Package ‘sechm’

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Type Package

Title sechm: Complex Heatmaps from a SummarizedExperiment

Version 1.12.0

Description sechm provides a simple interface between SummarizedExperiment objects and the ComplexHeatmap package.
It enables plotting annotated heatmaps from SE objects, with easy access to rowData and colData columns,
and implements a number of features to make the generation of heatmaps easier and more flexible.
These functionalities used to be part of the SEtools package.

Depends R (>= 4.0), SummarizedExperiment, ComplexHeatmap

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Description

Plot a multi-panel heatmap from a list of SummarizedExperiment-class.

Usage

crossHm(
  ses,
  features,
  do.scale = TRUE,
  uniqueScale = FALSE,
  assayName = .getDef("assayName"),
  sortBy = seq_along(ses),
  only.common = TRUE,
  cluster_cols = FALSE,
  cluster_rows = is.null(sortBy),
  toporder = NULL,
  hmcols = NULL,
  breaks = .getDef("breaks"),
  gaps_at = .getDef("gaps_at"),
  gaps_row = NULL,
  name = NULL,
  top_annotation = .getDef("anno_columns"),
  left_annotation = .getDef("anno_rows"),
  anno_colors = list(),
)
show_rownames = NULL,
merge_legends = FALSE,
show_colnames = FALSE,
rel.width = NULL,
...
}

Arguments

ses
A (named) list of `SummarizedExperiment-class` objects, with some matching row.names between them.

features
A vector of features (i.e. row.names) to plot.

do.scale
Logical; whether to scale rows in each SE (default TRUE).

uniqueScale
Logical; whether to force the same colorscale for each heatmap.

assayName
The name of the assay to use; if multiple names are given, the first available will be used. Defaults to "logcpm", "lognorm".

sortBy
Names or indexes of `ses` to use for sorting rows (default all)

only.common
Logical; whether to plot only rows common to all SEs (default TRUE).

cluster_cols
Logical; whether to cluster columns (default FALSE).

cluster_rows
Logical; whether to cluster rows (default TRUE if `do.sortRows=FALSE`, FALSE otherwise).

toporder
Optional vector of categories on which to supra-order when sorting rows, or name of a 'rowData' column to use for this purpose.

hmcols
Colors for the heatmap.

breaks
Breaks for the heatmap colors. Alternatively, symmetrical breaks can be generated automatically by setting ‘breaks’ to a numerical value between 0 and 1. The value is passed as the ‘split.prop’ argument to the `getBreaks` function, and indicates the proportion of the points to map to a linear scale, while the more extreme values will be plotted on a quantile scale. ‘breaks=FALSE’ will disable symmetrical scale and quantile capping, while retaining automatic breaks. ‘breaks=1’ will produce a symmetrical scale without quantile capping.

gaps_at
Columns of `colData` to use to establish gaps between columns.

gaps_row
A named vector according to which rows will be split.

name
The title of the heatmap key.

top_annotation
Columns of `colData` to use for top annotation.

left_annotation
Columns of `rowData` to use for left annotation.

anno_colors
List of colors to use for annotation.

show_rownames
Whether to show row names (default TRUE if 50 rows or less).

merge_legends
Logical; passed to `draw-HeatmapList-method`

show_colnames
Whether to show column names (default FALSE).

rel.width
Relative width of the heatmaps

... Any other parameter passed to each call of `Heatmap`.
Value

A Heatmap list.

Examples

data("Chen2017", package="sechm")
se1 <- Chen2017[,1:6]
se2 <- Chen2017[,7:15]
se3 <- crossHm(list(se1=se1, se2=se2), row.names(se1)[1:10] )

Description

A `SummarizedExperiment-class` containing (a subset of) hippocampus RNAseq of mice treated with Forskolin.

Value

A `SummarizedExperiment-class`.

References


getBreaks

Description

Produces symmetrical breaks for a color scale, with the scale steps increasing for large values, which is useful to avoid outliers influencing too much the color scale.

