Package 'RUtil'

February 17, 2011

Version 0.03
License GPL (>= 2)
Description A package collecting useful R code snippets from different projects made by Raivo Kolde.
Title Raivos util functions
Author Raivo Kolde < rkolde@gmail.com>
Maintainer Raivo Kolde <rkolde@gmail.com></rkolde@gmail.com>
Depends R (>= 2.10)
Imports plyr, reshape, stringr, ggplot2, RCurl, grid
Collate 'bubbleplot.r' 'ggplot2_themes.r' 'gprofiler.r' 'make_unique_comb.r' 'ROC.r' 'RUtil-package.r'
R topics documented: RUtil-package
bubbleplot 2 calcAUC 2 calcCurve 3
gconvert
generate_gcocoa_query
gprofiler 6 make_unique_comb 7 plotROC 8 theme_bw_raivo 9
Index 10
RUtil-package A set of useful functions by Raivo

Description

A set of useful functions by Raivo

2 calcAUC

bubbleplot

Function to draw the bubbleplots - an alternative to Venn diagram...

Description

Function to draw the bubbleplots - an alternative to Venn diagram

Usage

```
bubbleplot(lists, percentage=TRUE)
```

Arguments

lists input element lists

percentage logical showing if percentages or raw numbers are displayed

Details

Function to draw the bubbleplots - an alternative to Venn diagram

Value

Returns nothing only draws the picture

Author(s)

Raivo Kolde <rkolde@gmail.com>

Examples

```
bubbleplot(list(Asadsa = sample(letters, 14), Gadsa = sample(letters, 9)))
bubbleplot(list(Badsad = sample(letters, 16), Asadsa = sample(letters, 14), Gadsa = sample
```

calcAUC

Calculate AUC score based on the given curves...

Description

Calculate AUC score based on the given curves

Usage

```
calcAUC(curves, bootstrap=0)
```

Arguments

curves as calculated by calcCurve

bootstrap number of bootstrap samples for finding confidence intervals

calcCurve 3

Details

Calculate AUC score based on the given curves

Value

returns a data frame with auc values for each class, optionally also with bootstrapped confidence intervals

Author(s)

Raivo Kolde <rkolde@gmail.com>

calcCurve

Calculate ROC curves based on ordered class data...

Description

Calculate ROC curves based on ordered class data

Usage

calcCurve(data)

Arguments

data

Data as described in data

Details

Calculate ROC curves based on ordered class data

Value

Returns a data frame like data, but with additional columns TP and TN

Author(s)

Raivo Kolde <rkolde@gmail.com>

4 gconvert

gconvert

Convert gene ID-s.

Description

Convert gene ID-s.

Usage

```
gconvert(ids, organism="hsapiens", target="ENSG", df=T)
```

Arguments

ids gene list.

organism gene list origin.

target ID.

df logical showing if the output will be a data.frame or list.

Details

Wrapper for g:Convert web toolkit.

Value

The output can be either list or data.frame. List has an entry for every input gene. Data frame is just a two column table with inputs and corrsponding outputs. The input names may be duplicated.

Author(s)

Juri Reimand < jyri.reimand@ut.ee>, Raivo Kolde < rkolde@gmail.com>

References

J. Reimand, M. Kull, H. Peterson, J. Hansen, J. Vilo: g:Profiler - a web-based toolset for functional profiling of gene lists from large-scale experiments (2007) NAR 35 W193-W200

Examples

```
gconvert(c("Klf4", "Pax5", "Sox2", "Nanog"), organism = "mmusculus")
```

generate_gcocoa_query 5

```
generate_gcocoa_query
```

Generate query for g:Cocoa web tool...

Description

Generate query for g:Cocoa web tool

Usage

```
generate_gcocoa_query(glist)
```

Arguments

glist

a list of vectors of gene names

Details

Function that prints out a g:Gogoa query string given a list of gene name vectors.

Value

Prints out the query string

Author(s)

Raivo Kolde <rkolde@gmail.com>

Examples

```
glist = list(a = c("pax6", "klf9"), b = c("nanog", "Pou5f1")) generate_gcocoa_query(glist)
```

gorth

Find orthologs.

Description

Find orthologs.

Usage

```
gorth(genelist, source_organism="mmusculus", target_organism="hsapiens", mthresh
```

6 gprofiler

Arguments

Details

Wrapper for g:Orth web toolkit. The organism names are constructed, by combining the first letter of the name and family name. For example human - 'hsapiens' and mouse- 'mmusculus'

To alleviate the problem of having many orthologs per gene (most of them uninformative) one can set a threshold of how many results to show. The programs tries to find the most informative by selecting the most popular ones.

