

coRNAi

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BoxPlotShorth

Boxplot with horizontal bars at the midpoint of the shorth

Description

Produces a boxplot, but instead of horizontal bars at the median, the bars are at the midpoint of the shorth.

Usage

```
BoxPlotShorth(formula, data = NULL, ...)
```

Arguments

formula	formula for how the boxplot should be drawn.
data	the data to be used
...	other arguments to be passed to the plot function

Value

a boxplot object

Author(s)

Elin Axelsson

See Also

See Also [boxplot.formula](#)

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral= "Fluc")
BoxPlotShorth(value~replicate, df)
```

InteractGraph *functions to visualize interactions as a graph*

Description

visualizes significant interactions as a graph

Usage

```
InteractGraph(toptable, thresh, sizecutoff=0, by, key=FALSE, file="interactions", color
```

Arguments

toptable	toptable table from function topTable
thresh	thresh numeric, threshold for significance
sizecutoff	sizecutoff a minimal absolute size of a interaction for it to be included in the graph as an edge.
by	by column in topTable that thresh should be applied to
key	key optional, data frame with groupings of the genes in the toptable
file	file name of the file the results will be outputed to.
colors	colors list with colors to be used for pos interactions, neg interactions, key (nodes in key) and normal nodes.

Value

pdf file with graph

Author(s)

Elin Axelsson

See Also

[levelplot](#)

Examples

```
#see vignette
```

InteractLevelPlot *function to visualize interactions as a levelplot*

Description

visualizes significant interactions as a levelplot

Usage

```
InteractLevelPlot(toptable, thresh = 0.001, by = "P.Value", key = FALSE, col.reg
```

Arguments

toptable	toptable table from function topTable
thresh	thresh numeric, threshold for significance
by	by column in topTable that thresh should be applied to
key	key optional, data frame with groupings of the genes in the toptable
col.regions	col.regions colors to be used
zerolimit	zerolimit threshold below which interactions should be colored as 0.

Value

a levelplot, pdf files with graphs

Author(s)

Elin Axelsson

See Also

[levelplot](#)

Examples

```
# simulated data
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
tt = data.frame("ID"=(unique(df$Pair[df$Type=="comb"])), "size"=runif(length(unique(df$Pair)))
InteractLevelPlot(tt, thres=0, by="size")
```

MainFitPlot

Diagnostic plot

Description

Plots residuals vs fitted values after fitting of main effects.

Usage

```
MainFitPlot(fit, xlab = "Fitted values", ylab = "Residuals", sd.fit = TRUE, main
```

Arguments

<code>fit</code>	a fit from <code>lmmain</code> , <code>rlmmain</code> or similar
<code>xlab</code>	label for x-axis
<code>ylab</code>	label for y-axis
<code>sd.fit</code>	logical, should the local estimator of the standard deviation be plotted
<code>main</code>	main title for the plot
<code>...</code>	arguments to be passed on to the plot function

Value

a plot

Author(s)

Elin Axelsson

See Also

[locfit](#)

Examples

```
## simulated data

fitted.value = rnorm(100,2,1)
residuals = rnorm(100,0,1)
fit = list(fitted.value=fitted.value, residuals = residuals)
class(fit) = "lm"
MainFitPlot(fit)
```

PlotHeatmap *Plot a heatmap of interactions*

Description

Plots a heatmap of the mean residuals for each interaction pair.

Usage

```
PlotHeatmap(toptable, colpal = colorRampPalette(c("blue", "white", "yellow")),  
key=FALSE, margins=c(7, 7), na.color="grey", breaks=seq(-1, 1, by=0.01), ...)
```

Arguments

toptable	a data frame created by with the interaction estimates as "logFC" and pair id as "ID". Usually created by topTable function in limma
colpal	color palette to be used in the plot
key	logical should a color key be included
margins	margins for plot
na.color	color for NA values
breaks	mapping data to colors in colpal
...	additional arguments to be passed to heatmap.2 call

Value

a plot

Author(s)

Elin Axelsson

See Also

[heatmap.2](#)

Examples

