# Package 'profileplyr'

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Type Package

**Title** Visualization and annotation of read signal over genomic ranges with profileplyr

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**Depends** R (>= 3.6), BiocGenerics, SummarizedExperiment

# Description

Quick and straightforward visualization of read signal over genomic intervals is key for generating hypotheses from sequencing data sets (e.g. ChIP-seq, ATAC-seq, bisulfite/methyl-seq). Many tools both inside

and outside of R and Bioconductor are available to explore these types of data, and they typically start

with a bigWig or BAM file and end with some representation of the signal (e.g. heatmap). profileplyr leverages

many Bioconductor tools to allow for both flexibility and additional functionality in work-flows that end with

visualization of the read signal.

**License** GPL (>= 3) **RoxygenNote** 7.1.2

biocViews ChIPSeq, DataImport, Sequencing, ChipOnChip, Coverage

Imports GenomicRanges, stats, soGGi, methods, utils, S4Vectors,

R.utils, dplyr, magrittr, tidyr, IRanges, rjson,

ChIPseeker, GenomicFeatures, TxDb. Hsapiens. UCSC.hg19.knownGene, TxDb. Hsapiens. UCSC.hg38.knownGene, TxDb. Mrusculus. UCSC.mm9.knownGene, org. Hs.eg.db, org. Mm.eg.db, rGREAT,

pheatmap, ggplot2, EnrichedHeatmap, ComplexHeatmap, grid,

circlize, BiocParallel, rtracklayer, GenomeInfoDb, grDevices,

rlang, tiff, Rsamtools

Suggests BiocStyle, testthat, knitr, rmarkdown, png, Cairo

VignetteBuilder knitr

2 annotateRanges

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# R topics documented:

annotateRanges	2
annotateRanges_great	4
as_profileplyr	5
BamBigwig_to_chipProfile	6
clusterRanges	7
convertToEnrichedHeatmapMat	10
export_deepToolsMat	11
generateEnrichedHeatmap	12
generateProfilePlot	17
gene_list_character	18
gene_list_dataframe	18
groupBy	19
inherit_group_function	21
K27ac_GRlist_hind_liver_top5000	22
orderBy	
params	23
profileplyr-class	
sampleData	24
subsetbyGeneListOverlap	
subsetbyRangeOverlap	27
subset_GR_GL_common_top	28
summarize	29
	31

annotate Ranges

Annotate profileplyr ranges to genes using ChIPseeker

# Description

Index

The ranges from the deepTools matrix will be subset based on whether they overlap with specified annotated regions (using ChIPseeker) throughout the genome

annotateRanges 3

#### Usage

```
annotateRanges(
  object = "profileplyr",
  annotation_subset = "character",
  annoDb = "character"
  tssRegion = "numeric",
  changeGroupToAnnotation = "logical",
  heatmap_grouping = "character",
)
## S4 method for signature 'profileplyr'
annotateRanges(
  object = "profileplyr";
  annotation_subset = NULL,
  TxDb = NULL.
  annoDb = NULL
  tssRegion = c(-3000, 3000),
  changeGroupToAnnotation = FALSE,
  heatmap_grouping = "group",
)
```

# Arguments

object A profileplyr object

annotation\_subset

If specific annotations (from ChIPseeker package) are desired, specify them here in a character vector. Can be one or any combination of "Promoter", "Exon", "Intron", "Downstream", "Distal Intergenic", "3p UTR", or "5p UTR". This argument is optional and all annotation types will be included if argument is left out.

TxDb

This must be either a TxDb object, a character string that is a path to a GTF file, or character string indicating genome if one of the following - "hg19", "hg38", "mm9", "mm10".

annoDb

The annotation package to be used. If the 'TxDb' argument is set to "hg19", "hg38", "mm9", or "mm10" this will automatically be set and this can be left as NULL.

tssRegion

This needs to be a vector of two numbers that will define promoter regions. The first number must be negative, while the second number must be positive. Default values are c(-3000, 3000)

changeGroupToAnnotation

If the grouping should be changed to the annotations (typically when the ranges will be exported for visualization based on this annotation), this should be TRUE. The default if FALSE, which will keep the grouping that existed before annotat-

ing the object. This is typical if the output will be used for finding overlaps with gene lists in the 'groupBy' function.

heatmap\_grouping

Only relevant if 'keepAnnotationAsGroup' is set to TRUE. This argument needs to be either "group", or "annotation". This will determine how the ranges are grouped in the resulting object. Default is heatmap\_grouping = "Group". If there are no groups in the deepTools matrix that was used in the function, this argument is unnecessary

... pass to annotatePeak

#### **Details**

tbd

#### Value

A profileplyr object

#### Methods (by class)

• profileplyr: Annotate profileplyr ranges to genes using ChIPseeker

#### **Examples**

```
library(SummarizedExperiment)
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
object <- object[1:2, , ]

# annotate ranges with genes using ChIPseeker
# (NOTE: can choose subset of annotations with 'annotation_subset' argument)
annotateRanges(object, TxDb = "mm10")</pre>
```

annotateRanges\_great Annotate profileplyr ranges to genes using rGREAT

### **Description**

The ranges from the deepTools matrix will be subset based on whether they overlap with specified annotated regions related to a user defined gene list.

