## Package 'aveytoolkit'

October 28, 2014

#### Version 0.1

**Description** Provides functionality for simple functions like Pause(), and resetPar() as well as complex functions like sigHeatmap, PlotTimeCourse,ComputeTimeLag, ggSmartBoxplot, etc.

## **Depends** R (>= 3.0.2)

License Stefan Avey wrote most of these functions

## LazyData true

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 ${\tt Average Replicates}$ 

AverageReplicates

## Description

This function averages replicates in a matrix or data.frame

## Usage

```
AverageReplicates(eSubSet, numRep)
```

## **Arguments**

eSubSet a matrix or data.frame of values with samples as columns

numRep the number of replicates

#### Value

a data.frame of averaged values with column names coming from the first of each of the replicates with .avg appended

#### Note

Assumes that the replicates are all next to each other

#### Author(s)

Stefan Avey

#### **Examples**

```
mat <- matrix(rnorm(1000), ncol=10) ## 10 columns of random uniform numbers
avgMat <- AverageReplicates(mat, numRep=2) ## average adjacent pairs of columns</pre>
```

aveytoolkit

aveytoolkit.

## Description

aveytoolkit.

browseIndex 3

rowseIndex browseIndex

## Description

Simple function to open HTML page of index in the default browser

#### Usage

```
browseIndex(package, lib.loc = NULL)
```

#### **Arguments**

package a string with the name of package to use

1ib.1oc a string of the directory name of the R library, or <e2><80><98>NULL<e2><80><99>.

The default value of <e2><80><98>NULL<e2><80><99> corresponds to all li-

braries currently known.

#### **Details**

Only works for a single package. Could improve to list an HTML page with multiple packages that you could then choose from. Not currently implemented. Borrows the conecpts from utils::browseVignettes

#### Author(s)

Stefan Avey

#### **Examples**

```
browseIndex("utils")
```

cbind.fill resetPar

## Description

Simple function to combine multiple objects by column while filling in NAs into extra rows created from differing lengths

#### Usage

```
cbind.fill(...)
```

#### **Arguments**

... the objects, that will be converted to a list, to bind column-wise

#### Value

the cbind'ed objects passed in

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#### Author(s)

Dimitris Rizopoulos and Tyler Rinker

#### References

```
http://stackoverflow.com/questions/7962267/cbind-a-df-with-an-empty-df-cbind-fill
```

#### See Also

cbind

#### **Examples**

```
x<-matrix(1:10,5,2)
y<-matrix(1:16, 4,4)
z<-matrix(1:12, 2,6)

cbind.fill(x,y)
cbind.fill(x,y,z)
cbind.fill(mtcars, mtcars[1:10,])</pre>
```

collapseDataset

collapseDataset

#### **Description**

Collapses a dataset from probes to gene symbols.

ulated probes

#### Usage

```
collapseDataset(exprsVals, platform = NULL, mapVector = NULL, oper = max,
    prefer = c("none", "up", "down"), singleProbeset = FALSE,
    returnProbes = FALSE, deProbes = NULL)
```

#### **Arguments**

exprsVals	a matrix or data.frame of expression values with rownames denoting the probes.
platform	the microarray platform the data comes from for extracting the gene symbols
mapVector	a named character vector with names specififying the current identifiers (probes matching the rownames of exprsVals) and the values of the vector specifying the gene symbols (or other identifier to collapse to).
oper	the operation used to choose which probe when multiple probes map to the same gene. Default is max which will calculate the maximum of the average.
prefer	one of "none", "up", or "down", can be abbreviated.
singleProbeset	If TRUE, the operation applies to the average of each sample. Otherwise, if FALSE, the operation applies to the probesets over all samples and only one probeset will be selected. Default is FALSE.
returnProbes	if TRUE, a list of the collapsed expression matrix and the probes are both returned (see return).
deProbes	a list with named vectors "up" and "down" giving the names of up and downreg-

fishersMethod 5

#### **Details**

If singleProbeset is set to TRUE, untested and not recommended, the values for each sample will be taken from the maximum across any probe that maps to that gene. This means that a gene's expression values may be a composition of values from different probes rather than a single probe. if prefer is "up", when multiple deProbes match the same gene, the upregulated will be chosen. Similary for "down". Default is "none" and the probe with the oper will be chosen.