```
# simulated data  
data(screen1_raw)  
df = cellHTS2df(screen1_raw, neutral="Fluc")  
tt = data.frame("ID"=(unique(df$Pair[df$Type=="comb"])), "size"=runif(length(unique(df$Pair))  
PlotHeatmap(tt)
```

Pplot

Function to plot cumulative p-values

Description

Given a vector of p-values a cumulative p-value plot is produced

Usage

```
Pplot(x, col = "darkblue", maintitle="", nrpoints = 100, ...)
```

Arguments

x	vector with p-values
col	color to be used
maintitle	character, main plot title
nrpoints	numeric, how many points should be plotted
...	additional arguments passed on to the plot

Value

a plot

Author(s)

Wolfgang Huber

Examples

```
x = runif(1000,0,1)
Pplot(x,col="red",maintitle="uniform dist.", nrpoints=200)
```

cellHTS2df

converts cellHTS objects to dataframes

Description

converts a cellHTS2 object into a data.frame object and check that all mandatory meta data is included. Adds Directions, replicates and Pair columns.

Usage

```
cellHTS2df(x, neutral)
```

Arguments

x	a cellHTS object with correct annotations
neutral	string stating which RNAi is neutral (negative control)

Value

data.frame, with the data from the cellHTS object in column "value". Meta data from annotation file and the new columns; Directions, Replicate and Pair

Author(s)

Elin Axelsson

See Also

[cellHTS](#)

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
head(df)
```

cortestmatrices	<i>Function to extract correlations and corresponding p-values from interaction matrix.</i>
-----------------	---------------------------------------------------------------------------------------------

Description

This is a wrapper function for `cor.test`, given a matrix of interaction values, correlations and corresponding p-values for the genewise interaction profiles are calculated.

Usage

```
cortestmatrices(mat, method = c("pearson", "kendall", "spearman"))
```

Arguments

mat	mat interaction matrix
method	character deciding which correlation method should be used

Value

List of two matrices

cor.matrix	matrix with correlations
p.matrix	matrix with p-values

Author(s)

Elin Axelsson

See Also

[cor.test](#)

Examples

```
## simulate data with 2 genes with similar profiles

mat = matrix(rnorm(100*100,0,1),100,100)
pr = sample(2:10,100,replace=TRUE)
mat[1:2,] = mat[1:2,] + matrix(pr,ncol=100,nrow=2,byrow=TRUE)
mat = mat+t(mat)
diag(mat) = NA
dimnames(mat)=list(1:100,1:100)
res = cortestmatrices(mat,method="spearman")
cors= res[[1]]
ps = res[[2]]
print(which(ps==min(ps,na.rm=TRUE),arr.ind=TRUE))
```

data2graph

Function to create .dot files for graph representation of data

Description

From a interaction table or list of data matrices a .dot file is created for visualisation of the interaction/correlation network

Usage

```
data2graph(indata, sizethres=0, thres, thresBy = "P.Value", cols = c("blue", "wh
```

Arguments

indata	
sizethres	numerical, lower treshold on the absolute effect size for edges
thres	threshold that should be used for interactions/correlations to be included in graph
thresBy	what data should the the threshold by used at. By default the p value from the moderated t test is used but one could also use e.g. the ordinary t or the size
cols	colors to be used in the plot, should be a character vector with the colors for low, neutral and hig values
gamma.col	Factor used to scale the colors
scaleFactor	Scale factor to adjust the distances beeten nodes in the graph
nodecolor	character or character vector, which color(s) should the nodes have. Should either be of lenth 1 (all nodes same color) or same length as the number of nodes.
writedot	logical, should a .dot file be created.
filename	charcter string with name of .dot file
width	width of the nodes
penwidth	width of the lines in the plot
shape	shape of the nodes in the plot
fixedsize	should all nodes have the same size
fontsize	size of the font in the plot

Value

a .dot file is written if writefile argument is TRUE

ninf Dataframe with information about the nodes in the network
einfo Dataframe with information about the edges in the network

Author(s)

Greg Pau, Elin Axelsson

See Also

Also see Graphviz documantation <http://www.graphviz.org/>

Examples

```
#see vignette
```

df2array *function to do go from data frame to an array with values*

Description

The function transforms a float column in a data frame (from cHTS2df) to an array.