#### Usage

```
annotateRanges_great(object = "profileplyr", species = "character", ...)
## S4 method for signature 'profileplyr'
annotateRanges_great(object = "profileplyr", species = "character", ...)
```

as\_profileplyr 5

# Arguments

```
object A profileplyr object species GREAT accepts "hg19", "mm10", "mm9", "danRer7" (zebrafish)
```

... pass to submitGreatJob

#### **Details**

tbd

#### Value

A profileplyr object

# Methods (by class)

• profileplyr: Annotate profileplyr ranges to genes using rGREAT

# **Examples**

```
library(SummarizedExperiment)
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
object <- object[1:5, , ]

# annotate ranges with genes using GREAT with following command:
annotateRanges_great(object, species = "mm10")</pre>
```

as\_profileplyr

Import ChIP profile object to profile plyr

# Description

Function to convert soGGi ChIPprofile objects to profileplyr object.

#### Usage

```
as_profileplyr(chipProfile, names = NULL)
```

### **Arguments**

chipProfile A ChIPprofile object as created by soGGi regionPlot() function.

names Column to select row IDs/names from ChIPprofile mcols.

### Value

A profileplyr object

#### **Examples**

```
library(soGGi)
data("ik_Profiles")
proplyr <- as_profileplyr(ik_Profiles,names="ID")
export_deepToolsMat(proplyr,con=file.path(tempdir(),"ik_Profiles.MAT"))</pre>
```

BamBigwig\_to\_chipProfile

BamBigwig\_to\_chipProfile

#### **Description**

Generate a soGGi ChIPprofile object with multiple BAM/bigWig files or multiple BED files as the input

#### Usage

```
BamBigwig_to_chipProfile(
    signalFiles,
    testRanges,
    format,
    style = "percentOfRegion",
    nOfWindows = 100,
    bin_size = 20,
    distanceAround = NULL,
    distanceUp = 1000,
    distanceDown = 1000,
    ...,
    quant_params = NULL
)
```

#### **Arguments**

signalFiles paths to either BAM files or bigwig files. More than one path can be in this

character vector, but all paths in one function call must point to be either all

BAM files or all bigWig files, not a combination of the two.

testRanges A character vector with paths to BED files.

format character string of "bam", "bigwig", "RleList" or "PWM"

style a character string, "percentOfRegion" (default) for normalized length divided

into bins set by the 'nOfWindows' argument, "point" for per base pair plot where the number of base pairs per bin is set by the 'bin\_size' argument, and "region"

for combined plot

nOfWindows The number of windows/bins the normalised ranges will be divided into if 'style'

is set to 'percentOfRegion'. Default is 100.

bin\_size If 'style' is set to 'point' then this will determine the size of each bin over which

signal is quantified. The default is 20 base pairs.

clusterRanges 7

distanceAround This controls the distance around the region that is included. If 'style' is 'per-

centOfRegion', then the default is 100, meaning that a distance equal to 100 percent of that particular region on either side of the region will be included in the heatmap. If 'style' is 'point',then this is the number of basepairs from the center of each range, in either direction, that the heatmap will show. If style is 'point' and 'distanceAround' is NULL, then distanceUp and distanceDown will

be used.

upstream from the center of each peak signal will be quantified. If the 'distanceAround' argument is set (i.e. not NULL), that will be used for the quantifi-

cation range and 'distanceUp will be ignored.

distanceDown If 'style' is set to 'point' then this will determine the distance (in base pairs)

downstream from the center of each peak signal will be quantified. If the 'distanceAround' argument is set (i.e. not NULL), that will be used for the quantifi-

cation range and 'distanceDown' will be ignored.

... pass to regionPlot() within the soGGi package

quant\_params An optional BiocParallelParam instance determining the parallel back-end

to be used during evaluation. When this argument is set to NULL (default) SerialParam() will be used. For parallelization, MulticoreParam() can be used.

#### Value

A profileplyr object

8 clusterRanges

### **Description**

Cluster the ranges in a deepTools object based on signal within each range

#### Usage

```
clusterRanges(
  object = "profileplyr",
  fun = "function",
  scaleRows = "logical",
  kmeans_k = "integer",
  clustering_callback = "function",
  clustering_distance_rows = "ANY",
  cluster_method = "function",
  cutree_rows = "integer",
  silent = "logical",
  show_rownames = "logical",
  cluster_sample_subset = "ANY"
)
## S4 method for signature 'profileplyr'
clusterRanges(
  object = "profileplyr",
  fun = rowMeans,
  scaleRows = TRUE,
  kmeans_k = NULL,
  clustering_callback = function(x, ...) {
                                                return(x) },
  clustering_distance_rows = "euclidean",
  cluster_method = "complete",
  cutree_rows = NULL,
  silent = TRUE,
  show_rownames = FALSE,
  cluster_sample_subset = NULL
)
```

#### **Arguments**

object A profileplyr object

fun The function used to summarize the ranges (e.g. rowMeans or rowMax). This

is ignored when only one sample is used for clustering; in this case the lone

heatmap is clustered based the signal across the bins.

scaleRows If TRUE, the rows of the matrix containing the signal in each bin that is used as

the input for clustering will be scaled (as specified by pheatmap)

kmeans\_k The number of kmeans groups used for clustering

clustering\_callback

Clustering callback function to be passed to pheatmap

clustering\_distance\_rows

distance measure used in clustering rows. Possible values are "correlation" for Pearson correlation and all the distances supported by dist, such as "euclidean",

clusterRanges 9

etc. If the value is none of the above it is assumed that a distance matrix is provided.

cluster\_method clustering method used. Accepts the same values as hclust

cutree\_rows The number of clusters for hierarchical clustering

silent Whether or not a heatmap (from pheatmap) is shown with the output. This will

not change what is returned with the function as it will always be a profileplyr object. If silent = FALSE, the heatmap will be shown which may be helpful in quick evaluation of varying numbers of clusters before proceeding with downstream analysis. The default is silent = TRUE, meaning no heatmap will be

shown

show\_rownames for any heatmaps printed while running this function, set to TRUE if rownames

should be displayed. Default is FALSE.

cluster\_sample\_subset

Either a character or numeric vector indicating the subset of heatmaps to be used for clustering. If a character vector, all elements of the vector must match names of the samples of the profileplyr object (found with 'rownames(sampleData(object))'). For an numeric vector, the profileplyr object will be subset based on the samples that correspond to these numbers (i.e. the numeric index of that sample within 'rownames(sampleData(object))'). When only sample is chosen, the lone heatmap selected will be clustered by signal across the bins of that sample. When more than one sample are selected, the 'fun' argument will be used to summarize the ranges and cluster across these selected samples.

