# Package 'trqwe'

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allDups

All duplicates.

# Description

Finds all duplicates in a vector including first instances of duplicates.

# Usage

allDups(vec)

# Arguments

vec

A vector.

# Value

A boolean vector of the same length as vec. TRUE if the element is duplicated, FALSE if not.

```
allDups(sample(1:100, size=100, replace=T)
```

append<-

append<-

Append to a vector.

# Description

Appends the 2nd argument to the 1st.

# Usage

```
append(x) \leftarrow value
```

# Arguments

Х

A vector.

value

The element to append.

# **Examples**

```
x <- 1:5
append(x) <- 6
print(x)
[1] 1 2 3 4 5 6</pre>
```

bioc

Install Bioconductor package.

# Description

Utility function for installing packages from bioconductor easily.

#### Usage

```
bioc(package)
```

# Arguments

package

unquoted package name.

```
\code{bioc(DESeq2)}
```

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chop

Cleans leading and trailing whitespace.

### **Description**

Removes leading and trailing whitespace in a vector of strings.

# Usage

```
chop(x)
```

# Arguments

Х

A character vector.

#### Value

A vector with leading and trailing whitespace removed.

# **Examples**

```
chop(c(" hello ", " 123 \t"))
[1] "hello" "123"
```

cindex

Concordance Index.

# Description

Calculates the concordance index from the results of a censored survival model. Very fast compared to other packages.

#### Usage

```
cindex(probs, time, event)
```

# Arguments

probs The prognostic score of each patient.

time The time of each patient.

event The death of each patient: 1 for a patient death, 0 for censored.

#### Value

The concordance index.

cosineDist 5

cosineDist

Cosine distance.

#### **Description**

Calculates the cosine distance of rows of a matrix.

# Usage

```
cosineDist(x)
```

#### **Arguments**

Χ

A matrix.

#### Value

Cosine distance as a dist object.

#### See Also

http://stackoverflow.com/questions/2535234/find-cosine-similarity-in-r

# **Examples**

f1score

F1 score.

### Description

Calculates F1 score from the results of a classification model.

#### Usage

```
f1score(probs, class)
```

#### **Arguments**

probs A numeric vector where 1 is predicted positive and 0 is predicted negative.

class A numeric vector where 1 is positive and 0 is negative.

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#### Value

The F1 score.

#### See Also

https://en.wikipedia.org/wiki/F1\_score

fastAUC

Fast AUC

#### **Description**

This function calculates the Area Under the Reciever-Operator Curve from the results of a classifcation model.

# Usage

```
fastAUC(probs, class)
```

#### **Arguments**

probs A numeric vector of probabilities or likelihoods for each data point.

class A numeric vector where 1 is positive and 0 is negative.

#### Value

The AUC.

#### See Also

 $Reference\ https://stat.ethz.ch/pipermail/r-help/2005-September/079872.html$ 

```
library(AppliedPredictiveModeling)
data(abalone)
library(glmnet)
class <- factor(as.numeric(abalone$Type == "M"))
data <- data.matrix(abalone[,-1])
fit <- cv.glmnet(data, class, family="binomial")
probs <- predict(fit, newx=data)[,1]
fastAUC(probs, class)</pre>
```

fastPR 7

#### **Description**

Calculates the points in a Precision-Recall from the results of a classification model.

#### Usage

```
fastPR(probs, class)
```

#### **Arguments**

probs A numeric vector of probabilities or likelihoods for each data point.

class A numeric vector where 1 is positive and 0 is negative.

#### Value

a list containing the ROC curve.

#### **Examples**

```
library(AppliedPredictiveModeling)
data(abalone)
library(glmnet)
class <- factor(as.numeric(abalone$Type == "M"))
data <- data.matrix(abalone[,-1])
fit <- cv.glmnet(data, class, family="binomial")
probs <- predict(fit, newx=data)[,1]
with(fastPR(probs, class), plot(recall, precision, type="l"))</pre>
```

fastReadLines Fast readLines.

#### **Description**

Replacement for readLines, faster.