#### Value

If returnProbes is TRUE, a list containing the collapsed dataset in \$exprsVals and the probes chosen in \$probeSets. Otherwise, if returnProbes is FALSE, only the expression matrix is returned.

#### Author(s)

Christopher Bolen, Modified by Stefan Avey

#### **Examples**

## ??

fishersMethod

fishersMethod

## Description

This function combines multiple p-values according to Fisher's Method

## Usage

```
fishersMethod(x)
```

#### Value

a single combined p-value

#### Author(s)

Mike Love

#### References

http://mikelove.wordpress.com/2012/03/12/combining-p-values-fishers-method-sum-of-p-values-bind

```
x \leftarrow c(runif(1000, 0, 1), runif(100, .1, .2))
fishersMethod(x)
```

FoldChange

FoldChange	FoldChange		
------------	------------	--	--

## Description

Calculate the fold change between pairs of conditions in a matrix or data frame

## Usage

FoldChange(x, condNum, condDen, conditions, grouping, log2Transform = FALSE)

## Arguments

X	matrix or data.frame from which to calculate fold changes with samples in columns.
condNum	a vector of $condition(s)$ to be used as the numerator in the fold change calculation
condDen	a vector of condition(s) to be used as the denominator in the fold change calculation
conditions	a vector with length equal to the number of columns of x containing the condition labels between which to find the fold changes.
grouping	a vector with length equal to the number of columns of x containing a grouping of the samples (e.g. subjects, cell lines, strains).
log2Transform	when 'TRUE', log2 transformation will be applied to x before taking the FC. If 'FALSE' (default) no transformation is applied and x is ASSUMED to be already log transformed.

## **Details**

FoldChange takes the fold change of log2 Transformed data by subtracting columns of the x dataframe or matrix depending on the conditions passed in.

#### Value

a data.frame of the fold changes with one column for each fold change

## Author(s)

Stefan Avey

getLoginDetails 7

getLoginDetails

getLoginDetails

#### **Description**

Uses teltk to display a prompt for a loginID and password

#### Usage

```
getLoginDetails()
```

#### **Details**

This function displays a window for a user to enter a loginID and password without showing the password.

#### Value

an invisible named vector of loginID and password

#### Author(s)

Markus Gesmann, Barry Rowlingson

#### References

```
http://www.r-bloggers.com/simple-user-interface-in-r-to-get-login-details/http://r.789695.n4.nabble.com/tkentry-that-exits-after-RETURN-tt854721.html#none
```

## See Also

tcltk

## Examples

```
credentials <- getLoginDetails()
## Do what needs to be done with loginID and password
rm(credentials) # Delete credentials</pre>
```

ggSmartBoxplot

ggSmartBoxplot

## **Description**

Boxplot wrapper for ggplot

#### Usage

```
ggSmartBoxplot(x, mat, splitRowBy = NA, splitColBy = NA, colorBy = NULL,
rows, cols = NA, whichCols = NA, sep = ".", outlier.shape = 17,
outlier.color = NULL, fileName = NA, ...)
```

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#### **Arguments**

the variable to group by for boxplots Χ data.frame or matrix of values to plot with samples in columns mat splitRowBy a factor used to split the data by row in facet\_grid a factor used to split the data by col in facet\_grid splitColBy colorBy a factor used for coloring. No coloring will be done if NULL (default) rows row names or row indices of the items to be plotted substring to search for with "grep" in column names to be plotted cols the column indices or full column names whichCols a separator used in searching for cols in the column names sep outlier.shape shape of outliers (default is 17, filled triangle) outlier.color color of outliers (default is NULL) fileName other arguments that are passed to qplot filename the name of a file to write a PDF to or NA to plot in standard graphics device.

