Package ‘epistack’

May 29, 2024

Title Heatmaps of Stack Profiles from Epigenetic Signals

Version 1.10.0

Description The epistack package main objective is the visualizations of stacks of genomic tracks (such as, but not restricted to, ChIP-seq, ATAC-seq, DNA methylation or genomic conservation data) centered at genomic regions of interest. epistack needs three different inputs: 1) a genomic score objects, such as ChIP-seq coverage or DNA methylation values, provided as a `GRanges` (easily obtained from `bigwig` or `bam` files). 2) a list of feature of interest, such as peaks or transcription start sites, provided as a `GRanges` (easily obtained from `gtf` or `bed` files). 3) a score to sort the features, such as peak height or gene expression value.

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Encoding UTF-8

LazyData false

Imports GenomicRanges, SummarizedExperiment, BiocGenerics, S4Vectors, IRanges, graphics, plotrix, grDevices, stats, methods

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addBins

Description

Add an optional bin metadata column to gr, to serve as annotations for the epistack plots.

Usage

addBins(rse, nbins = 5L, bin = NULL)

Arguments

- `rse` a SummarizedExperiment or a GRanges object.
- `nbins` an integer number, the number of bins.
- `bin` a vector containing pre-determined bins, in the same order as gr.
addMetricAndArrangeGRanges

Details

nbins is taken into account only if bin is NULL. rse should be sorted first, usually with the addMetricAndArrangeGRanges() function. addBin(rse, bin = vec) is equivalent to rse$bin <- vec, while addBin(rse, nbins = 5) will create 5 bins of equal size based on rse order.

Value

the RangedSummarizedExperiment or GRanges object with a new bin metadata column

See Also

addMetricAndArrangeGRanges plotBinning

Examples

data("stackepi")
addBins(stackepi)

# 3 bins instead of 5
addBins(stackepi, nbins = 3)

# assign bins using a vector
addBins(stackepi, bin = rep(c("a", "b", "c"), length.out = length(stackepi)))

addMetricAndArrangeGRanges

Description

Perform an inner join between a GRanges object and a data.frame. Sort the resulting GRanges based on a metric column.

Usage

addMetricAndArrangeGRanges(
  gr,
  order,
  gr_key = "name",
  order_key = "name",
  order_value = "exp",
  shuffle_tie = TRUE
)
addMetricAndArrangeRSE

Arguments

- `gr`: a GRanges object.
- `order`: a data.frame with at least two columns: keys and values.
- `gr_key`: name of the gr metadata column containing unique names for each genomic region in gr. Usually gene names/id or peak id.
- `order_key`: name of the order column that will be used as key for the inner join.
- `order_value`: name of the order column that contain value used for sorting.
- `shuffle_tie`: a boolean Value (TRUE / FALSE). When TRUE, shuffle the GRanges before sorting, mixing the ties.

Details

This utility function allow the addition of a metric column to genomic regions of interest. One of its common use case is to add gene expression values on a set of transcription start sites. The resulting GRanges object will only contain regions presents in both gr and order.

Value

a GRanges sorted in descending order.

Examples

data("stackepi_gr")
randomOrder <- data.frame(gene_id = stackepi_gr$gene_id,
                          value = rnorm(length(stackepi_gr)))
addMetricAndArrangeGRanges(stackepi_gr,
                          randomOrder, gr_key = "gene_id",
                          order_key = "gene_id", order_value = "value")

addMetricAndArrangeRSE

Description

Perform an inner join between a rangedSummarizedExperiment object and a data.frame. Sort the resulting rangedSummarizedExperiment based on a metric column.
addMetricAndArrangeRSE

Usage

addMetricAndArrangeRSE(
  rse,
  order,
  rse_key = "name",
  order_key = "name",
  order_value = "exp",
  shuffle_tie = TRUE
)

Arguments

- rse: a rangedSummarizedExperiment object.
- order: a data.frame with at least two columns: keys and values.
- rse_key: name of the gr metadata column containing unique names for each genomic region in rowRanges(rse). Usually gene names/id or peak id.
- order_key: name of the order column that will be used as key for the inner join.
- order_value: name of the order column that contain value used for sorting.
- shuffle_tie: a boolean Value (TRUE / FALSE). When TRUE, shuffle the GRanges before sorting, mixing the ties.

Details

This utility function allow the addition of a metric column to genomic regions of interest. One of its common use case is to add gene expression values on a set of transcription start sites. The resulting GRanges object will only contain regions presents in both rse and order.