### **Details**

tbd

#### Value

A profileplyr object

#### Methods (by class)

• profileplyr: Cluster Ranges

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)

# k-means clustering
clusterRanges(object, fun = rowMeans, kmeans_k = 3)

# hierarchical clustering, print heatmap, yet still return profileplyr object
clusterRanges(object, fun = rowMeans, cutree_rows = 3, silent = FALSE)</pre>
```

convertToEnrichedHeatmapMat

export a profileplyr object to a list of matrices that can be used as an input for EnrichedHeatmap

### **Description**

export a profileplyr object to a list of matrices that can be used as an input for EnrichedHeatmap

### Usage

```
convertToEnrichedHeatmapMat(object = "profileplyr", sample_names = "character")
## S4 method for signature 'profileplyr'
convertToEnrichedHeatmapMat(object = "profileplyr", sample_names = NULL)
```

#### **Arguments**

object A profileplyr object

sample\_names A character vector that will set the names of the heatmap components that are

generated from the profileplyr assays() matrices. This argument is optional, by default the names will be the name of the samples in the profileplyr object

rownames(sampleData(object)).

#### **Details**

Takes a profileplyr object and converts all of the matrices in the assays() section of the object to matrices that can be used as an input for EnrichedHeatmap

### Value

A list of normalized matrices that can be used for generating visualizations with EnrichedHeatmap

# Methods (by class)

• profileplyr: export a profileplyr object to a list of matrices that can be used as an input for EnrichedHeatmap

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
library(EnrichedHeatmap)
EH_mat <- convertToEnrichedHeatmapMat(object)
EnrichedHeatmap(EH_mat[[1]], name = names(EH_mat[1]), column_title = names(EH_mat[1]))</pre>
```

export\_deepToolsMat 11

 $export\_deepToolsMat$ 

Export and import profileplyr from/to deeptools

#### **Description**

Export and Import files

### Usage

```
export_deepToolsMat(
  object = "profileplyr",
  con = "character",
  decreasing = "logical",
  overwrite = "logical"
)

## S4 method for signature 'profileplyr'
export_deepToolsMat(
  object = "profileplyr",
  con = "character",
  decreasing = FALSE,
  overwrite = FALSE
)

import_deepToolsMat(con)
```

# **Arguments**

object A profileplyr object

con Connection to write/read deeptools data to/from.

decreasing If object@params\$mcolToOrderBy has been set and not NULL, then the ranges

will be ordered by the column indicated in this slot of the metadata. By default, the order will be increasing for the factor or numeric value. For decreasing order,

choose decreasing = TRUE.

overwrite Logical specifying whether to overwrite output if it exists.

#### **Details**

A profileplyr object

### Value

The path to deepTools matrix file

A profileplyr object

#### Methods (by class)

• profileplyr: Export and import profileplyr from/to deeptools

### **Examples**

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
export_deepToolsMat(object,file.path(tempdir(),"ATAC_Example.MAT"))</pre>
```

generateEnrichedHeatmap

generateEnrichedHeatmap

# **Description**

export a profileplyr object directly to an object of the EnrichedHeatmap class

#### Usage

```
generateEnrichedHeatmap(
 object,
  include_group_annotation = TRUE,
  extra_annotation_columns = NULL,
  sample_names = NULL,
  return_ht_list = FALSE,
  ylim = "common_max",
  top_anno_height = unit(2, "cm"),
  samples_to_sortby = NULL,
  decreasing = FALSE,
  all_color_scales_equal = TRUE,
 matrices_color = NULL,
  color_by_sample_group = NULL,
 matrices_pos_line = FALSE,
 matrices_pos_line_gp = gpar(lty = 2),
  top_anno_axis_font = gpar(fontsize = 8),
 matrices_column_title_gp = gpar(fontsize = 8, fontface = "bold"),
 matrices_axis_name = NULL,
 matrices_axis_name_gp = gpar(fontsize = 8),
  group_anno_color = NULL,
  group_anno_width = 3,
  group_anno_row_title_gp = gpar(fontsize = 10),
  group_anno_column_names_gp = gpar(fontsize = 10),
 extra_anno_color = vector(mode = "list", length = length(extra_annotation_columns)),
  extra_anno_top_annotation = TRUE,
  extra_anno_width = (rep(6, length(extra_annotation_columns))),
  only_extra_annotation_columns = FALSE,
  gap = 2,
```

