

Package ‘sechm’

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Type Package

Title sechm: Complex Heatmaps from a SummarizedExperiment

Version 1.17.1

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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crossHm	<i>crossHm</i>
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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] )
```

data	<i>Example dataset</i>
------	------------------------

Description

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

Value

a [SummarizedExperiment-class](#).

References

Chen et al. 2017. Mapping Gene Expression in Excitatory Neurons during Hippocampal Late-Phase Long-Term Potentiation *Frontiers in Molecular Neuroscience*. DOI: 10.3389/fnmol.2017.00039

getBreaks	<i>getBreaks</i>
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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)

Value

A vector of breaks of length = 'n'

Examples

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

getDEA

getDEA

Description

Extracts (standardized) DEA results from the rowData of an SE object.

Usage

```
getDEA(se, dea = NULL, homogenize = FALSE, sort = TRUE)
```

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
sort	Logical; whether to return the table sorted by significance

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

```
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

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

homogenizeDEA

homogenizeDEA

Description

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

Usage

```
homogenizeDEA(x)
```

Arguments

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

Value

A standardized data.frame.

log2FC	<i>log2FC</i>
--------	---------------

Description

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

Usage

```
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

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

meltSE

*meltSE***Description**

Melts a SE object into a [ggplot](#)-ready long data.frame.

Usage

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

Arguments

<code>x</code>	An object of class SummarizedExperiment-class
<code>features</code>	A vector of features (i.e. row.names) to include. Use ‘features=NULL’ to include all.
<code>assayName</code>	The name(s) of the assay(s) to use. If NULL and the assays are named, all of them will be included.
<code>colDat.columns</code>	The colData columns to include (defaults includes all). Use ‘colDat.columns=NA’ in order not to include any.
<code>rowDat.columns</code>	The rowData columns to include (default all). Use ‘rowData=NA’ to not include any.
<code>flatten</code>	Logical, whether to flatten nested data.frames.
<code>baseDF</code>	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

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

qualitativeColors	<i>qualitativeColors</i>
-------------------	--------------------------

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	<i>resetAllSechmOptions</i>
----------------------	-----------------------------

Description

Resets all package options

Usage

```
resetAllSechmOptions()
```

Value

None

Examples

```
resetAllSechmOptions()
```

safescale	<i>safescale</i>
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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)
```

sechm	<i>sechm</i>
-------	--------------

Description

ComplexHeatmap wrapper for [SummarizedExperiment-class](#).

Usage

```
sechm(
  se,
  features,
  do.scale = FALSE,
  assayName = NULL,
  name = NULL,
  sortRowsOn = NULL,
  cluster_cols = FALSE,
  cluster_rows = NULL,
  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",
    annorow_title_side = ifelse(show_colnames, "bottom", "top"),
    annocol_title_side = "right",
    includeMissing = FALSE,
    sort.method = "MDS_angle",
    ...
)

```

Arguments

<code>se</code>	A SummarizedExperiment-class .
<code>features</code>	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.
<code>do.scale</code>	Logical; whether to scale rows (default FALSE).
<code>assayName</code>	An optional vector of assayNames to use. The first available will be used, or the first assay if NULL.
<code>name</code>	The name of the heatmap, eventually appearing as title of the color scale.
<code>sortRowsOn</code>	Sort rows by MDS polar order using the specified columns (default all)
<code>cluster_cols</code>	Whether to cluster columns (default F)
<code>cluster_rows</code>	Whether to cluster rows; default FALSE if 'do.sortRows=TRUE'.
<code>toporder</code>	Optional vector of categories on which to supra-order when sorting rows, or name of a 'rowData' column to use for this purpose.
<code>hmcols</code>	Colors for the heatmap.
<code>breaks</code>	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.
<code>gaps_at</code>	Columns of 'colData' to use to establish gaps between columns.
<code>gaps_row</code>	Passed to the heatmap function; if missing, will be set automatically according to toporder.
<code>left_annotation</code>	Columns of 'rowData' to use for left annotation. Alternatively, an 'HeatmapAnnotation' object.
<code>right_annotation</code>	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
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	<i>Set rowData attribute of given rows</i>
------------	--

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))
```

setSechmOption	<i>setSechmOption</i>
----------------	-----------------------

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("hmcsls", value=c("blue","black","yellow"))
```

sortRows

*sortRows***Description**

sortRows

Usage

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

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	Serialization 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

```
# 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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