#### Usage

```
fastReadLines(fname, newlinechar = "\n", nchars = NULL)
```

# Arguments

fname Filename to read.

newlinechar The new line character in the file.

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#### Value

A vector containing all the lines in the file.

#### **Examples**

fastROC

Fast ROC

#### **Description**

Calculates the points in a Reciever-Operator Curve from the results of a classification model.

#### Usage

```
fastROC(probs, class)
```

#### **Arguments**

probs A numeric vector of probabilities or likelihoods for each data point.

class A numeric vector where 1 is positive and 0 is negative.

#### Value

a list containing the ROC curve.

```
library(AppliedPredictiveModeling)
data(abalone)
library(glmnet)
class <- factor(as.numeric(abalone$Type == "M"))
data <- data.matrix(abalone[,-1])
fit <- cv.glmnet(data, class, family="binomial")
probs <- predict(fit, newx=data)[,1]
with(fastROC(probs, class), plot(fpr, tpr, type="l"))</pre>
```

fpkmFromCounts 9

|--|

#### **Description**

Calculates FPKM from a count matrix.

#### Usage

```
fpkmFromCounts(mat, gene_lengths, uq_norm = T)
```

#### **Arguments**

mat An integer matrix representing the counts of a RNA-Seq dataset.

gene\_lengths A named vector with the length of each gene.

uq\_norm If TRUE, will perform upper-quartile normalization.

#### Value

A matrix containing FPKM with the same dimensions as mat.

#### See Also

FPKM-UQ - A normalized read count in which gene expression values, in FPKM, are divided by the 75th percentile value.

https://gdc-docs.nci.nih.gov/Data/Bioinformatics\_Pipelines/Expression\_mRNA\_Pipeline/http://vinaykmittal.blogspot.com/2013/10/fpkmrpkm-normalization-caveat-and-upper.html

rix.
------

#### **Description**

Shortcut function for previewing a matrix or data.frame by displaying the upper-left corner. Similar to head.

#### Usage

```
head2(x, n = 10, ncols = 10)
```

# Arguments

x A wide matrix or data.frame.

n Number of lines to display. Default 10.ncols Number of columns to display. Default 10.

logit

# Value

n by ncols subset of the matrix taken from the upper-left corner.

install

Install CRAN package.

# Description

Utility function for installing packages from CRAN easily.

### Usage

```
install(package, repos = "http://cran.us.r-project.org")
```

# **Arguments**

package

unquoted package name.

# **Examples**

```
\code{install(Rcpp)}
```

logit

Logit Transformation.

# **Description**

Calculates the results of the Logit Transformation.

# Usage

logit(x)

### **Arguments**

Х

Input to the function (e.g., probabilities from 0 to 1).

#### Value

Logit(x)

matrixFactor 11

matrixFactor

Matrix Factor Design.

#### **Description**

From a factor, returns a design matrix with a column for each level.

#### Usage

```
matrixFactor(x, names = NULL)
```

#### **Arguments**

x A factor.

names The name of each instance in the resulting matrix (i.e., the rownames).

#### Value

The design matrix.

#### **Examples**

```
matrixFactor(factor(letters))
```

mccscore

Matthew's correlation coefficient

# Description

Calculates Matthew's correlation coefficient. Requires the Rmpfr package.

#### Usage

```
mccscore(probs, class)
```

#### **Arguments**

probs A numeric vector where 1 is predicted positive and 0 is predicted negative.

class A numeric vector where 1 is positive and 0 is negative.

#### Value

The MCC score.

#### See Also

https://en.wikipedia.org/wiki/Matthews\_correlation\_coefficient

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mcreadRDS

Multi-threaded readRDS

#### **Description**

Uses the pigz utility to improve loading large R objects. This is compatible with saveRDS and readRDS functions. Requires pigz (sudo apt-get install pigz on Ubuntu).

#### Usage

```
mcreadRDS(file, mc.cores = min(parallel::detectCores(), 4))
```

#### **Arguments**

file The filename of the rds object.

mc.cores How many cores to use in pigz. The program does not seem to benefit after more

than about 4 cores.

#### Value

The R object.