```
genes_to_label = NULL,
  gene_label_font_size = 6,
  show_heatmap_legend = NULL,
  legend_params = list(),
  use_raster = length(object) > 2000,
  raster_device = "CairoPNG",
  raster_quality = 2,
  raster_device_param = list()
)
```

#### **Arguments**

object A profileplyr object

include\_group\_annotation

If TRUE (default value) then the Heatmap will be grouped based on the range metadata column specified by 'rowGroupsInUse'

extra\_annotation\_columns

A character vector of names that match column names of mcols(object). Extra annotation columns will be added to the heatmap based on the values of these indicated range metadata columns.

sample\_names

A character vector that will set the names of the heatmap components that are generated from the profileplyr assays() matrices. This argument is optional, by default the names will be the name of the samples in the profileplyr object rownames(sampleData(object)).

return\_ht\_list Whether the returned object is the heatmap list and not the actual figure. This will be a list of the various components (heatmaps and annotation columns) that can be added to with additional columns in a customized manner.

ylim

A numeric vector of two numbers that specifies the minimum and maximum of the yaxis of all the heatmaps generated for the matrices. The default is to use the max of the heatmap with the highest signal (ylim = 'common\_max'). If ylim = NULL, different ranges will be inferred for each heatmap. If ylim is a single numeric vector, then that range will be used for all heatmaps. Different ranges can be set for each heatmap by making ylim a list that is the same length as the number of heatmaps/matrices, with each element of the list corresponding to each heatmap. Lastly, ylim can be a character string matching a column name in sampleData(object), and this will make the heatmaps with the same grouping have the same ylims as determined by the common max within groups.

top\_anno\_height

The height (as a unit object) of the top annotation of all heatmaps representing the matrices

samples\_to\_sortby

Only relevant if object@params\$mcolToOrderBy is NULL (i.e it hasn't been changed), meaning that the rows are sorted by the mean signal of all heatmaps. This argument allows sorting by the mean of a subset of samples, and should be either a character or numeric vector. If numeric, then the samples/matrices that have that index in the profileplyr object will be used to order the rows of the heatmap. If a character vector, then the elements must match the name of a sample in the object (rownames(sampleData(object))), and these samples will be used to order the heatmap.

decreasing

If object@params\$mcolToOrderBy has been changed and is not NULL, then the ranges will be ordered by the column indicated in this slot of the metadata. By default, the order will be increasing for the factor or numeric value. For decreasing order, choose decreasing = TRUE.

#### all\_color\_scales\_equal

If TRUE (default value) then the same color scale will be used for each separate heatmap. If FALSE, color scales will be inferred for each heatmap as indicated by the legends.

matrices\_color Either a single character vector, a numeric vector, a function call to colorRamp2 from the circlize package, or a list. For anything but a list, all the heatmaps generated for the matrices of the profileplyr object will be the same and will be colored as specified here. The character and numeric vector inputs must be either two or three elements in length (denoting color progressions - three elements will give a middle color break), and each element must be a character string or number that points to a color. By default, numeric vectors use the colors in palette(), however this can be expanded with other R color lists(e.g. colors()). If this argument is a list then it's length must equal the number of matrices/samples that exist in the input profileplyr object. The components of the list can be either a numeric vector, character vector, or color function (they do not have to all be the same type of specification). Each element in the list will be the color mapping to the corresponding element in the profileplyr object.

#### color\_by\_sample\_group

A character vector that is identical to a column name in sampleData(object), and if set, the heatmaps will be colored based on that column (should be a factor, if not it will be converted to one)

### matrices\_pos\_line

A logical for whether to draw a vertical line(s) at the position of the target (for both a single point or a window). Default is true.

#### matrices\_pos\_line\_gp

Graphics parameters for the vertical position lines. Should be set with the gpar() function from the grid() package.

#### top\_anno\_axis\_font

The fontsize of the y-axis labels for the top annotation of all heatmaps representing the matrices

### matrices\_column\_title\_gp

Graphics parameters for the titles on top of each range/matrix. Should be set with the gpar() function from the grid() package.

#### matrices\_axis\_name

Names for axis which is below the heatmap. For profileplyr object made from BamBigwig\_to\_chipProfile/as\_profileplyr functions, the names will be of length three, with the middle point being the midpoint of each range. If the profileplyr object was made from a deeptools matrix with import\_deepToolsMat(), the names will be length three if matrix was generated with 'computeMatrix reference-point', or length of four if matrix was generated with 'computeMatrix scale-regions' corresponding to upstream, start of targets, end of targets and downstream (or length of two if no upstream/downstream included).

#### matrices\_axis\_name\_gp

Graphics parameters for the text on the x-axis of each matrix heatmap. Should be set with the gpar() function from the grid() package.

#### group\_anno\_color

This will specify colors for the grouping column if the 'include\_group\_annotation' argument is set to TRUE. Since the group column of the range metadata should always be a discrete value, this should be either a numeric vector or character vector with color names. By default, numeric vectors use the colors in palette(), however this can be expanded with other R color lists(e.g. colors()). The length of this vector must equal the number of groups.

#### group\_anno\_width

A numeric value that is used to will set the width of the column bar (in mm using the unit() function from the grid package) for the grouping annotation column.

#### group\_anno\_row\_title\_gp

Graphics parameters for the labels of the groups on the side of the heatmap. Should be set with the gpar() function from the grid() package.

#### group\_anno\_column\_names\_gp

Graphics parameters for the label of the grouping annotation column. Should be set with the gpar() function from the grid() package.

#### extra\_anno\_color

This will specify colors for the annotation columns added by the 'extra\_annotation\_columns' argument. This must be a list that is of equal length to the 'extra\_annotation\_columns' argument. Each element of this list will be used to specify the color scheme for the corresponding element of the 'extra\_annotation\_columns' vector. If an element is NULL, the default colors will be used for the column annotation. For a column with discrete variables this will typically be a vector of numbers or a vector of color names. By default, numeric vectors use the colors in palette(), however this can be expanded with other R color lists(e.g. colors()). For columns with continuous variables, this can also be a a vector of numbers or a vector of color names to signify the color progression, or it can be color mapping function using colorRamp2() from the circlize package.

#### extra\_anno\_top\_annotation

This is a logical vector that determines whether annotation plots are shown on top of the heatmaps for the extra annotations. This must either be a length of 1, in which case all of the heatmaps will abide by this value. Otherwise this must be a vector of equal length to the 'extra\_annotation\_columns' argument and the elements of this vector will correspond to the equivalent elements in 'extra annotation\_columns'

#### extra\_anno\_width

This will set the width of the individual extra annotation columns on the right side of the figure. This must be a numeric vector with each element setting the width for the corresponding element in the 'extra\_annotation\_columns' argument.

#### only\_extra\_annotation\_columns

If set to TRUE, only the heatmaps representing the extra annotation columns

gap

will be shown, and the range based heatmaps from the assay matrices will be excluded.

CACIUG

The size of the gap between heatmaps and annotations. Only relevant if re-

turn\_ht\_list = FALSE

genes\_to\_label A character vector of gene symbols that should match character strings in the

'SYMBOL' column that results from either 'annotateRanges' or 'annotateRanges\_great'.

Genes that are both in this vector and in the 'SYMBOL' column will be labeled

on the heatmap.

gene\_label\_font\_size

The size of the text for the labels for genes specified in 'genes\_to\_label' argu-

ment.

show\_heatmap\_legend

A logical vector with each position corresponding to each matrix heatmap (not including the 'extra\_annotation\_columns') that determines whether a legend is produced for that heatmap. By default a single legend is made if all heatmaps use the same color scale, or separate legends are made for each matrix heatmap

if the scales are different.

legend\_params A list that contains parameters for the legend. See color\_mapping\_legend-ColorMapping-method

for all available parameters.

use\_raster Whether render the heatmap body as a raster image. It helps to reduce file size

when the matrix is huge.

raster\_device Graphic device which is used to generate the raster image. Options are "png",

"jpeg", "tiff", "CairoPNG", "CairoJPEG", "CairoTIFF"

raster\_quality A value set to larger than 1 will improve the quality of the raster image.

raster\_device\_param

A list of further parameters for the selected graphic device. For raster image support, please check https://jokergoo.github.io/ComplexHeatmap-reference/book/a-single-heatmap.html#heatmap-as-raster-image.

#### **Details**

Takes a profileplyr object and generates a heatmap that can be annotated by group or by range metadata columns of the profileplyr object

# Value

By default a customized version of a heatmap from EnrichedHeatmap, if return\_ht\_list = TRUE then a heatmap list is returned that can be modified and then entered as an input for the EnrichedHeatmap function

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
generateEnrichedHeatmap(object, include_group_annotation = FALSE)</pre>
```

