

Package ‘rhcoclust’

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Title Robust Hierarchical Co-Clustering to Identify Significant Co-Cluster

Version 1.1.0

Description Here we performs robust hierarchical co-clustering between row and column entities of a data matrix in absence and presence of outlying observations. It can be used to explore important co-clusters consisting of important samples and their regulatory significant features. Please see Hasan, Badsha and Mollah (2020) <[doi:10.1101/2020.05.13.094946](https://doi.org/10.1101/2020.05.13.094946)>.

License GPL (>= 2)

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plot_rhcoclust	<i>Plot of the 'rhcoclust' objects</i>
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Description

This function used for two plots from output of rhcoclust (i) plot results for gene (row) and compound (column) co-cluster graph, and (ii) plot graph of QCC for identification of biomarker co-cluster.

Usage

```
plot_rhcoclust(CoClustObj, plot.cocluster = FALSE, plot.ccim = FALSE,
cex.xaxis = 0.7, cex.yaxis = 0.5)
```

Arguments

CoClustObj	Output objects from rhcoclust
plot.cocluster	To set no plotting as the default for cocluster.
plot.ccim	To set no plotting as the default for icc.
cex.xaxis	A numerical value giving to control/annotation text size in x-axis. Default is 0.7.
cex.yaxis	A numerical value giving to control/annotation text size in y-axis. Default is 0.5.

Value

Plots

Author(s)

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See Also

[rhcoclust](#) for generating a graph objects for clustering network

Examples

```
# Load necessary library
library(rhcoclust)
library(fields)

# Load real data
data("toxygates_data")

# Load predefined simulated data
data("simu_data")

# Data analysis using real data by rhcoclust. Note, for analysis of predefined
# simulated data use data <- simu_data
data <- toxygates_data

# Apply rhcoclust to identify significant co-cluster of samples and their regulatory features
CoClustObj <- rhcoclust(data, rk = 3, ck = 3, method.dist = "manhattan", method.hclust = "ward.D")

# Plot co-cluster
# Please use par(mar=c(6, 10, 3, 6)) or modify if needed for best fit of the graph
# mar order: bottom, left, top, and right
plot_rhcoclust (CoClustObj, plot.coclust = TRUE, plot.ccim = FALSE,
cex.xaxis = 0.7, cex.yaxis = 0.5)

# Plot CCIM
# Please use dev.off() to avoid the figure margin from previous plot
plot_rhcoclust (CoClustObj, plot.coclust = FALSE, plot.ccim = TRUE)
# Please add legend with change or add any parameters if needed.
legend("topleft",
      legend = c("Upper-significant", "Insignificant", "Down-significant"),
      col = c("red", "black", "blue"),
      bty = "n",
      pch = c(20, 20, 20),
      pt.cex = 2,
      cex = 1.2,
      x.intersp = 0.2,
      y.intersp = 0.4,
      text.col = "black",
      horiz = FALSE,
      inset = c(0.3, -0.08))
```

reversestring

Reverse given string

Description

This is the function for reverse string

Usage

```
reversestring(string, n = 1)
```

Arguments

string	Given string or seq
n	By which n-plets we should reverse the given string

Details

This function is used to reverse given string or seq

Value

Reversed string or seq

Author(s)

Md. Bahadur Badsha <mbbadshar@gmail.com>

Examples

```
x1 <- c("R1C1", "R2C2", "R3C3")
reversestring(x1, 2)
```

rhcoclust

The function for co-clustering sample and feature to explore significant samples and their regulatory features

Description

Toxicogenomic studies require co-clustering to identify biomarker genes for the assessment of chemical toxicity from gene expression levels. It is also essential in the drug discovery experiments. However, gene expression datasets are often contaminated by outliers due to several steps involve in the data generating process. This package performs robust hierarchical co-clustering between row and column entities of a data matrix by reducing the influence of outlying observations. It can be used to explore biomarker genes those are divided into upregulatory and downregulatory groups by the influence of different chemical compounds groups more accurately. It can also provide the statistical significance of the identified co-clusters.

Usage

```
rhcoclust(data, rk, ck, method.dist = "manhattan", method.hclust= "ward.D")
```

Arguments

data	A data matrix containing data having the characteristics of interval and ratio level of measurement or continuous data
rk	Number of clusters in the row entities of the data matrix
ck	Number of clusters in the column entities of the data matrix

- `method.dist` The distance measure to be used. The default is "manhattan". The other options are "euclidean", "maximum", "canberra", "binary" or "minkowski". Any unambiguous substring can be given.
- `method.hclust` The agglomeration method to be used. The default is "ward.D". The other options are "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC).

Value

A [list](#) of object that containing the following:

`Coclust_MeanMat` : A data frame containing combination of row and column cluster number in the first column and their ranked co-cluster mean in the second column. In the first column first number indicates row cluster index and second number indicates column cluster index, respectively.

`CoClSDtMat` : The reorganized transformed data matrix to generate co-cluster graph.