Usage

```
df2array(df, what)
```

Arguments

df Data frame (from cellHTS2df) with additional column weights.
what which column contains the data, eg. value or residuals

Value

An array with the data from the data frame.

Author(s)

Elin Axelsson

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral = "Fluc")
df$weights = as.numeric(df$type=="comb")

aa = df2array(df, what="value")

## see head for first replicate

head(aa[, , 1])
```

df2fitmatrix

matrix for lmFit from dataframe

Description

converts a dataframe into the right format for lmFit function

Usage

```
df2fitmatrix(df)
```

Arguments

df dataframe from cellHTS2df function

Value

A matrix with the genepairs as rows and the replicates as columns. This matrix is in the right format for the lmFit.

Author(s)

Elin Axelsson

See Also

[df2lmFit](#)

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
df = weightDf(df)
lmm = lmmain(df)
df = updateDf(df, lmm)
mfit = df2fitmatrix(df)
head(mfit)
```

df2lmFit	<i>lmFit from dataframe</i>
----------	-----------------------------

Description

converts a dataframe into the right format for lmFit function, calls the lmFit from limma and returns the result.

Usage

```
df2lmFit(df)
```

Arguments

df dataframe from cellHTS2df function

Value

Object of class 'MArrayLM'

Author(s)

Elin Axelsson

See Also

[lmFit](#)

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
df = weightDf(df)
lmm = lmmain(df)
df = updateDf(df, lmm)
mfit = df2lmFit(df)
str(mfit)
```

estmodel	<i>Function to do estimate the main effects from data using median, mean or shorth.</i>
----------	-----------------------------------------------------------------------------------------

Description

The function estimates the main effect i from all data with the RNi against i . It can be done by median, mean or shorth.

Usage

```
estmodel(df, estimate = c("median", "mean", "shorth"), per = NULL)
```

Arguments

`df` data frame from cellHTS2df function with extra column weight (see weightDf)

`estimate` estimate median, mean or shorth, decides how the main effects will be estimated.

`per` per for which factor should the analysis be done separately, eg. batch or replicate.

Value

for `per = NULL`, a list with

`coefficient` the estimated main effects

`residuals` the residual after the main effects have been subtracted from the observations

for other `per`, a list of lists like the once described above, one for each level of the factor `per`.

Author(s)

Elin Axelsson

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
df$weight = as.numeric(df$type=="comb")
main = estmodel(df, estimate="median")
str(main)
```

faultyscreen *faulty screen*

Description

A screen with both systematic errors and sporadic contaminations.

Usage

```
data(faultyscreen)
```

Format

The format is: chr "cellHTS"

Examples

```
data(faultyscreen)
## maybe str(faultyscreen) ; plot(faultyscreen) ...
```

interactiontable *Returns a list of interactions with associated statistics.*

Description

This is an extended wrapper around the `topTable` function from the `limma` package, as an option the ordinary t statistics can be calculated as well.

Usage

```
interactiontable(ebfit, sort = "none", ord.t = FALSE, correction = "BH")
```

Arguments

<code>ebfit</code>	ebfit a MArrayLM object produced by the <code>eBayes</code> function
<code>sort</code>	character string specifying which statistic to rank genes by, possible arguments are <code>none</code> , <code>ID</code> , <code>size</code> , <code>t</code> , <code>B</code> , <code>adj.P.val</code> , <code>P.Value</code> , and if <code>ord.t = TRUE</code> : <code>ord.t</code> , <code>ord.p</code> and <code>ord.p.adj</code> .
<code>ord.t</code>	Logical, should ordinary t statistics be calculated? Default is <code>FALSE</code> .
<code>correction</code>	method used to adjust the p-values for multiple testing. Default is <code>BH</code> . See <code>p.adjust</code> for the complete list of options.