Value

- a rangedSummarizedExperiment sorted in descending order.

Examples

data("stackepi")
randomOrder <- data.frame(   
  gene_id = SummarizedExperiment::rowRanges(stackepi)$gene_id, 
  value = rnorm(length(stackepi))
)
addMetricAndArrangeRSE(stackepi, 
  randomOrder, rse_key = "gene_id", 
  order_key = "gene_id", order_value = "value")
GRanges2RSE

**Description**

Convert objects from the old input format (GRanges object) to the new recommended input format RangedSummarizedExperiment.

**Usage**

GRanges2RSE(gr, patterns, names = patterns)

**Arguments**

- `gr` a GRanges object with matrix embeded as metadata columns.
- `patterns` A character vector of column prefixes (can be regular expressions) that should match columns of gr.
- `names` specify the desired names of the assays (if different from patterns).

**Details**

Mostly used for backward compatibilities and unit testing.

**Value**

a RangedSummarizedExperiment.

**Examples**

data("stackepi_gr")
GRanges2RSE(stackepi_gr, patterns = c("window"))
GRanges2RSE(stackepi_gr, patterns = c("^window_"), names = c("DNAme"))

---

meanColor

**Description**

Return the average color of a vector of colors, computed in the RGB space.

**Usage**

meanColor(colors)

**Arguments**

- `colors` a vector of colors
Details

Input colors can be either in html or color name formats. The alpha channel is supported but optional.

Value

a single color value

See Also

redimMatrix

Examples

meanColor(c("#000000FF", "#FFFFFF00", "#FFFF00FF", "#FF0000FF"))

# works with color names
meanColor(c("blue", "red"))

# Mix color names and HTML codes
meanColor(c("blue", "red", "#FFFF00FF"))

# works without alpha channel in inputs (but outputs an alpha channel):
meanColor(c("#000000", "#FFFFFF", "#FFFF00", "#FF0000"))

Description

Plot the average stack profiles +/- error (sd or sem). If a bin column is present in rowRanges(rse), one average profile is drawn for each bin.

Usage

plotAverageProfile(
  rse,
  assay = NULL,
  x_labels = c("Before", "Anchor", "After"),
  palette = colorRampPalette(c("#DF536B", "black", "#61D04F")),
  alpha_for_se = 0.25,
  error_type = c("sd", "sem", "ci95"),
  reversed_z_order = FALSE,
  ylim = NULL,
  y_title = NULL,
  pattern = NULL
)
Arguments

- **rse**: a RangedSummarizedExperiment input. Alternatively: can be a GRanges object (for backward compatibility, pattern will be required).
- **assay**: specify the name of the assay to plot, that should match one of assayNames(rse).
- **x_labels**: x-axis labels.
- **palette**: A vector of colors, or a function that returns a palette of n colors.
- **alpha_for_se**: the transparency (alpha) value for the error band.
- **error_type**: can be either "sd" (standard deviation), "sem" (standard error of the mean), or "ci95" (95% confidence interval). Default: "sd".
- **reversed_z_order**: should the z-order of the curves be reversed (i.e. first or last bin on top?)
- **ylim**: a vector of two numbers corresponding to the y-limits of the plot.
- **y_title**: the y-axis title.
- **pattern**: only if rse is of class GRanges. A single character that should match metadata of rse (can be a regular expression).

Value

Display a plot.

Examples

data("stackepi")
plotAverageProfile(stackepi)

plotBinning()
Arguments

rse a RangedSummarizedExperiment input with a column bin in rowRanges(rse). Alternatively (for backward compatibility), a GRanges object or any object such as rse$bin exists.

target_height an integer, the approximate height (in pixels) of the final plot. Used to avoid overplotting artefacts.

palette A vector of colors, or a function that returns a palette of n colors.

Value

Display a plot.

Examples

data("stackepi")
rse <- stackepi
rse <- addBins(rse, nbins = 3)
plotBinning(rse)

gr2 <- data.frame(bin = rep(c(1,2,3,4), each = 5))
plotBinning(gr2, palette = colorRampPalette(c("blue4", "forestgreen", "coral3", "goldenrod")))
plotEpistack

Arguments

- **rse**: a RangedSummarizedExperiment input. Alternatively: can be a GRanges object (for backward compatibility).
- **metric**: name of the column in rse metadata containing scores.
- **title**: title of the plot.
- **trans_func**: A function to transform value of x before plotting. Useful to apply log10 transformation (i.e. with trans_func = function(x) log10(x+1)).
- **ylim**: limit of the y axis; format: ylim = c(min, max)
- **ylab**: y-axis title
- **palette**: A vector of colors, or a function that returns a palette of n colors.