generateProfilePlot 17

 ${\tt generateProfilePlot} \qquad {\tt \textit{Import ChIP profile object to profileplyr}}$ 

# Description

Function to convert soGGi ChIPprofile objects to profileplyr object .

# Usage

```
generateProfilePlot(
  object,
  sampleNames = rownames(sampleData(object)),
  colorGroup = params(object)$rowGroupsInUse,
  colorlist = NULL,
  facet_nrow = 1,
  facet_ncol = NULL,
  facet_scales = "fixed"
)
```

# Arguments

object	A profileplyr object
sampleNames	The names used to label the samples in the profileplyr object. By default, the names stored in rownames(sampleData(object)) are used.
colorGroup	The name of the column in mcols(object) that will be used for color grouping in the plot. By default the column name in params(object)\$rowGroupsInUse is used. If this column is not a factor variable, then it will be converted into one.
colorlist	A vector containing the colors to be used. The positions in the vector will be matched with the levels of the factor variable chosen in the 'colorGroup' argument.
facet_nrow	The number of rows when making the plot panels. This argument is passed to 'nrow' of the ggplot2 function facet_wrap.
facet_ncol	The number of columns when making the plot panels. This argument is passed to 'ncol' of the ggplot2 function facet_wrap.
facet_scales	Whether the scales of all plot panels should be fixed ("fixed", default), free ("free"), or free in one dimension ("free_x" or "free_y"). This argument is passed to 'scales' of the ggplot2 function facet_wrap.

# Value

A profileplyr object

18 gene\_list\_dataframe

### **Examples**

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
generateProfilePlot(object)</pre>
```

gene\_list\_character

Character vector of the top differentially expressed genes from hindbrain versus liver as measured by RNA-seq

### **Description**

This dataset contains a character vector of the top differentially expressed genes in the hindbrain versus liver as measured by RNA-seq (both genes that go up and those that go down). Data was downloaded from ENCODE.

### Usage

```
data(gene_list_character)
```

#### **Details**

• gene\_list\_character

#### Value

A character vector of the top differentially expressed genes in the hindbrain versus liver as measured by RNA-seq/

gene\_list\_dataframe

Dataframe of top differentially expressed genes from hindbrain versus liver as measured by RNA-seq

#### **Description**

This dataset contains a dataframe of the top differentially expressed genes in the hindbrain versus liver as measured by RNA-seq (both genes that go up and those that go down). The gene names are the rownames, and the first column is the 'stat' column from DESeq2. Data was downloaded from ENCODE.

# Usage

```
data(gene_list_dataframe)
```

groupBy 19

#### **Details**

• gene\_list\_dataframe

#### Value

A dataframe of top differentially expressed genes from hindbrain versus liver as measured by RNA-seq/

groupBy

group the rows and ranges of the profileplyr object

# **Description**

group the rows and ranges of the profileplyr object

# Usage