#### Value

invisibly returns a named list of the data frame(s) used for plotting the boxplot(s). The names come from converting the rows argument to a character vector.

#### Author(s)

Stefan Avey

#### See Also

```
ggplot2, qplot
```

```
data(OrchardSprays)
## Example of functionality
ggSmartBoxplot(x=OrchardSprays$treatment,
               mat=t(OrchardSprays[,1]),
              rows=1, whichCols=1:ncol(t(OrchardSprays)),
              colorBy=factor(OrchardSprays$rowpos+OrchardSprays$colpos > 9),
              xlab="Treatment")
## NOT RUN:
cellType <- "PBMC"
geneSub <- grep("HLA-A29.1", rownames(expr))</pre>
age <- "Young"
ages <- c("Young", "Old")</pre>
responses <- c("NR", "R")
subset <- targetFClist[[cellType]]$Age %in% ages &</pre>
  targetFClist[[cellType]]$Response %in% responses
ggSmartBoxplot(x=targetFClist[[cellType]][subset, "Time"],
               mat=exprFClist[[cellType]], ylim=c(-1,1),
               rows=geneSub, whichCols=which(subset),
```

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```
colorBy=targetFClist[[cellType]][subset,"Response"],
splitRowBy=targetFClist[[cellType]][subset,"Age"],
xlab="Days (Post Vaccination)",
fileName=NA)
```

MakeDF

MakeDF

## Description

Creates a data frame from a list. Useful for when the list elements have unequal lengths and as.data.frame fails.

#### Usage

```
MakeDF(list, names)
```

#### **Arguments**

list the list to convert names the names of the list

#### Value

a data frame of the converted list.

## Author(s)

Josh O'Brien

#### References

http://stackoverflow.com/questions/15753091/convert-mixed-length-named-list-to-data-frame and the state of the convert-mixed of the state of the s

#### See Also

gsub

10 Pause

 ${\tt makeTransparent}$ 

makeTransparent

#### **Description**

Simple function to make some colors transparent

#### Usage

```
makeTransparent(..., alpha = 0.5)
```

#### **Arguments**

```
... vector or list of colors
```

alpha transparency factor in range [0,1]

#### Value

a vector of new colors made transparent

#### Author(s)

Ricardo Oliveros-Ramos

#### References

http://stackoverflow.com/questions/8047668/transparent-equivalent-of-given-color

#### See Also

```
rgb, col2rgb
```

## **Examples**

```
makeTransparent("red", "blue")
##[1] "#FF00007F" "#0000FF7F"
makeTransparent("red", "blue", alpha=0.8)
## [1] "#FF0000CC" "#0000FFCC"
```

Pause

Pause

## Description

This function prompts for return key and waits until the return is pushed to continue execution. It is used often to view plots coded in a loop one at a time allowing the user to control when the next plot should be displayed

## Usage

```
Pause(str = "continue", quiet = FALSE)
```

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#### **Arguments**

str optional string to display. Defaults to "continue". quiet if TRUE, no prompt is displayed. Default is FALSE

#### **Details**

The Pause function uses readline to wait until a newline character (produced by the Enter key) is given. Instead of pressing Enter, a newline character can be used to automate this waiting time.

#### Value

NULL is returned by invisible

#### Author(s)

Stefan Avey

#### See Also

```
readline, invisible
```

## **Examples**

```
for(p in 1:10) {
  plot(-10:10, (-10:10)^p, type=b)
  Pause(paste0(see plot of x^,p+1))
}
```

PlotTimeCourse

**PlotTimeCourse** 

#### **Description**

Plot helper function for PlotPCATimeCourse

## Usage

```
PlotTimeCourse(x, y, colors, groups, sampleNames, pch = 19, plotTitle = "",
  legend.loc = "topleft", plotType = c("times", "points"), alpha = 0.15,
  cex.pt = 1, cex.time = 2, time.adj = c(-0.3, -0.3), arrLen = 0.1,
  lwd = 3, numRep = 3, plotFont = NULL, ctrl = TRUE, hourMarks = TRUE,
  legend.cex = 2, ...)
```

#### **Arguments**