Value

Display a plot.

Examples

data("stackepi")
plotBoxMetric(
  stackepi,
  trans_func = function(x) x,
  metric = "exp",
  title = "Metric"
)

plotEpistack

plotEpistack()

Description

Given a list of genomic regions, epigenetic signals surrounding these regions, and a score for each region, plot epigenetic stacks depending on the score. An optional bin column allow the grouping of several genomic regions to produce average profiles per bins.

Usage

plotEpistack(
  rse,
  assays = NULL,
  tints = "gray",
  titles = NULL,
  legends = "",
  main = NULL,
  x_labels = c("Before", "Anchor", "After"),
  zlim = c(0, 1),
  ylim = NULL,
  palette = ""
metric_col = "exp",
metric_title = "Metric",
metric_label = "metric",
metric_ylab = NULL,
metric_transfunc = function(x) x,
bin_palette = colorRampPalette(c("#DF536B", "black", "#61D04F")),
npix_height = 650,
n_core = 1,
high_mar = c(2.5, 0.6, 4, 0.6),
low_mar = c(2.5, 0.6, 0.3, 0.6),
error_type = c("ci95", "sd", "sem"),
reversed_z_order = FALSE,
rel_widths = c(score = 0.35, bin = 0.08, assays = 0.35),
rel_heights = c(1, 0.14, 0.3),
patterns = NULL,
...)
)

Arguments

tse a RangedSummarizedExperiment input. Alternatively: can be a GRanges object (for backward compatibility, patterns will be required).

assays specify the name(s) and order of assay(s) to plot. A vector of names that should match assayNames(rse).

tints a vector of colors to tint the heatmaps. Can also be a function returning n colors, or a list of such palette functions.
titles titles of each heatmap. Defaults to assays.
legends legend names for the epistacks.
main Main title for the figure.
x_labels a character vector of length 3 used as x-axis labels.
zlim the minimum and maximum z values the heatmap. Format: zlim = c(min, max). zlim can also be specified of as a list of pairs of limits, on for each assay.
ylim limits of the y axis for bottom plots. ylim can also be specified of as a list of pairs of limits, on for each assay. Format: ylim = c(min, max)

metric_col a character, name of a column in gr such as expression value, peak height, pvalue, fold change, etc.

metric_title title to be display on the leftmost plots.
metric_label label of the leftmost plots.
metric_ylab y axis label of the top left plot.
metric_transfunc a function to transform value of metric_col before plotting. Useful to apply log10 transformation (i.e. with trans_func = function(x) log10(x+1)).
bin_palette A vector of colors, or a function that returns a palette of n colors. Used to color average profiles per bin in the bottom plots.
plotEpistack

npix_height  The matrix height is reduced to this number of rows before plotting. Useful to limit overplotting artefacts. It should roughly be set to the pixel height in the final heatmaps.

n_core     number of core used to speedup the matrix resizing.

high_mar   a vector of numerical values corresponding to the margins of the top figures. c(bottom, left, top, right)

low_mar    a vector of numerical values corresponding to the margins of the bottom figures. c(bottom, left, top, right)

error_type error_type, can be either "sd" (standard deviation), "sem" (standard error of the mean), or "ci95" (95% confidence interval). Default: "ci95".

reversed_z_order For the bottom panels: should the z-order of the curves be reversed (i.e. first or last bin on top)?

rel_widths  A named vector of three elements of relative panel widths: score is the leftmost panel, bin is the optionnal binning panels, and assays are the panels of the stacked-matrices. Default to c(score = .35, bin = .08, assays = .35)

rel_heights A vector of three elements of relative panel heights. Default to c(1, .14, .3)

patterns only if rse is of class GRanges. A character vector of column prefixes (can be regular expressions) that should match columns of rse.

... Arguments to be passed to par such as cex

Details

This function produce a comprehensive figure including epigenetic heatmaps and average epigenetic profiles from a well formated RangedSummarizedExperiment object with expected rowData metadata columns. It scales resonably well up to hundreds of thousands of genomic regions.

The visualisation is centered on an anchor, a set of genomic coordinated that can be transcription start sites or peak center for example. Anchor coordinates are those of the GRanges used as a rowData in the input RangedSummarizedExperiment object (hereafter rse).