```
groupBy(
 object = "profileplyr",
  group = "ANY",
 GRanges_names = "character",
  levels = "ANY",
  include_nonoverlapping = "logical",
  separateDuplicated = "logical",
  inherit_groups = "logical"
)
## S4 method for signature 'profileplyr'
groupBy(
 object = "profileplyr",
  group = "ANY",
  GRanges_names = NULL,
  levels = NULL,
  include_nonoverlapping = FALSE,
  separateDuplicated = TRUE,
  inherit_groups = FALSE
)
```

# **Arguments**

object

A profileplyr object

group

How the ranges will be grouped. If this is a character string, then it must match a column name of the range metadata, and this column will be used for grouping of any exported deepTools matrix. If this is a GRanges, or GRangesList, then the ranges will be subset based on overlap with these GRanges. If this is a list, each element should contain ether 1) a character vector of genes, and ranges will

20 groupBy

> be subset based on overlap with these genes, as determined by the annotations made by annotateRanges() or annotateRanges\_great() functions, or 2) a data frame with the gene symbols as the rownames. Any additional columns of this dataframe will be added to the range metadata.

GRanges\_names

The names of the GRanges that were used for the "GRanges" argument. This will be used to label these groups in the construction of the resulting profileplyr object.

levels

This will set the levels of the grouping column set by 'rowGroupsInUse' (if the grouping column is not a factor, it will be converted to one). If levels are not provided, they will remain unchanged if the grouping column was already a factor, or will use default leveling (e.g. alphabetical) if grouping column is not already a factor variable.

include\_nonoverlapping

A logical argument, if FALSE (default) the regions from the original deepTools matrix that do not overlap with the user defined regions will be left out of the returned profileplyr object.

separateDuplicated

A logical argument, if TRUE (default) then regions that overlap multiple inputs to 'GRanges' argument will be separated and made into their own group. All possible combinations of region overlaps will be tested, so it is not recommended to have more than 3 groups if this option is TRUE. If FALSE, then regions that overlap each individual 'GRanges' input will be in the output, and if one region overlaps multiple 'GRanges' inputs, then it will be duplicated in the output and will show up in the section for each group.

inherit\_groups A logical whether that groups the exist in the profileplyr object in the 'object' argument should be included in the default grouping scheme for the output object of this function. The default is TRUE. If false, only the GRanges or gene list overlap annotation will be used for heatmap grouping.

### **Details**

Takes a SE object and groups rows

#### Value

A profileplyr object

### Methods (by class)

• profileplyr: group the rows and ranges of the profileplyr object

```
# group by gene list or list of data frames with genes as rownames
## not shown here but see vignette for grouping by gene lists
# group by GRanges
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
```

inherit\_group\_function 21

```
object <- import_deepToolsMat(example)
data("K27ac_GRlist_hind_liver_top5000") # load pre-made GRanges
K27ac_groupByGR <- groupBy(object, group = K27ac_GRlist_hind_liver_top5000)
# switch rowGroupsInUse
switchGroup <- groupBy(K27ac_groupByGR, group = "GR_overlap_names")
params(switchGroup)$rowGroupsInUse</pre>
```

inherit\_group\_function

Redundant code for inheriting grouping wrapped into subsetbyRangeOverlap() or subsetbyGeneListOverlap() functions

#### **Description**

Redundant code for inheriting grouping wrapped into subsetbyRangeOverlap() or subsetbyGeneListOverlap() functions

#### **Usage**

```
inherit_group_function(object, rowGroupsInUse_input, type, separateDuplicated)
```

#### **Arguments**

object A profileplyr object

rowGroupsInUse\_input

the inherited rowGroupsInUse

type

Either "GR" for subsetbyRangeOverlap() function or "GL" for subsetbyGeneLis-

tOverlap() function

separateDuplicated

A logical argument, if TRUE then regions that overlap multiple inputs to 'GRanges' argument will be separated and made into their own group. All possible combinations of region overlaps will be tested, so it is not recommended to have more than 3 groups if this option is TRUE. If FALSE, then regions that overlap each individual 'GRanges' input will be in the output, and if one region overlaps multiple 'GRanges' inputs, then it will be duplicated in the output and will show up in the section for each group.

### **Details**

tbd

#### Value

A profileplyr object

22 orderBy

```
K27ac_GRlist_hind_liver_top5000
```

GRangesList of the top 5000 H3K27ac peaks from hindbrain and liver downloaded from ENCODE

### **Description**

This dataset contains a GRangesList of the H3K27ac peaks in either the hindbrain or the liver with the highest signal. Data was downloaded from ENCODE.

#### Usage

```
data(K27ac_GRlist_hind_liver_top5000)
```

### **Details**

• K27ac\_GRlist\_hind\_liver\_top5000

### Value

A GRangesList of the top 5000 H3K27ac peaks from hindbrain and liver downloaded from ENCODE/

orderBy

choose the column by which to order the ranges by within each group

# Description

choose the column by which to order the ranges by within each group

#### Usage

```
orderBy(object = "profileplyr", column = "ANY")
## S4 method for signature 'profileplyr'
orderBy(object = "profileplyr", column = "ANY")
```

#### **Arguments**

object A profileplyr object

column Which column of mcols(proplyrObject) should be used for ordering the ranges.

If NULL removes any previous setting for row ordering.

### **Details**

Takes a profileplyr object and orders the rows based on a user defined metadata column of rowRanges

params 23

# Value

A profileplyr object

### Methods (by class)

• profileplyr: choose the column by which to order the ranges by within each group

#### **Examples**

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)

library(SummarizedExperiment)
cluster <- clusterRanges(object, fun = rowMeans, cutree_rows = 3)
cluster_order <- orderBy(cluster, column = "hierarchical_order")
params(cluster_order)$mcolToOrderBy</pre>
```

params

Retrieve and set parameters in profileplyr object

#### **Description**

Retrieve and set parameters in profileplyr object

# Usage

```
params(object)
```

### **Arguments**

object

A profileplyr object

### Value

A list containing parameters for profileplyr object.