`NG_Cocls` : The index of gene/row names.

`NC_Cocls` : The index of column names.

`rowclust` :The gene/row entity clusters.

`colclust` : The column entity clusters.

`colorsG`: Colors of genes/row entity clusters to generate co-cluster graph.

`colorsC`: Colors of DCCs/column entity clusters to generate co-cluster graph.

`CentralLine`: Central Line of individual control chart to generate graph of control chart and to identify significant co-clusters.

`UpContLimit`: Upper Control Limit to generate graph of control chart and to identify significant co-clusters.

`LowrContLimit`: Lower Control Limit to generate graph of control chart and to identify significant co-clusters.

`color`: Colors to generate individual control chart.

`pchmark`: Shape of points to generate individual control chart.

Author(s)

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Examples

```
# Load necessary library
library(rhcoclust)
library(fields)

# Load real data
data("toxygates_data")

# Load predefined simulated data
data("simu_data")

# Data analysis using real data by rhcoclust. Note, for analysis of predefined
```

```
# simulated data use data <- simu_data
data <- toxygates_data

# Apply rhcoclust to identify significant co-cluster of samples and their regulatory features
CoClustObj <- rhcoclust(data, rk=3, ck=3, method.dist= "manhattan", method.hclust = "ward.D")

# A data frame containing combination of row and column cluster number in the first
# column and their ranked co-cluster mean in the second cluster.
GC_cls_MeanMat <- CoClustObj$NGC_cls_MeanMat

# The reorganized transformed data matrix to generate co-cluster graph.
CoClsDtMat <- CoClustObj$CoClsDtMat

# The gene/row entity clusters.
rowclust <- CoClustObj$rowclust

# The column entity clusters.
colclust <- CoClustObj$colclust

# Colors of genes/row entity clusters to generate co-cluster graph
colorsG <- CoClustObj$colorsG

# Colors of DCCs/column entity clusters to generate co-cluster graph
colorsC <- CoClustObj$colorsC

# Central Line of individual control chart to generate graph of control chart and to
# identify significant co-clusters.
CntrLine_QC <- CoClustObj$CentralLine

# Upper Control Limit to generate graph of control chart and to identify significant
# co-clusters.
UCL_QC <- CoClustObj$UpContLimit

# Lower Control Limit to generate graph of control chart and to identify significant
# co-clusters.
LCL_QC <- CoClustObj$LowrContLimit

# Colors to generate individual control chart.
ColorQC <- CoClustObj$color

# Shape of points to generate individual control chart.
PcmQC <- CoClustObj$pchmark

# Plot co-cluster
# par(mar=c(6,10,3,6)) # Modify if needed
# mar order: bottom, left, top, and right
plot_rhcoclust (CoClustObj, plot.coclust = TRUE, plot.ccim = FALSE)

# Plot CCIM
# use dev.off() to avoid the figure margin from previous plot
plot_rhcoclust (CoClustObj, plot.coclust = FALSE, plot.ccim = TRUE)
```

rhcoclust_internet *Interaction network (internet) of the 'rhcoclust' objects*

Description

This function is used for visualization of clustering interaction network plot for the objects that are generated by rhcoclust and list of up-regulated and down-regulated variables list. There are three layers in the network plot, (i) genes (rows) are shown in the first layer, (ii) co-cluster are in the second layer and (iii) columns (compounds) are in the last layer. Red and blue color indicates up and down regulated respectively.

Usage

```
rhcoclust_internet(data, CoClustObj, plot.internet = FALSE,  
cex.nodes = 0.7, edge.width = 1)
```

Arguments

data	A data matrix containing data having the characteristics of interval and ratio level of measurement or continuous data
CoClustObj	A list of output objects from rhcoclust
plot.internet	To set no plotting as the default for interaction network (internet)
cex.nodes	A numerical value giving to control/annotation node size in the network. Default is 0.7.
edge.width	A numerical value giving to control/annotation edge width in the network. Default is 1.

Value

Plot A [list](#) up-regulated and down-regulated variables.

Author(s)

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See Also

[rhcoclust](#) for generating a graph objects for clustering network

Examples

```
# Load necessary library  
library(rhcoclust)  
library(fields)  
  
# Load real data  
data("toxygates_data")
```

```
# Load predefined simulated data
data("simu_data")

# Data analysis using real data by rhcoclust. Note, for analysis of predefined
# simulated data
data <- simu_data
# Real data use: data <- toxygates_data

# Apply rhcoclust to identify significant co-cluster of samples and their regulatory features
CoClustObj <- rhcoclust(data, rk=4, ck=3, method.dist = "manhattan", method.hclust = "ward.D")

# Plot interaction network (internet)
# Please use dev.off() to avoid the figure margin from previous plot
# mar order: bottom, left, top, and right
# please use par(mar=c(5,2,5,2)) or modify when necessary to best fit for the plot
Netrhcoclust <- rhcoclust_internet(data, CoClustObj = CoClustObj, plot.internet =
TRUE, cex.nodes = 0.7, edge.width = 1)
# Please change or add any parameter if needed.
text(x = -1, y = 1.1, "Row Cluster", cex = 0.7)
# Please change or add any parameter if needed.
text(x = 0, y = 1.1, "Co-Cluster", cex = 0.7)
# Please change or add any parameter if needed.
text(x = 1, y = 1.1, "Column Cluster", cex = 0.7)
```

rhcoclust_network *Visualization of clustering network plot*

Description

This function is used for visualization of clustering network plot, the plot objects are generated by rhcoclust.