Value

Returns a dataframe where the rows are the interaction pairs and the columns the statistics:

`ID`: Interaction pair id

`size`: the average interaction size

`t`: the moderated t statistics

`P.Value`: p-value for the moderated t statistics

`adj.P.Val`: adjusted p-value

`B`: the b statistics

if the `ord.t=TRUE`, the ordinary t statistics (`ord.t`), with corresponding p-values (`ord.p`) and adjusted p-values (`ord.p.adj`)

Warning

usage of the ordinary t statistics is not recommended for data sets with few replicates.

Author(s)

Elin Axelsson

See Also

[p.adjust](#), [topTable](#)

Examples

```
## simulated data
y <- matrix(rnorm(50*4, sd=1), 50, 4)
rownames(y) <- paste("Pair", 1:50)

# fit and eBayes
fit <- lmFit(y)
fit <- eBayes(fit)
tt = interactiontable(fit, sort="size")
head(tt)
```

key	<i>A key to data set screen1, contains (additional) information about the genes in the screen.</i>
-----	----------------------------------------------------------------------------------------------------

Description

Contains information about which of the 16 genes in screen1 are cell cycle related. This is used in interaction graphs/plots.

Usage

```
data(key)
```

Format

A data frame with 16 observations on the following 2 variables.

```
GeneID a factor with levels AnnIX CG12785 CG16935 CG3165 CG7889 CG8108 CSN3
      CSN4 CSN5 fwd pbl Rbf Rho1 sos trbl zip
cellCycle a numeric vector
```

Examples

```
data(key)
table(key$cellCycle)
```

replots	<i>reproducibility plots</i>
---------	------------------------------

Description

plots reproducibility of replicates within/between screens

Usage

```
BetweenScreenPlot(df, what="value", names, smooth=TRUE)
WithinScreenPlot(df, what="value", main="within-screen replicates", ylab="technic
```

Arguments

<code>df</code>	<code>df</code> dataframe created by the <code>cellHTS2df</code> function
<code>names</code>	<code>names</code> optional, character vector with names of the different screens.
<code>what</code>	<code>what</code> what should be plotted, eg value or residuals
<code>main</code>	<code>main</code> main title
<code>ylab</code>	<code>ylab</code> label for y-axis
<code>xlab</code>	<code>xlab</code> label for x-axis
<code>smooth</code>	<code>smooth</code> should the <code>smoothScatter</code> function be called. Default is TRUE.
<code>...</code>	<code>...</code> further argument to be passed to the plot function

Value

pairs plot

Author(s)

Elin Axelsson

See Also

[pairs,plot](#)

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
BetweenScreenPlot(df)
WithinScreenPlot(df)
```

LS main

main effect estimation

Description

for `rlmmain` the main effects are estimated using `rlm` function from MASS package, with `lmmain` the OLS is used.

Usage

```
rlmmain(df, per=NULL)
lmmain(df, per=NULL)
```

Arguments

<code>df</code>	<code>df</code> dataframe created by function <code>cellHTS2df</code>
<code>per</code>	string argument for which factor the analysis should be done separately, eg. replicate or batch

Value

lm,rlm

Author(s)

Elin Axelsson

See Also

[rlm,lm](#)

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw,neutral="Fluc")
df = weightDf(df,exclude=c("controlP1","controlP2","controlN1","controlN2","controlP1N1",
mains = rlmmain(df)
hist(coef(mains))
```

screen1_raw

screen1 raw data

Description

cellHTS2 object containing the raw data from screen1 (cellcycle related)

Usage

```
data(screen1_raw)
```

Format

The format is: chr "cellHTS"

Examples

```
data(screen1_raw)
state(screen1_raw)
```

screen2_raw	<i>screen2 raw data</i>
-------------	-------------------------

Description

cellHTS2 object containing the raw data from screen2 (phospatases)

Usage

```
data(screen2_raw)
```

Format

The format is: chr "cellHTS"

Examples

```
data(screen2_raw)
state(screen2_raw)
```

signalplots	<i>plot variation vs signal intensity</i>
-------------	-------------------------------------------

Description

plots the variation of replicates vs the mean intensity either by within screen replicate separately or over all screen replicates.