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
params(object)</pre>
```

24 sampleData

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profil	rebry	r-class

Join, subset and manipulate ChIPprofile objects

# Description

Join, subset and manipulate ChIPprofile objects

# Usage

```
## S4 method for signature 'profileplyr'
c(x, ...)
## S4 method for signature 'profileplyr,ANY,ANY,ANY'
x[i, j, k, ..., drop = FALSE]
```

# Arguments

X	profileplyr object
	Additional arguments.
i	An integer or character scalar indicating ranges of profileplyr object to return
j	An integer or character scalar indicating columns of profileplyr object to return or a An integer or character scalar indicating which profileplyr object samples to return
k	An integer or character scalar indicating samples of profileplyr object to return.
drop	A logical whether to drop empty samples

# Value

A profileplyr object

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sam	Lαι	.eu	ata	

Retrieve and set sample data in profileplyr object

# Description

Retrieve and set sample data in profileplyr object

#### Usage

```
sampleData(object = "profileplyr")

## S4 method for signature 'profileplyr'
sampleData(object = "profileplyr")

sampleData(object) <- value

## S4 replacement method for signature 'profileplyr,DataFrame'
sampleData(object) <- value</pre>
```

# Arguments

object A profileplyr object

value DataFrame of sample information

#### Value

A DataFrame containing sample data

A DataFrame containing sample data to replace current sample data

### **Examples**

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)
sampleData(object)
sampleData(object)$scale <- c(1,10,1)</pre>
```

subsetbyGeneListOverlap

Subset ranges based on overlap with lists of Gene sets

# **Description**

The ranges from the deepTools matrix will be subset based on whether they overlap with user defined gene sets

# Usage

```
subsetbyGeneListOverlap(
  object,
  group,
  include_nonoverlapping = FALSE,
  separateDuplicated = TRUE,
  inherit_groups = FALSE
)
```

#### **Arguments**

object

A profileplyr object

group

How the ranges will be grouped. If this is a character string, then it must match a column name of the range metadata, and this column will be used for grouping of any exported deepTools matrix. If this is a GRanges, or GRangesList, then the ranges will be subset based on overlap with these GRanges. If this is a list, each element should contain ether 1) a character vector of genes, and ranges will be subset based on overlap with these genes, as determined by the annotations made by annotateRanges() or annotateRanges\_great() functions, or 2) a data frame with the gene symbols as the rownames. Any additional columns of this dataframe will be added to the range metadata.

include\_nonoverlapping

A logical argument, if FALSE the regions from the original deepTools matrix that do not overlap with the user defined regions will be left out of the returned profileplyr object.

separateDuplicated

A logical argument, if TRUE (default) then regions that overlap multiple inputs to 'GRanges' argument will be separated and made into their own group. All possible combinations of region overlaps will be tested, so it is not recommended to have more than 3 groups if this option is TRUE. If FALSE, then regions that overlap each individual 'GRanges' input will be in the output, and if one region overlaps multiple 'GRanges' inputs, then it will be duplicated in the output and will show up in the section for each group.

inherit\_groups A logical whether that groups the exist in the profileplyr object in the 'object' argument should be included in the default grouping scheme for the output object of this function. The default is TRUE. If false, only the gene list overlap annotation will be used for heatmap grouping.

#### **Details**

tbd

#### Value

A profileplyr object

#### **Examples**

# see the groupby function within profileplyr for examples

subsetbyRangeOverlap Subset ranges based on overlap with a GRanges object

Description

The ranges from the deepTools matrix will be subset based on whether they overlap with user defined ranges

#### **Usage**

```
subsetbyRangeOverlap(
  object,
  group,
  GRanges_names = NULL,
  include_nonoverlapping = FALSE,
  separateDuplicated = TRUE,
  inherit_groups = FALSE
)
```

#### **Arguments**

object

A profileplyr object

group

How the ranges will be grouped. If this is a character string, then it must match a column name of the range metadata, and this column will be used for grouping of any exported deepTools matrix. If this is a GRanges, or GRangesList, then the ranges will be subset based on overlap with these GRanges. If this is a list, each element should contain ether 1) a character vector of genes, and ranges will be subset based on overlap with these genes, as determined by the annotations made by annotateRanges() or annotateRanges\_great() functions, or 2) a data frame with the gene symbols as the rownames. Any additional columns of this dataframe will be added to the range metadata.

GRanges\_names

The names of the GRanges that were used for the "GRanges" argument. This will be used to label these groups in the construction of the resulting profileplyr object.

include\_nonoverlapping

A logical argument, if FALSE the regions from the original deepTools matrix that do not overlap with the user defined regions will be left out of the returned profileplyr object.

separateDuplicated

A logical argument, if TRUE then regions that overlap multiple inputs to 'GRanges' argument will be separated and made into their own group. All possible combinations of region overlaps will be tested, so it is not recommended to have more than 3 groups if this option is TRUE. If FALSE, then regions that overlap each individual 'GRanges' input will be in the output, and if one region overlaps multiple 'GRanges' inputs, then it will be duplicated in the output and will show up in the section for each group.

inherit\_groups A logical whether that groups the exist in the profileplyr object in the 'object'

argument should be included in the default grouping scheme for the output object of this function. The default is TRUE. If false, only the GRanges overlap

annotation will be used for heatmap grouping.

