4 method for ModBamResult

Examples

```
nmr <- load_example_nanomethresult()
plot_gene(nmr, "Peg3")
```

plot_gene_heatmap *Plot gene methylation heatmap*

Description

Plot the methylation heatmap of a gene symbol specified within the exon(x) slot.

Usage

```
plot_gene_heatmap(x, gene, ...)

## S4 method for signature 'NanoMethResult,character'
plot_gene_heatmap(
  x,
  gene,
  window_prop = 0.3,
  pos_style = c("to_scale", "compact"),
  subsample = 50
)

## S4 method for signature 'ModBamResult,character'
plot_gene_heatmap(
  x,
  gene,
  window_prop = 0.3,
  pos_style = c("to_scale", "compact"),
  subsample = 50
)
```

Arguments

x	the NanoMethResult or ModBamResult object.
gene	the gene symbol for the gene to plot.
...	additional arguments
window_prop	the size of flanking region to plot. Can be a vector of two values for left and right window size. Values indicate proportion of gene length.
pos_style	the style for plotting the base positions along the x-axis. Defaults to "to_scale", plotting (potentially) overlapping squares along the genomic position to scale. The "compact" options plots only the positions with measured modification.
subsample	the number of read of packed read rows to subsample to.

Value

a ggplot object of the heatmap
a ggplot plot containing the heatmap.

Examples

```
nmr <- load_example_nanomethresult()  
plot_gene_heatmap(nmr, "Peg3")
```

plot_grange*Plot GRanges***Description**

Plot GRanges

Usage

```
plot_grange(  
  x,  
  grange,  
  anno_regions = NULL,  
  binary_threshold = NULL,  
  avg_method = c("mean", "median"),  
  spaghetti = FALSE,  
  heatmap = FALSE,  
  span = NULL,  
  window_prop = 0,  
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),  
  line_size = 2  
)
```

Arguments

<code>x</code>	the NanoMethResult object.
<code>grange</code>	the GRanges object with one entry.
<code>anno_regions</code>	the data.frame of regions to be annotated.
<code>binary_threshold</code>	the modification probability such that calls with modification probability above the threshold are set to 1 and probabilities equal to or below the threshold are set to 0.
<code>avg_method</code>	the average method for pre-smoothing at each genomic position. Data is pre-smoothed at each genomic position before the smoothed aggregate line is generated for performance reasons. The default is "mean" which corresponds to the average methylation fraction. The alternative "median" option is closer to an average within the more common methylation state.

<code>spaghetti</code>	whether or not individual reads should be shown.
<code>heatmap</code>	whether or not read-methylation heatmap should be shown.
<code>span</code>	the span for loess smoothing.
<code>window_prop</code>	the size of flanking region to plot. Can be a vector of two values for left and right window size. Values indicate proportion of gene length.
<code>palette</code>	the ggplot colour palette used for groups.
<code>line_size</code>	the size of the lines.

Value

a patchwork plot containing the methylation profile in the specified region.

Examples

```
nmr <- load_example_nanomethresult()
plot_grange(nmr, GenomicRanges::GRanges("chr7:6703892-6730431"))
```

`plot_grange_heatmap` *Plot GRanges heatmap*

Description

Plot GRanges heatmap

Usage

```
plot_grange_heatmap(
  x,
  grange,
  pos_style = c("to_scale", "compact"),
  window_prop = 0,
  subsample = 50
)
```

Arguments

<code>x</code>	the NanoMethResult object.
<code>grange</code>	the GRanges object with one entry.
<code>pos_style</code>	the style for plotting the base positions along the x-axis. Defaults to "to_scale", plotting (potentially) overlapping squares along the genomic position to scale. The "compact" options plots only the positions with measured modification.
<code>window_prop</code>	the size of flanking region to plot. Can be a vector of two values for left and right window size. Values indicate proportion of gene length.
<code>subsample</code>	the number of read of packed read rows to subsample to.

Value

a ggplot plot containing the heatmap.

Examples

```
nmr <- load_example_nanomethresult()
plot_grange_heatmap(nmr, GenomicRanges::GRanges("chr7:6703892-6730431"))
```

plot_mds

*Plot MDS***Description**

Plot multi-dimensional scaling plot using algorithm of limma::plotMDS(). It is recommended this be done with the log-methylation-ratio matrix generated by bsseq_to_log_methy_ratio().

Usage

```
plot_mds(
  x,
  top = 500,
  plot_dims = c(1, 2),
  labels = colnames(x),
  groups = NULL
)
```

Arguments

<code>x</code>	the log-methylation-ratio matrix.
<code>top</code>	the number of top genes used to calculate pairwise distances.
<code>plot_dims</code>	the numeric vector of the two dimensions to be plotted.
<code>labels</code>	the character vector of labels for data points. By default uses column names of <code>x</code> , set to <code>NULL</code> to plot points.
<code>groups</code>	the character vector of groups the data points will be coloured by. Colour palette can be adjusted using <code>scale_colour_*</code> () functions from ggplot2.