Usage

```
SDplot(df, xlab="intensity mean", ylab="sd", add=FALSE, main, ...)
MAplot(df, main, rank=FALSE)
```

Arguments

df	df dataframe created by cellHTS2df function
main	main character string to be used as main title
xlab	xlab label for x-axis
ylab	ylab label for y-axis
add	add logical, should result be added to existing plot
rank	rank if TRUE the rank of the average intensities will be used
...	... further arguments to be passed to the plot function.

Value

plot

Author(s)

Elin Axelsson

See Also[plot](#)**Examples**

```
data(screen1_raw)
df=cellHTS2df(screen1_raw, neutral="Fluc")
MAplot(df, main="raw data")
SDplot(df, main="raw data")
```

`tt2matrix`*Extracting data from a toptable and format it to matrix*

Description

Given an dataframe with data, typically from the `interactiontable`, the gene pair data is converted to a symmetric matrix.

Usage

```
tt2matrix(toptable, what)
```

Arguments

<code>toptable</code>	a dataframe with data for the pairwise interactions. Typically from the <code>interactiontable</code> function.
<code>what</code>	character indicating which of the columns in the dataframe should be used in the matrix.

Value

a symmetric matrix with the selected data for gene pair i,j in `matrix[i,j]` and `matrix[j,i]`

Author(s)

Elin Axelsson

Examples

```
## simulated data
mytoptable = data.frame("ID" = c("A B", "A C", "B C"), "size"=c(1:3), stringsAsFactors=FALSE)
mat = tt2matrix(mytoptable, what="size")
```

 updateDf

updates dataframe after fitting of main estimates

Description

adds residuals (value-main effects) to a dataframe

Usage

```
updateDf(df, lm, per=NULL)
```

Arguments

df	df dataframe created by cellHTS2df function
lm	lm lm objects, residuals from fitting main effects to data
per	per string argument, for which factor the analysis was done separately, eg. replicate.

Value

data frame with a new column "residuals"

Author(s)

Elin Axelsson

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
df = weightDf(df, exclude=c("double", "controlP1", "controlP2", "controlN1", "controlN2", "controlN3"))
lmain = lmmain(df)
df = updateDf(df, lmain)
hist(df$residuals)
```

 weightDf

Function to indicate which data points should be involved in downstream analysis.

Description

Function to do add weights to the data points in a data frame. At the time being 0 means excluded and everything >0 means included.

Usage

```
weightDf(df, exclude = c("double", "controlN2", "controlP2", "controlP1N1", "controlP1N2", "controlP1N3"))
```

Arguments

df data frame from cellHTS2df
exclude which type of data should be excluded from analysis.

Details

See also vignette for information about different "Type" types.

Value

A data frame with an added column 'weight'

Author(s)

Elin Axelsson

Examples

```
data(screen1_raw)
df = cellHTS2df(screen1_raw, neutral="Fluc")
##stupid example
df = weightDf(df, exclude="controlN2")
head(df)
```

Index

*Topic **datasets**

faultyscreen, 12
key, 14

*Topic **hplot**

MainFitPlot, 4
replots, 14

*Topic **models**

df2lmFit, 11

BetweenScreenPlot (*replots*), 14

boxplot.formula, 1
BoxPlotShorth, 1

cellHTS, 7
cellHTS2df, 6
cor.test, 7
cortestmatrices, 7

data2graph, 8
df2array, 9
df2fitmatrix, 10
df2lmFit, 10, 11

estmodel, 11

faultyscreen, 12

heatmap.2, 5

InteractGraph, 2
interactiontable, 13
InteractLevelPlot, 3

key, 14

levelplot, 2, 3
lm, 16
lmFit, 11
lmmain (*LS main*), 15
locfit, 4
LS main, 15

MainFitPlot, 4
MAplot (*signalplots*), 17

p.adjust, 13

pairs, 15
plot, 15, 18
PlotHeatmap, 5
Pplot, 6

replots, 14
rlm, 16
rlmmain (*LS main*), 15

screen1_raw, 16
screen2_raw, 17
SDplot (*signalplots*), 17
signalplots, 17

topTable, 13
tt2matrix, 18

updateDf, 19

weightDf, 19
WithinScreenPlot (*replots*), 14