```
    x x-values for plotting
    y y-values for plotting
    colors named vector specifying colors for each sample
    groups the virus strain names for the conditions of interest
```

sampleNames names of the samples

12 PlotTimeCourse

pch the plotting character. Default is 19 (a closed circle).

plotTitle a string used for the plotting title

legend.loc location of the legend. Default is topleft.

plotType one of "times" or "points". See Details

alpha transparency factor passed to the alpha function (scales library)

cex.pt size of points. Default is 1

cex.time size of time labels. Default is 2

time.adj the ammount to adjust the time labels. Default is c(-.3, -.3) which moves them

to the lower left

arrLen length of the arrows plotted at the average of each time point. Default is 0.1

lwd line width. Default is 3

numRep the number of replicates. Default is 3

plotFont which font to use for plotting text

ctrl should the control time points be included?

hourMarks should the 4 and 8 hour time points be marked on the plot?

legend.cex size expansion for the legened. Default is 2.

... other arguments passed to heatmap.2

## **Details**

If plotType is "times", ??? Also used to plot 2 genes expression against each other over time. If legend.loc is "none", no legend is plotted. ctrl flag indicates whether or not first numRep values in x and y are from a control measurement

## Value

Nothing is returned

#### Note

Colors are assumed to have as the names attribute some part of the sampleName which can uniquely identify it.

#### Author(s)

Stefan Avey

PrepareExpression 13

|--|--|--|

#### **Description**

Takes in an expression matrix or data frame and prepares it for further analysis

#### Usage

```
PrepareExpression(eset, target, returnProbes = TRUE, labelColumn = "Label",
   select = colnames(target), collapse = ".")
```

#### **Arguments**

eset expression information and (potentially) other columns

target target file where the column names of eset can be matched to 'Label'

returnProbes whether probe mapping should be returned along with expression values in a

list. This will only be returned correctly if there is a column of eset matching

SYMBOL in any case.

labelColumn the column name in the target file to use for matching the column names of eset.

Default is "Label"

select the column names of target to select and merge as the new column names of eset

collapse

#### Details

Wrote this to automate the few lines I always perform to "prepare" an expression set for further processing. I always want to remove the symbols column, rename the column names based on the target file, and (usually) change the rownames to be gene symbols. This function takes in the matrix format that I use to store processed expression files (in a pacakge or file).

#### Value

if returnProbes is FALSE: a list of the prepared expression data frame (exprDat) and the (potentially modified) target data frame (target) . if returnProbes is TRUE (default): a list of three elements including the two above and probeMap (a vector mapping from gene symbols to probe names).

#### Author(s)

Stefan Avey

```
## Creating fake expression matrix
dat <- matrix(rnorm(1000, mean=8, sd=1), nrow=100, ncol=10)
colnames(dat) <- sample(letters[1:10], size=10)
fakeGenes <- as.vector(outer(LETTERS[1:26], LETTERS[1:26], paste0))
x <- data.frame(symbol=fakeGenes[1:nrow(dat)], dat, row.names=paste0("Probe_", 1:nrow(dat)))
head(x) # look at first 6 rows of toy data set
target <- data.frame(Label=letters[1:26], Class=rep(1:3, length.out=26))
head(target)</pre>
```

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**ProcessNames** 

ProcessNames

#### **Description**

Cleans up strings to make them pretty names by removing punctuation, whitespace, and specified substrings

#### Usage