Usage

`getBreaks(x, n, split.prop = 0.98, symmetric = TRUE)`

Arguments

- `x` A matrix of log2FC (or any numerical values centered around 0)
- `n` The desired number of breaks.
- `split.prop` The proportion of the data points to plot on a linear scale; the remaining will be plotted on a scale with regular frequency per step (quantile).
- `symmetric` Logical; whether breaks should be symmetric around 0 (default TRUE)
getDEA

Value

A vector of breaks of length = 'n'

Examples

dat <- rnorm(100, sd = 10)
getBreaks(dat, 10)

getDEA(se, dea = NULL, homogenize = FALSE)

Arguments

se A SummarizedExperiment-class, with DEAs each saved as a rowData column of 'se', with the column name prefixed with "DEA."
dea The optional name of the DEA to extract
homogenize Logical; whether to homogenize the DEA

Value

The DEA data.frame if 'dea' is given, otherwise a named list of data.frames.

Examples

# loading example SE
data("Chen2017", package="sechm")
# this ones doesn't have saved DEAs in the standard format:
getDEA(Chen2017)
getDEGs

*Get DEGs from a SE or list of DEA results*

**Description**

Get DEGs from a SE or list of DEA results

**Usage**

```r
getDEGs(
  x,
  dea = NULL,
  lfc.th = log2(1.3),
  fdr.th = 0.05,
  direction = 0,
  merge = TRUE
)
```

**Arguments**

- `x`: A ‘SummarizedExperiment’ object with DEA results in rowData, or a list of DEA result data.frames.
- `dea`: Which DEA(s) to use (default all). Used only if ‘x’ is a ‘SummarizedExperiment’.
- `lfc.th`: Absolute log-foldchange threshold.
- `fdr.th`: FDR threshold.
- `direction`: If !=0, specifies whether to fetch only upregulated or downregulated features
- `merge`: Logical; whether to take the union of DEGs from the different DEAs (when more than one).

**Value**

A character vector with the significant features, or a list of such vectors.

**Examples**

```r
# loading example SE
data("Chen2017", package="sechm")
# this ones doesn't have saved DEAs in the standard format:
getDEGs(Chen2017)
```
**homogenizeDEA**

---

**Description**

Standardizes the outputs of differential expression methods (to an edgeR-like style)

**Usage**

```r
homogenizeDEA(x)
```

**Arguments**

- `x`: A data.frame containing the results of a differential expression analysis

**Value**

A standardized data.frame.

---

**log2FC**

---

**Description**

Generates log2(foldchange) matrix/assay, eventually on a per-batch fashion.

**Usage**

```r
log2FC(
  x,
  fromAssay = NULL,
  controls,
  by = NULL,
  isLog = NULL,
  agFun = rowMeans,
  toAssay = "log2FC",
  pseudocount = 1L,
  ndigits = 2
)
```
Arguments

- **x**: A numeric matrix, or a ‘SummarizedExperiment’ object
- **fromAssay**: The assay to use if ‘x’ is a ‘SummarizedExperiment’
- **controls**: A vector of which samples should be used as controls for foldchange calculations.
- **by**: An optional vector indicating groups/batches by which the controls will be averaged to calculate per-group foldchanges.
- **isLog**: Logical; whether the data is log-transformed. If NULL, will attempt to figure it out from the data and/or assay name.
- **agFun**: Aggregation function for the baseline (default rowMeans)
- **toAssay**: The name of the assay in which to save the output. If left to the default value, both a log2FC assay as well as a scaled log2FC assay (scaled by unit-variance, but not centered) will be saved in the object.
- **pseudocount**: If the origin assay is not log-transformed, ‘pseudocount’ will be added to the values before calculating a log-transformation. This prevents infinite fold-changes and moderates them.
- **ndigits**: Number of digits after the decimal of the log2FC (and scaledLFC).

Value

An object of same class as ‘x’; if a ‘SummarizedExperiment’, will have the additional assay named from ‘toAssay’.

Examples

```R
log2FC( matrix(rnorm(40), ncol=4), controls=1:2 )
```

meltSE

Description

Melts a SE object into a ggplot-ready long data.frame.