Anchors are plotted from top to bottom in the same order as in rse. One should sort rse before plotting if needed.

rse’s rowData should have a metric column that is used in the leftmost plots. The name of the metric column must be specified to metric_col. The metric can be transformed before plotting if needed using the metric_transfunc parameter.

The matrix or matrices used to display the heatmap(s) should be passed as assay(s) in rse. Such matrix can be obtained using EnrichedHeatmap::normalizeToMatrix() for example. The assay names are then specified through assays.

If an optionnal bin column is present in rse’s rowData, it will be used to group genomic regions to performed average profile per bins in the bottom plots.

Epistack are multipanel plots build using layout(). Margins for the panels can be specified using high_mar and low_mar parameters if needed, especially to avoid text overlaps. The default value should be appropriate in most situations. Individual component can be plotted using severa epistack functions such as plotStackProfile() or plotAverageProfile().
Plotting more than > 1000 regions can lead to overplotting issues as well as some plotting artefacts (such as horizontal white strips). Both issues can be resolved with fiddling with the npix_height parameter. npix_height should be smaller than the number of regions, and in the same order of magnitude of the final heatmap height in pixels. Last minutes call to the redimMatrix() function will happen before plotting using npix_height as target height. Parameter n_core is passed to redimMatrix() to speed up the down-scaling.

The input can also be a GRanges object for backward compatibility. See GRanges2RSE. patterns would then be required.

Value

Display a plot.

See Also

plotStackProfile, plotAverageProfile, redimMatrix, normalizeToMatrix, addMetricAndArrangeGRanges, addBins

Examples

data("stackepi")
plotEpistack(stackepi,  
metric_col = "exp",  
ylim = c(0, 1),  
metric_transfunc = function(x) log10(x+1))

plotMetric

Description

Plot a vertical line chart of the metric column, in the same order as the input.

Usage

plotMetric(  
  x,  
  trans_func = function(x) x,  
  title = "Metric",  
  ylim = NULL,  
  xlab = NULL,  
  ylab = NULL  
)
Arguments

- **x**: a numeric vector.
- **trans_func**: a function to transform x values before plotting. Useful to apply log10 transformation (i.e. with `trans_func = function(x) log10(x+1)`).
- **title**: Title of the plot.
- **ylim**: limit of the y axis; format: `ylim = c(min, max)`
- **xlab**: x-axis title
- **ylab**: y-axis title

Value

Display a plot.

See Also

- `plotEpistack`, `plotBoxMetric`

Examples

```r
data("stackepi")
plotMetric(SummarizedExperiment::rowRanges(stackepi)$exp)
```

---

**Description**

Display a heatmap of an epigenetic track centered at genomic anchors such as Transcription Start Sites or peak center.

**Usage**

```r
plotStackProfile(
    rse,
    assay = NULL,
    x_labels = c("Before", "Anchor", "After"),
    title = "",
    ylim = NULL,
    palette = function(n) grDevices::hcl.colors(n, rev = TRUE),
    target_height = 650,
    summary_func = function(x) mean(x, na.rm = TRUE),
    n_core = 1,
    pattern = NULL
)
```
Arguments

- **rse**
  - a RangedSummarizedExperiment input. Alternatively: can be a GRanges object (for backward compatibility, pattern will be required).
- **assay**
  - specify the name of the assay to plot, that should match one of assayNames(rse).
- **x_labels**
  - a character vectors of length 3 used to label the x-axis.
- **title**
  - The title of the heatmap
- **zlim**
  - The minimum and maximum z values to match color to values. Format: zlim = c(min, max)
- **palette**
  - a palette of color, (i.e. a function of parameter n that should return n colors).
- **target_height**
  - The matrix height is reduced to this number of rows before plotting. Useful to limit overplotting artefacts. It should roughly be set to the pixel height in the final heatmap.
- **summary_func**
  - function passed to redimMatrix(). Usualy mean, but can be set to median or max for sparse matrices.
- **n_core**
  - multicore option, passed to redimMatrix().
- **pattern**
  - only if rse is of class GRanges. A character vector of length 1 of a column prefixe (can be regular expressions) that should match columns of rse.

Details

The visualisation is centered on an anchor, a set of genomic coordinates that can be transcription start sites or peak center for example. Anchor coordinates are those of the RangedSummarizedExperiment object used as an input (hereafter rse).

Anchors are plotted from top to bottom in the same order as in rse. One should sort rse before plotting if needed.

The matrix used to display the heatmap should be passed as assay of rse. Such matrix can be obtained using EnrichedHeatmap::normalizeToMatrix() for example.