Value

ggplot object of the MDS plot.

Examples

```
nmr <- load_example_nanomethresult()
bss <- methy_to_bsseq(nmr)
lmr <- bsseq_to_log_methy_ratio(bss)
plot_mds(lmr)
```

`plot_pca`*Plot PCA***Description**

Plot multi-dimensional scaling plot using algorithm of BiocSingular::runPCA(). It is recommended this be done with the log-methylation-ratio matrix generated by bsseq_to_log_methy_ratio().

Usage

```
plot_pca(x, plot_dims = c(1, 2), labels = colnames(x), groups = NULL)
```

Arguments

- `x` the log-methylation-ratio matrix.
- `plot_dims` the numeric vector of the two dimensions to be plotted.
- `labels` the character vector of labels for data points. By default uses column names of `x`, set to `NULL` to plot points.
- `groups` the character vector of groups the data points will be coloured by.

Value

ggplot object of the MDS plot.

Examples

```
nmr <- load_example_nanomethresult()
bss <- methy_to_bsseq(nmr)
lmr <- bsseq_to_log_methy_ratio(bss)
plot_pca(lmr)
```

`plot_region`*Plot region methylation***Description**

Plot the methylation of a genomic region.

Usage

```
plot_region(x, chr, start, end, ...)

## S4 method for signature 'NanoMethResult,character,numeric,numeric'
plot_region(
  x,
  chr,
  start,
  end,
  anno_regions = NULL,
  binary_threshold = NULL,
  avg_method = c("mean", "median"),
  spaghetti = FALSE,
  heatmap = FALSE,
  span = NULL,
  window_prop = 0,
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),
  line_size = 2
)

## S4 method for signature 'ModBamResult,character,numeric,numeric'
plot_region(
  x,
  chr,
  start,
  end,
  anno_regions = NULL,
  binary_threshold = NULL,
  avg_method = c("mean", "median"),
  spaghetti = FALSE,
  heatmap = FALSE,
  span = NULL,
  window_prop = 0,
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),
  line_size = 2
)

## S4 method for signature 'NanoMethResult,factor,numeric,numeric'
plot_region(
  x,
  chr,
  start,
  end,
  anno_regions = NULL,
  binary_threshold = NULL,
  avg_method = c("mean", "median"),
  spaghetti = FALSE,
  heatmap = FALSE,
```

```

plot_region

  span = NULL,
  window_prop = 0,
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),
  line_size = 2
)

## S4 method for signature 'ModBamResult,factor,numeric,numeric'
plot_region(
  x,
  chr,
  start,
  end,
  anno_regions = NULL,
  binary_threshold = NULL,
  avg_method = c("mean", "median"),
  spaghetti = FALSE,
  heatmap = FALSE,
  span = NULL,
  window_prop = 0,
  palette = ggplot2::scale_colour_brewer(palette = "Set1"),
  line_size = 2
)

```

Arguments

<code>x</code>	the NanoMethResult or ModBamResult object.
<code>chr</code>	the chromosome to plot.
<code>start</code>	the start of the plotting region.
<code>end</code>	the end of the plotting region.
<code>...</code>	additional arguments.
<code>anno_regions</code>	the data.frame of regions to be annotated.
<code>binary_threshold</code>	the modification probability such that calls with modification probability above the threshold are set to 1 and probabilities equal to or below the threshold are set to 0.
<code>avg_method</code>	the average method for pre-smoothing at each genomic position. Data is pre-smoothed at each genomic position before the smoothed aggregate line is generated for performance reasons. The default is "mean" which corresponds to the average methylation fraction. The alternative "median" option is closer to an average within the more common methylation state.
<code>spaghetti</code>	whether or not individual reads should be shown.
<code>heatmap</code>	whether or not read-methylation heatmap should be shown.
<code>span</code>	the span for loess smoothing.
<code>window_prop</code>	the size of flanking region to plot. Can be a vector of two values for left and right window size. Values indicate proportion of gene length.
<code>palette</code>	the ggplot colour palette used for groups.
<code>line_size</code>	the size of the lines.

Value

a patchwork plot containing the methylation profile in the specified region.

Examples

```
nmr <- load_example_nanomethresult()
plot_region(nmr, "chr7", 6703892, 6730431)
```

plot_region_heatmap *Plot region methylation heatmap*

Description

Plot the methylation heatmap of a genomic region.