Usage

```R
meltSE(
  x,
  features,
  assayName = NULL,
  colDat.columns = NULL,
  rowDat.columns = NULL,
  flatten = TRUE,
  baseDF = TRUE
)
```
`qualitativeColors`

**Arguments**

- `x`: An object of class `SummarizedExperiment-class`
- `features`: A vector of features (i.e. row.names) to include. Use `features=NULL` to include all.
- `assayName`: The name(s) of the assay(s) to use. If NULL and the assays are named, all of them will be included.
- `colDat.columns`: The colData columns to include (defaults includes all). Use `colDat.columns=NA` in order not to include any.
- `rowDat.columns`: The rowData columns to include (default all). Use `rowData=NA` to not include any.
- `flatten`: Logical, whether to flatten nested data.frames.
- `baseDF`: Logical, whether to return a base data.frame (removing columns containing other objects such as atomic lists). Filtering is applied after flattening.

**Value**

A data.frame (or a DataFrame).

**Examples**

```r
data("Chen2017", package="sechm")
head(meltSE(Chen2017,"Fos"))
```

---

`qualitativeColors` `qualitativeColors`

**Description**

`qualitativeColors`

**Usage**

`qualitativeColors(names, ...)`

**Arguments**

- `names`: The names to which the colors are to be assigned, or an integer indicating the desired number of colors
- `...`: passed to `randomcoloR::distinctColorPalette`

**Value**

A vector (eventually named) of colors
resetAllSechmOptions

**Description**
Resets all package options

**Usage**
resetAllSechmOptions()

**Value**
None

**Examples**
resetAllSechmOptions()

safescale

**Description**
Equivalent to 'base::scale', but handling missing values and null variance a bit more elegantly.

**Usage**
safescale(x, center = TRUE, byRow = FALSE)

**Arguments**
- `x`: A matrix.
- `center`: Logical, whether to center values.
- `byRow`: Logical, whether to scale by rows instead of columns.

**Value**
A scaled matrix.

**Examples**
m <- matrix(rnorm(100), nrow=10)
m.scaled <- safescale(m)
Description

ComplexHeatmap wrapper for `SummarizedExperiment-class`.

Usage

```r
sechm(
  se,
  features,
  do.scale = FALSE,
  assayName = NULL,
  name = NULL,
  sortRowsOn = seq_len(ncol(se)),
  cluster_cols = FALSE,
  cluster_rows = is.null(sortRowsOn),
  toporder = NULL,
  hmcols = NULL,
  breaks = .getDef("breaks"),
  gaps_at = NULL,
  gaps_row = NULL,
  left_annotation = NULL,
  right_annotation = NULL,
  top_annotation = NULL,
  bottom_annotation = NULL,
  anno_colors = list(),
  show_rownames = NULL,
  show_colnames = FALSE,
  isMult = FALSE,
  show_heatmap_legend = !isMult,
  show_annotation_legend = TRUE,
  mark = NULL,
  na_col = "white",
  anrorow_title_side = ifelse(show_colnames, "bottom", "top"),
  annocol_title_side = "right",
  includeMissing = FALSE,
  sort.method = "MDS_angle",
  ...
)
```