#### **Details**

tbd

#### Value

A profileplyr object

#### **Examples**

# see the groupby function within profileplyr for examples

subset\_GR\_GL\_common\_top

Redundant code wrapped into subsetbyRangeOverlap() or subsetby-GeneListOverlap() functions

### Description

Redundant code wrapped into subsetbyRangeOverlap() or subsetbyGeneListOverlap() functions

#### Usage

subset\_GR\_GL\_common\_top(object, overlap, input\_names, type, separateDuplicated)

#### **Arguments**

object A profileplyr object

overlap hits object from subsetByOverlap function

input\_names names of either the gene list of the granges that go into function

type Either "GR" for subsetbyRangeOverlap() function or "GL" for subsetbyGeneLis-

tOverlap() function

separateDuplicated

A logical argument, if TRUE (default) then regions that overlap multiple inputs to 'GRanges' argument will be separated and made into their own group. All possible combinations of region overlaps will be tested, so it is not recommended to have more than 3 groups if this option is TRUE. If FALSE, then regions that overlap each individual 'GRanges' input will be in the output, and if one region overlaps multiple 'GRanges' inputs, then it will be duplicated in the output and will show up in the section for each group.

summarize 29

#### **Details**

tbd

#### Value

A list of profileplyr objects

summarize

summarize the rows of a deepTools matrix

# Description

summarize the rows of a deepTools matrix

# Usage

```
summarize(
  object = "profileplyr",
  fun = "function",
  output = "character",
  keep_all_mcols = "logical",
  sampleData_columns_for_longPlot = "character"
)

## S4 method for signature 'profileplyr'
summarize(
  object = "profileplyr",
  fun = "function",
  output = "character",
  keep_all_mcols = FALSE,
  sampleData_columns_for_longPlot = NULL
)
```

# **Arguments**

object A profileplyr object

fun the function used to summarize the ranges (e.g. rowMeans or rowMax)

output Must be either "matrix", "long", or "object".

output indicate the final in the first of th

keep\_all\_mcols if output is 'long' and this is set to TRUE, then all metadata columns in the rowRanges will be included in the output. If FALSE (default value), then only the column indicated in the 'rowGroupsInUse' slot of the metadata will be included in the output dataframe.

sampleData\_columns\_for\_longPlot

If output is set to 'long', then this argument can be used to add information stored in sampleData(object) to the summarized data frame. This needs to be a character vector with elements matching coumn names in sampleData(object).

30 summarize

#### **Details**

Takes a SE object and outputs a summarized experiment object with a matrix containing ranges as rows and each sample having one column with summary statistic

#### Value

If output="matrix" returns a matrix, if output="long" returns a data.frame in long format, if output="object" returns a SummarizedExperiment object with the summarized matrix.

# Methods (by class)

• profileplyr: summarize the rows of a deepTools matrix

```
example <- system.file("extdata", "example_deepTools_MAT", package = "profileplyr")
object <- import_deepToolsMat(example)

# output matrix (can be used to make a heatmap)
object_sumMat <- summarize(object, fun = rowMeans, output = "matrix")

# output long dataframe for ggplot
object_long <- summarize(object, fun = rowMeans, output = "long")
object_long[1:3, ]

library(ggplot2)
ggplot(object_long, aes(x = Sample, y = log(Signal))) + geom_boxplot()

# output profileplyr object containing summarized matrix
summarize(object, fun = rowMeans, output = "object")</pre>
```

# **Index**

```
* datasets
                                                                                                                       groupBy, 19
           gene_list_character, 18
                                                                                                                       groupBy, profileplyr-method (groupBy), 19
           gene_list_dataframe, 18
                                                                                                                       import_deepToolsMat
          K27ac_GRlist_hind_liver_top5000,
                                                                                                                                             (export_deepToolsMat), 11
                                                                                                                       inherit_group_function, 21
[,profileplyr,ANY,ANY,ANY-method
                     (profileplyr-class), 24
                                                                                                                       K27ac_GRlist_hind_liver_top5000, 22
annotatePeak, 4
                                                                                                                       orderBy, 22
annotateRanges, 2
                                                                                                                       orderBy, profileplyr-method (orderBy), 22
annotateRanges, profileplyr-method
                     (annotateRanges), 2
                                                                                                                       params, 23
annotateRanges_great, 4
                                                                                                                       profileplyr-class, 24
annotate Ranges\_great, \verb|profileplyr-method|
                     (annotateRanges_great), 4
                                                                                                                       sampleData, 24
as_profileplyr, 5
                                                                                                                       sampleData,profileplyr-method
                                                                                                                                             (sampleData), 24
BamBigwig_to_chipProfile, 6
                                                                                                                        sampleData<- (sampleData), 24</pre>
BiocParallelParam, 7
                                                                                                                        sampleData<-,profileplyr,DataFrame-method</pre>
                                                                                                                                            (sampleData), 24
c,profileplyr-method
                                                                                                                       submitGreatJob, 5
                     (profileplyr-class), 24
                                                                                                                       subset_GR_GL_common_top, 28
clusterRanges, 7
                                                                                                                       subsetbyGeneListOverlap, 25
clusterRanges,profileplyr-method
                                                                                                                       subsetbyRangeOverlap, 27
                     (clusterRanges), 7
                                                                                                                       summarize, 29
convertToEnrichedHeatmapMat, 10
convert To Enriched Heatmap Mat, profile plyr-method {\tt nummarize}, profile plyr-met
                                                                                                                                            (summarize), 29
                     (convertToEnrichedHeatmapMat),
                     10
EnrichedHeatmap, 16
export_deepToolsMat, 11
export_deepToolsMat,profileplyr-method
                     (export_deepToolsMat), 11
facet_wrap, 17
gene_list_character, 18
gene_list_dataframe, 18
generateEnrichedHeatmap, 12
generateProfilePlot, 17
```