```
ProcessNames(strs, stringsToRm = NULL, rmPunct = TRUE, sep = "_")
```

#### **Arguments**

strs vector or strings to process

stringsToRm a vector or list of strings to search for and remove from strs

rmPunct should punctuation be removed? Default is TRUE.

sep character to replace whitespace

## **Details**

stringsToRm are replaced by "in the order they are given using gsub. After this, punctuation is removed if rmPunct is TRUE. Then, leading and/or trailing whitespace will be removed and the sep will be used to separate words. This function is useful when reading in other people's data and you want to change the row or column names to legal R names or just shorten the names.

## Value

a vector of modified strings from strs

#### Author(s)

Stefan Avey

#### See Also

gsub

RepeatBefore 15

#### **Examples**

```
badNames <- c("Whos Birthday?", "[Date]", "gift Received")
## Remove the string "Whos", remove punctuation, and separate words by _
goodNames <- ProcessNames(badNames, stringsToRm="Whos", rmPunct=TRUE, sep=_)
goodNames
## Remove the string "Whos", dont remove punctuation, and put no separation between words
goodNames <- ProcessNames(badNames, stringsToRm="Whos", rmPunct=FALSE, sep=)
goodNames</pre>
```

RepeatBefore

RepeatBefore

#### **Description**

Replaces NAs with the latest non-NA value

#### Usage

RepeatBefore(x)

## **Arguments**

Х

a vector of values

#### **Details**

NA values will be replaced by the most recent value with a lower index. If there is no non-NA value before the NA appears, it will remain NA.

#### Value

a vector of values

## Author(s)

Ruben

## References

http://stackoverflow.com/questions/7735647/replacing-nas-with-latest-non-na-value

## See Also

rep

```
x = c(NA,NA,a,NA,NA,NA,NA,NA,NA,b,c,d,NA,NA,NA,NA,NA,e)
newX <- RepeatBefore(x)
show(newX)</pre>
```

16 resetPar

resetPar

resetPar

## Description

Simple function to reset plotting parameters for when things get wonky

## Usage

```
resetPar()
```

#### **Details**

This function resets the graphical parameters from the par function. It flashes a new device on the screen but works to reset parameters. Meant to be used when things get hairy and not coded in scripts

## Value

an invisible named list of parameters returned by calling par

## Author(s)

Gavin Simpson

## References

http://stackoverflow.com/questions/5789982/reset-par-to-the-default-values-at-startup

#### See Also

par

```
par(oma=c(4,10,2,1))
plot(1,1)
## paramter settings werent saved so do a reset
resetPar()
plot(1,1)
```

runLimma 17

#### **Description**

This function performs a basic LIMMA analysis on the given expression set

#### Usage

```
runLimma(eset, labels, contrasts, covariates = NULL,
  filterReplicateGenes = TRUE, min.fold.change = 1, min.intensity = 4,
  p.cutoff = 0.05, fitOnly = FALSE)
```

#### **Arguments**

eset the expression matrix

labels the labels for each column of the eset

contrasts Vector of contrasts to make

block Vector of factors specifying a blocking variable (i.e. for paired samples or for ).

NOT IMPLEMENTED!

covariates data frame of covariates (of same length as labels) to include in the model. Use

this if there are paired samples, etc.

filterReplicateGenes

Only include one probeset for each gene (determined by symbol)

min.fold.change

Minimum log2 fold change to be differentially expressed. Default is 1.

min.intensity Minimum log2 intensity (at any time) to be differentially expressed. Default is

4.

p.cutoff FDR corrected cutoff for significant differential expression. Default is 0.05.

fitOnly If true, will return fit2, rather than the matrix of significant genes. Default is

FALSE.

#### **Details**

Generally, an expression matrix is made up of rows of genes (or any other features) and columns of samples. The matrix has data for multiple classes (which are denoted with the 'labels' parameter) and the classes are compared using the vector of contrasts.

#### Value

depends on fitOnly

#### Author(s)

Christopher Bolen

#### See Also

limma

18 sigHeatmap

#### **Examples**