Arguments

- `se` A `SummarizedExperiment-class`.
- `features` A vector of features (i.e. row names of `se`). Alternatively, can be a list of feature sets, in which case these will be plotted as different row chunks.
do.scale Logical; whether to scale rows (default FALSE).
assayName An optional vector of assayNames to use. The first available will be used, or the first assay if NULL.
name The name of the heatmap, eventually appearing as title of the color scale.
sortRowsOn Sort rows by MDS polar order using the specified columns (default all)
cluster_cols Whether to cluster columns (default F)
cluster_rows Whether to cluster rows; default FALSE if 'do.sortRows=TRUE'.
toporder Optional vector of categories on which to supra-order when sorting rows, or name of a ‘rowData’ column to use for this purpose.
hmcols Colors for the heatmap.
breaks Breaks for the heatmap colors. Alternatively, symmetrical breaks can be generated automatically by setting ‘breaks’ to a numerical value between 0 and 1. The value is passed as the ‘split.prop’ argument to the getBreaks function, and indicates the proportion of the points to map to a linear scale, while the more extreme values will be plotted on a quantile scale. ‘breaks=FALSE’ will disable symmetrical scale and quantile capping, while retaining automatic breaks. ‘breaks=1’ will produce a symmetrical scale without quantile capping.
gaps_at Columns of ‘colData’ to use to establish gaps between columns.
gaps_row Passed to the heatmap function; if missing, will be set automatically according to toporder.
left_annotation Columns of ‘rowData’ to use for left annotation. Alternatively, an ‘HeatmapAnnotation’ object.
right_annotation Columns of ‘rowData’ to use for left annotation. Alternatively, an ‘HeatmapAnnotation’ object.
top_annotation Columns of ‘colData’ to use for top annotation. Alternatively, an ‘HeatmapAnnotation’ object. To disable (overriding defaults), use ‘top_annotation=character()’. 
bottom_annotation Columns of ‘colData’ to use for bottom annotation. Alternatively, an ‘HeatmapAnnotation’ object.
anno_colors List of colors to use for annotation.
show_rownames Whether to show row names (default TRUE if less than 50 rows to plot).
show_colnames Whether to show column names (default FALSE).
isMult Logical; used to silence labels when plotting multiple heatmaps
show_heatmap_legend Logical; whether to show heatmap legend
show_annotation_legend Logical; whether to show the annotation legend.
mark An optional vector of gene names to highlight.
na_col Color of NA values
annorow_title_side Side (top or bottom) of row annotation names
setRowAttr

annocol_title_side
   Side (left or right) of column annotation names

includeMissing
   Logical; whether to include missing features (default FALSE)

sort.method
   Row sorting method (see sortRows)

... Further arguments passed to ‘Heatmap’

Value

A a Heatmap-class.

Examples

data("Chen2017", package="sechm")
sechm(Chen2017, row.names(Chen2017)[1:10], do.scale=TRUE)

---

setRowAttr

Set rowData attribute of given rows

Description

Set rowData attribute of given rows

Usage

setRowAttr(se, values, name = "cluster", clear = TRUE, other = NA)

Arguments

se
   A 'SummarizedExperiment' object

values
   A named vector of values, where the names correspond to rows of 'se'

name
   The name of the rowData column in which to store the attribute.

clear
   Logical; whether to clear out any pre-existing such column.

other
   The value for unspecified rows (default NA)

Value

The modified 'se' object.

Examples

data("Chen2017", package="sechm")
Chen2017 <- setRowAttr(Chen2017, c("Arc"=1,"Junb"=1,"Npas4"=2))
**Description**
Sets a package-wide option for `sechm`.

**Usage**
```
setSechmOption(variable, value)
```

**Arguments**
- `variable` : The name of the variable to set
- `value` : The parameter value to save

**Value**
None

**Examples**
```
setSechmOption("hmcols", value=c("blue","black","yellow"))
```

---

**Description**
sortRows

**Usage**
```
sortRows(  
  x,  
  z = FALSE,  
  toporder = NULL,  
  na.rm = FALSE,  
  method = "MDS_angle",  
  toporder.meth = "before"  
)
```
sortRows

Arguments

- **x**: A numeric matrix or data.frame.
- **z**: Whether to scale rows for the purpose of calculating order.
- **toporder**: Optional vector of categories (length=nrow(x)) on which to supra-order when sorting rows.
- **na.rm**: Whether to remove missing values and invariant rows.
- **method**: Seriation method; 'MDS_angle' (default) or 'R2E' recommended.
- **toporder.meth**: Whether to perform higher-order sorting 'before' (default) or 'after' the lower-order sorting.

Value

A reordered matrix or data.frame.

Examples

```r
# random data
m <- matrix( round(rnorm(100,mean=10, sd=2)), nrow=10,
             dimnames=list(LETTERS[1:10], letters[11:20]) )

m

sortRows(m)
```
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