This function scale reasonably well up to hundred thousands of regions. Overplotting issues are solved by last-minute reduction of the matrix size using redimMatrix().

Value

Display a plot.

See Also

plotAverageProfile, plotEpistack, normalizeToMatrix, plotStackProfileLegend

Examples

```r
data("stackepi")
plotStackProfile(stackepi,
    target_height = 650,
    zlim = c(0, 1),
    palette = colorRampPalette(c("white", "dodgerblue", "black")),
    title = "DNA methylation")
```
plotStackProfileLegend

\textit{plotStackProfileLegend}\()\)

\section*{Description}
Utility function to plot the corresponding legend key of \texttt{plotStackProfile()}’s plots.

\section*{Usage}
\begin{verbatim}
plotStackProfileLegend(
  zlim,
  palette = colorRampPalette(c("white", "grey", "black")),
  title = NA
)
\end{verbatim}

\section*{Arguments}
\begin{itemize}
  \item \texttt{zlim} \hspace{1cm} the limits of the values to be displayed. Format: c(min, max)
  \item \texttt{palette} \hspace{1cm} a palette of color, (i.e. a function of parameter n that should return n colors).
  \item \texttt{title} \hspace{1cm} an optional title to be display below the color legend.
\end{itemize}

\section*{Value}
Display a plot.

\section*{See Also}
\texttt{plotStackProfile}

\section*{Examples}
\begin{verbatim}
plotStackProfileLegend(zlim = c(0, 2),
  palette = colorRampPalette(c("white", "grey", "black")))
\end{verbatim}

\section*{redimMatrix}
\texttt{redimMatrix()}\)

\section*{Description}
Reduce the input matrix size by applying a summary function on cells to be fused.
redimMatrix

Usage

redimMatrix(
  mat,
  target_height = 100,
  target_width = 100,
  summary_func = function(x) mean(x, na.rm = TRUE),
  output_type = 0,
  n_core = 1
)

Arguments

mat                the input matrix.
target_height      height of the output matrix (should be smaller than or equal to nrow(mat)).
target_width       width of the output matrix (should be smaller than or equal to ncol(mat)).
summary_func       how to summerize cells? A function such has mean, median, max, or meanColors.
output_type        Type of the output, to be passed to vapply's FUN.VALUE.
n_core             number of core to use for parallel processing.

Details

This function is used to reduce matrix right before plotting them in order to avoid overplotting issues as well as other plotting artefacts.

Value

A resized matrix of size target_width x target_height where the summary_func was apply to adjacent cells.

See Also

meanColor

Examples

data("stackepi")
mat <- SummarizedExperiment::assay(stackepi, "DNAme")
dim(mat)
smallMat <- redimMatrix(mat, target_height = 10, target_width = ncol(mat))
dim(smallMat)

# changing the summary function
mat <- matrix(sample(1:40,100,replace=TRUE),nrow=10,ncol=10)
dim(mat)
smallMat <- redimMatrix(mat, target_height = 5, target_width = ncol(mat),
                        summary_func = function(x) max(x, na.rm = TRUE))
dim(smallMat)

# working with colors
```r
colmat <- matrix(
    c("red", "red", "blue", "blue", "red", "blue", "blue", "green"),
    ncol = 2
)
redimMatrix(colmat, target_height = 2, target_width = 2,
            summary_func = meanColor, output_type = "color")
```

---

**Description**

DNA methylation profiles (from MBD-seq data) around transcription start sites of the 693 chr18 genes annotated on the pig genome (Sscrofa11.1), as well as gene expression levels in Transcript Per Million (TPM) measured by RNA-seq in the same duodenum sample.

**Usage**

```r
data("stackepi")
```

**Format**

A RangedSummarizedExperiment of the 693 rows, 2 rows metadata columns, and one assay containing the DNA methylation signal.

**Source**

This dataset was generated from ENSEMBL annotation data and data generated by our lab (publicly available soon).

---

**Description**

DNA methylation profiles (from MBD-seq data) around transcription start sites of the 693 chr18 genes annotated on the pig genome (Sscrofa11.1), as well as gene expression levels in Transcript Per Million (TPM) measured by RNA-seq in the same duodenum sample.

**Usage**

```r
data("stackepi_gr")
```

**Format**

A GRanges of the 693 rows and 54 metadata columns, kept for unit-testing backward-compatibility.
Source
This dataset was generated from ENSEMBL annotation data and data generated by our lab (publicly available soon).

See Also
GRanges2RSE
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