```
## Example:
## If you have a m X 10 matrix eset, with 5 samples of class A and 5 of class B,
## you could compare class A to class B using the following code:
##
## results = runLimma(eset, c(A,A,A,A,B,B,B,B,B), "B-A")
##
## This will return to you a matrix with columns for each comparison and rows for each gene.
## The value in each cells will either be -1, 0, or 1, depending on whether the gene is
## significantly higher in B, not significant, or significantly higher in A, respectively.
## If you want information on p-values and fold changes, set "fitOnly=T", and you can access
## the fit object to get the information.
##
## For other comparisons, you can look at the LIMMA user guide.
```

sigHeatmap

sigHeatmap

#### **Description**

Draw heatmap with significance indicated on boxes

#### Usage

```
sigHeatmap(hm, pvals, pvalDisplayName = "P-value", cutoff = 0.05,
    showOnly = c("both", "positive", "negative", "all"), main = "",
    mainNewlines = 0, sigChar = "*", Rowv = T, hclustMethod = "ward.D",
    ...)
```

#### **Arguments**

hm a matrix of values used for drawing the heatmap

pvals a list or data frame of (possibly FDR corrected but this is not handled by the

function) positive p-values

pvalDisplayName

is printed on the heatmap as a legend. Default is "P-value" but might want to

change to "Q-value", "FDR", etc.

cutoff is threshold for significance of pvals. Default is 0.05

showOnly one of "both", "positive", "negative", or "all" can be abbreviated.

main a string giving the plot main title. Default is "" (i.e. no title is plotted).

mainNewlines a non-negative integer specifying the number of newline characters to plot be-

fore the main title. Used to make the title appear lower on the page. Default is

0

sigChar the character used for plotting on top of significant boxes Rowv should the rows be reordered, passed into heatmap.2

hclustMethod passed to the function stats::hclust. The agglomeration method to be used. This

should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median"

(= WPGMC) or "centroid" (= UPGMC). Default is "ward.D".

... other arguments passed to heatmap.2

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#### **Details**

Only rows with at least one significant column are plotted. If showOnly is "both", plots both positive and negative significant changes. If showOnly is "positive" or "negative", plots only rows of hm with significant positive or negative values respectively. If showOnly is "all", all rows of hm are shown.

#### Value

a vector indicating which of the rows of hm were determined to be significant and subsequently plotted

#### Author(s)

Stefan Avey

#### **Examples**

```
data(mtcars)
x <- as.matrix(mtcars)</pre>
alpha <- 10^-7 # significance threshold
## Caculate whether difference from mean is significant
## This is not done correctly but just to have some sort of significance
diffMean <- mtcars-matrix(colMeans(mtcars),</pre>
                           ncol=ncol(mtcars), nrow=nrow(mtcars), byrow=TRUE)
stdErr <- matrix(sapply(mtcars, sd)/sqrt(nrow(mtcars)),</pre>
                ncol=ncol(mtcars), nrow=nrow(mtcars), byrow=TRUE)
tstats <- diffMean/stdErr
pvals <- pt(as.matrix(tstats), nrow(mtcars)-2, lower=FALSE)</pre>
op <- par(oma=c(4,0,0,20))
sel <- sigHeatmap(x, pvals=pvals, cutoff=alpha, showOnly="b",</pre>
                  main="mtcars Example Heatmap", sigChar="*", notecol=black,
                  notecex=2, Colv=T, Rowv=T, dendrogram="row", trace="none")
par(op)
## Which cars werent selected
rownames(mtcars)[setdiff(1:nrow(mtcars), sel)]
```

VennDiagram

sigHeatmap

## **Description**

Draw a venn diagram of 2 or 3 sets

## Usage

```
VennDiagram(setList, mar = c(0, 0, 1, 0), \ldots)
```

## **Arguments**

setList

a (named) list of the sets to be plotted. The names will be used on the plot. If the list is unnamed, the default names in vennDiagram

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## **Details**

Wrapper around the limma vennDiagram function to make it simpler.

## Value

a data frame of binary values indicating membership in each set with rownames giving the set entries.

## Author(s)

Stefan Avey

#### References

 $Code\ modified\ from\ http://research.stowers-institute.org/mcm/venn.R$ 

## See Also

vennDiagram

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