Value

The output can be either list or data.frame. List has an entry for every input gene. Data frame is just a two column table with inputs and corrsponding outputs. The input names may be duplicated.

Author(s)

Raivo Kolde <rkolde@gmail.com>, Juri Reimand <jyri.reimand@ut.ee>

References

J. Reimand, M. Kull, H. Peterson, J. Hansen, J. Vilo: g:Profiler – a web-based toolset for functional profiling of gene lists from large-scale experiments (2007) NAR 35 W193-W200

Examples

```
gorth(c("Klf4", "Pax5", "Sox2", "Nanog"), source_organism = "mmusculus", target_organism
```

gprofiler

Annotate gene list functionally.

Description

Annotate gene list functionally.

Usage

```
gprofiler(organism="scerevisiae", query, ordered_query=0, significant=1)
```

7 make_unique_comb

Arguments

gene list origin organism vector of gene names or list of such vectors query ordered_query when output gene lists are ranked one can use this option to get GSEA style

p-values.

logical indicating if all or only statistically significant results should be returned. significant

Details

Wrapper for g:Profiler web toolkit for finding enrichments in gene lists.

Value

Data frame with the Enricment analysis results. If input consisted of several lists the corresponding list is indicated with a variable 'query number'

Author(s)

Juri Reimand < jyri.reimand@ut.ee>

References

J. Reimand, M. Kull, H. Peterson, J. Hansen, J. Vilo: g:Profiler - a web-based toolset for functional profiling of gene lists from large-scale experiments (2007) NAR 35 W193-W200

Examples

```
gprofiler(c("Klf4", "Pax5", "Sox2", "Nanog"), organism = "mmusculus")
```

Function for producing all unique combinations of elements from one make_unique_comb vector.

Description

Function for producing all unique combinations of elements from one vector.

Usage

```
make_unique_comb(1)
```

Arguments

1 vector of some values

Details

The function creates all unique combinations of different values in vector l.

8 plotROC

Value

A matrix with two columns. Rownames are the element names pasted together and separated by "_". The rows are ordered alphabetically according to rownames. The two values in each row are also ordered alphabetically.

Author(s)

Raivo Kolde <rkolde@gmail.com>

Examples

```
make_unique_comb(letters[1:3])
```

plotROC

Plot ROC curve...

Description

Plot ROC curve

Usage

```
plotROC(data, bootstrap=100, colours=NA, text_size=4)
```

Arguments

data input data.frame

colours what colours should the lines be

text_size AUC text size

bootstrap number of bootstrap samples

Details

Input data frame has to contain two columns Probability and Class. Probability refers to some score accordind to which the data is ordered and Class referes to if the observation was positive or negative. If several lines are needed, additional column algorithm should be provided. The ROC curves and AUC-s are calculated by functions calcCurve and calcAUC. By default the AUC values get also confidence intervals, which are calculated using bootstrap. The number of bootstrap samples taken can be set with parameter bootstrap. If desired one can also enter replicate experiments to calculate confidence intervals. For that the dataset has to contain also Column Replicate.

Value

Decription of return value

Author(s)

Raivo Kolde <rkolde@gmail.com>

theme_bw_raivo 9

Examples

```
data = data.frame(Probability = runif(300))
data$Class = rbinom(300, 1, prob = 1 - data$Probability)
#plotROC(data)

data$Algorithm = sample(c("A", "B", "C"), 300, replace = TRUE)
#plotROC(data)

data$Replicate = factor(sample(1:5, nrow(data), replace = TRUE))
#plotROC(data)

data$Replicate = factor(sample(1:5, nrow(data), replace = TRUE))
#plotROC(data)
```

theme_bw_raivo

Theme_bw without ugly whitespace on x axis...

Description

Theme_bw without ugly whitespace on x axis

Usage

```
theme_bw_raivo(base_size=12, base_family="")
```

Arguments

```
base_size base font size
base_family base font family
```

Details

Theme_bw without ugly whitespace on x axis

Author(s)

Raivo Kolde <rkolde@gmail.com>

Index

```
bubbleplot, 2
calcAUC, 2, 8
calcCurve, 2, 3, 8
data, 3
gconvert, 4
generate_gcocoa_query, 5
gorth, 5
gprofiler, 6
make_unique_comb, 7
plotROC, 8
RUtil-package, 1
theme_bw_raivo, 9
```