Usage

```
plot_region_heatmap(x, chr, start, end, ...)

## S4 method for signature 'NanoMethResult,character,numeric,numeric'
plot_region_heatmap(
  x,
  chr,
  start,
  end,
  pos_style = c("to_scale", "compact"),
  window_prop = 0,
  subsample = 50
)

## S4 method for signature 'ModBamResult,character,numeric,numeric'
plot_region_heatmap(
  x,
  chr,
  start,
  end,
  pos_style = c("to_scale", "compact"),
  window_prop = 0,
  subsample = 50
)

## S4 method for signature 'NanoMethResult,factor,numeric,numeric'
plot_region_heatmap(
  x,
  chr,
  start,
```

```

    end,
    pos_style = c("to_scale", "compact"),
    window_prop = 0,
    subsample = 50
  )

## S4 method for signature 'ModBamResult,factor,numeric,numeric'
plot_region_heatmap(
  x,
  chr,
  start,
  end,
  pos_style = c("to_scale", "compact"),
  window_prop = 0,
  subsample = 50
)

```

Arguments

<code>x</code>	the NanoMethResult or ModBamResult object.
<code>chr</code>	the chromosome to plot.
<code>start</code>	the start of the plotting region.
<code>end</code>	the end of the plotting region.
<code>...</code>	additional arguments.
<code>pos_style</code>	the style for plotting the base positions along the x-axis. Defaults to "to_scale", plotting (potentially) overlapping squares along the genomic position to scale. The "compact" option plots only the positions with measured modification.
<code>window_prop</code>	the size of flanking region to plot. Can be a vector of two values for left and right window size. Values indicate proportion of gene length.
<code>subsample</code>	the number of read packed read rows to subsample to.

Value

- a ggplot object of the heatmap.
 a ggplot plot containing the heatmap.

Examples

```

nmr <- load_example_nanomethresult()
plot_region_heatmap(nmr, "chr7", 6703892, 6730431)

```

query_exons

Query exons

Description

Query a data.frame of exons for a subset.

Usage

```
query_exons_region(exons, chr, start, end)  
query_exons_gene_id(exons, gene_id)  
query_exons_symbol(exons, symbol)
```

Arguments

exons	the data.frame of exons.
chr	the chromosome to query.
start	the start of the query region.
end	the end of the query region.
gene_id	the gene_id to query.
symbol	the gene_id to query.

Value

data.frame of queried exons.

Functions

- `query_exons_region()`: Query region.
- `query_exons_gene_id()`: Query gene ID.
- `query_exons_symbol()`: Query gene symbol.

<code>query_methy</code>	<i>Query methylation data</i>
--------------------------	-------------------------------

Description

Query methylation data

Usage

```
query_methy(x, chr, start, end, simplify = TRUE, force = FALSE)
```

Arguments

<code>x</code>	the NanoMethResults object or a path to the methylation data (tabix-bgzipped).
<code>chr</code>	the vector of chromosomes
<code>start</code>	the vector of start positions
<code>end</code>	the vector of end positions
<code>simplify</code>	whether returned results should be row-concatenated
<code>force</code>	whether to force empty output when query region 'chr' does not appear in data. Without 'force', an empty result indicates that the requested 'chr' appears in the data but no data overlaps with requested region, and an invalid 'chr' will cause an error.

Value

A table containing the data within the queried regions. If simplify is TRUE (default) then all data is contained within one table, otherwise it is a list of tables where each element is the data from one region.

Examples

```
nmr <- load_example_nanomethresult()
query_methy(methy(nmr), "chr7", 6703892, 6730431)
```

<code>raw_methy_to_tabix</code>	<i>Convert methylation file to tabix format</i>
---------------------------------	---

Description

Convert methylation file to tabix format

Usage

```
raw_methy_to_tabix(x)
```

Arguments

- x the path to the sorted methylation file

Value

invisibly returns the path to the tabix file

reexports*Objects exported from other packages*

Description

These objects are imported from other packages. Follow the links below to see their documentation.

[e1071 sigmoid](#)

region_methy_stats*Calculate region methylation statistics*

Description

Calculate the average methylation probability and prevalence based on specified probability threshold.

Usage

```
region_methy_stats(nmr, regions, threshold = 0.5)
```

Arguments

- | | |
|-----------|---|
| nmr | the NanoMethResult object. |
| regions | the table of regions to query statistics for. |
| threshold | the threshold to use for determining methylation calls for the calculation of prevalence. |

Value

table of regions with additional columns of methylation summary statistics.

Examples

```
nmr <- load_example_nanomethresult()
gene_anno <- exons_to_genes(NanoMethViz::exons(nmr))
region_methy_stats(nmr, gene_anno)
```

samples	<i>Get sample annotation</i>
---------	------------------------------

Description

Get sample annotation

Usage

```
samples(object)
```

samples<-	<i>Set sample annotation</i>
-----------	------------------------------

Description

Set sample annotation

Usage

```
samples(object) <- value
```

sort_methy_file	<i>Sort methylation file</i>
-----------------	------------------------------

Description

Sort methylation file

Usage

```
sort_methy_file(x)
```

Arguments

x the path to the methylation file to sort

Value

invisibly returns path of sorted file

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