Usage

```
rhcoclust_network(CoClustObj, scale.threshold = 10)
```

Arguments

CoClustObj Output objects from rhcoclust
scale.threshold Edge width scaling threshold, defaults is 10

Value

Plot

Author(s)

Md. Bahadur Badsha <mbbadshar@gmail.com>

See Also

[rhcoclust](#) for generating a graph objects for clustering network

Examples

```
# Load necessary library
library(rhcoclust)
library(fields)
library(igraph)

# Load real data
data("toxygates_data")

# Load predefined simulated data
data("simu_data")

# Data analysis using real data by rhcoclust. Note, for analysis of predefined
# simulated data use data <- simu_data
data <- toxygates_data

# Apply rhcoclust to identify significant co-cluster of samples and their regulatory features
CoClustObj <- rhcoclust(data, rk = 3, ck = 3, method.dist = "manhattan", method.hclust = "ward.D")

# Visualization of clustering network plot
rhcoclust_network(CoClustObj, scale.threshold = 10)
```

simulate_data	<i>Simulate data for robust hirarchical clustering to identify significant co-cluster</i>
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Description

We generate fold change gene expression (FCGE) data according to the characteristics of toxicogenic data.

Usage

```
simulate_data(no.gene, no.dcc)
```

Arguments

no.gene	Number of genes in the simulated data.
no.dcc	Numbe of doses of chemical compounds (dcc) in the simulated data.

Details

There are four gene groups and three DCCs groups in the simulated dataset. The gene group 1, 2, 3 and 4 consists the genes G1-G10, G11-G20, G21-G30 and G31-G50 respectively and DCCs group 1, 2 and 3 consists the DCCs C1_High-C5_High- C1_Middle-C5_Middle, C6_High-C10_High-C6_Middle-C10_Middle and C1_Low-C12_Low- C11_Middle- C12_Middle- C11_High- C12_High respectively. Where, G stands for gene and C stands for chemical compound arranged in the row and column of the simulated data matrix respectively. The error term $N(0,0.35)$ from normal distribution with mean 0 and variance 0.35 is added to each element of the simulated dataset. In the simulated dataset the gene group-1 is up-regulated by the DCCs group-1, gene group-2 is up and down-regulated by the DCCs group-2 and 1 respectively. The gene group-3 is down-regulated by the DCs group-2. The gene group-4 is not regulated by any of the DCCs groups and DCCs group-3 does not influence any of the genes in the dataset.

Value

A [list](#) of object that containing the following:

SimData: A simulated data matrix as generated.

SimDataRnd: Randomly distributed row and column entity of simulated data.

GCmat: Transformed simulated data.

Author(s)

Md. Bahadur Badsha <mbbadshar@gmail.com>

Examples

```
# Load library
library(rhcoclust)

# Number of genes in the simulated data.
no.gene <- 50

# Numbe of doses of chemical compounds (dcc) in the simulated data.
no.dcc <- 12

SimulteData <- simulate_data(no.gene,no.dcc)

# A simulated data matrix as generated.
SimulteData$SimData

# A randomly distributed row and column entity of simulated data.
SimulteData$SimDataRnd

# A transformed simulated data.
SimulteData$GCmat
```

simu_data	<i>A predefined simulated data for 'rhcoclust' package</i>
-----------	--

Description

This is the predefined simulated data matrix that used as an example in 'rhcoclust' package.

Details

The column of the simulated data matrix represents doses of chemical compounds (DCCs) and row represents genes. Each element in the dataset represents fold change gene expression data.

Value

Matrix

Author(s)

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See Also

[simulate_data](#)

Examples

```
library(rhcoclust)

# Load data
data("simu_data")
```

toxygates_data	<i>A real dataset for 'rhcoclust' package</i>
----------------	---

Description

This is the real data matrix that used as an example in 'rhcoclust' package.

Details

The real data matrix collected from Nyström-Persson et al., 2013 (<https://toxygates.nibiohn.go.jp/toxygates/#columns>). The column of the data matrix represents doses of chemical compounds (DCCs) and row represents genes. Each element in the dataset represents fold change gene expression data.

Value

Matrix

Author(s)

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References

1. Nyström-Persson et al., (2013). Toxygates: interactive toxicity analysis on a hybrid microarray and linked data platform. *Bioinformatics*, Volume 29, Issue 23, Pages 3080–3086.

Examples

```
# Load library
library(rhcoclust)

# Load real data
data("toxygates_data")
```

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