#### See Also

```
http://stackoverflow.com/questions/28927750/
```

#### **Examples**

```
x <- sample(1e4, 1e7, replace=T)
saveRDS(x, file="temp.Rds")
xmc <- mcreadRDS("temp.Rds")</pre>
```

mcsaveRDS

Multi-threaded saveRDS

#### **Description**

Uses the pigz utility to improve saving large R objects. This is compatible with saveRDS and readRDS functions. Requires pigz (sudo apt-get install pigz on Ubuntu).

#### Usage

```
mcsaveRDS(object, file, mc.cores = min(parallel::detectCores(), 4))
```

mcsplitapply 13

#### **Arguments**

object An r object to save.

file The filename to save to.

mc.cores How many cores to use in pigz. The program does not seem to benefit after more

than about 4 cores.

#### See Also

http://stackoverflow.com/questions/28927750/

#### **Examples**

mcsplitapply

Parallel split-matrix loop.

#### Description

Splits a matrix into subsets based on a factor, and applies a function to each subset. Typical use case: sum exon count data to gene count data.

#### Usage

```
mcsplitapply(mat, f, func, mc.cores = 4, .combine = rbind, ...)
```

#### **Arguments**

mat The matrix.

f A factor of length equal to nrow(mat). The levels of this factor will split the

matrix into subsets.

func The function to apply to each subset.

mc.cores The number of cores to use.

. combine The function to combine the results with. Default is rbind. Use NA to return a

list.

#### Value

A list or a combined object depending on the .combine parameter.

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#### **Examples**

```
library(pasilla)
library(DEXSeq)
library(trqwe)
data(pasillaDEXSeqDataSet)
exon_counts <- counts(dxd)
f <- rowData(dxd)$groupID
gene_counts <- mcsplitapply(exon_counts, f, colSums)</pre>
```

mgrepl

Multiple grepl.

# Description

Takes in a list of regex patterns and returns true if any pattern matches.

#### Usage

```
mgrepl(patterns, x, ...)
```

#### **Arguments**

x A vector of strings to search.

pattern A vector of regex patterns.

#### Value

A vector of the same length as x, TRUE if any pattern matches.

```
x \leftarrow fastReadLines("http://textfiles.com/ufo/ufobooks.ufo", newlinechar="\r\n", 1e5)
head(x[mgrepl(c("ALIENS", "UFO"), x)])
[1] "
                            UFOLOGY BOOKS (REVISION 2.1 343 books)"
[2] "
         H. S. Stewart on the subject of UFO's. The list is alphabetic"
[3] "
         Tom Mickus's most excellent board UFONET I. (416-237-1204)"
[4] "
         Bill Adler * LETTERS TO THE AIR FORCE ON UFOS 1967"
[5] "
         Gordon W. Allen
                                     OVERLORDS OLYMPIANS AND THE UFO
[6] "
         Robert B. Beard
                                     FLYING SAUCERS, UFO'S AND EXTRA"
```

nelson\_aelen\_surv 15

nelson\_aelen\_surv

Nelson Aelen estimator

#### **Description**

Nelson<e2><80><93>Aalen estimator is an estimator of the cumulative hazard function in survival data. It can be used to compare the overall risks of two groups or used to estimate the number of deaths before a certain time.

#### Usage

```
nelson_aelen_surv(time, event)
```

#### **Arguments**

time The time of each patient.

event The death of each patient: 1 for a patient death, 0 for censored.

#### Value

A list containing the cumulative hazard function.

#### See Also

#https://en.wikipedia.org/wiki/Nelson-Aalen\_estimator

#### **Examples**

```
library(survival)
data(veteran)
with(nelson_aelen_surv(veteran$time, veteran$status), plot(ti, Hi, type="b"))
```

posteriorBalance

Posterior probability adjustment.

#### **Description**

Adjusts the posterior probability of a classifier based on unbalanced datasets. In classification model where the negative data is randomly under-sampled and all the positive data is used, the adjustment factor (beta) is p(s=1|-) - i.e., the probability that a negative datapoint is selected in the classifier.

#### Usage

```
posteriorBalance(probs, beta)
```

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#### **Arguments**

probs The original posterior probability.

beta The adjustment factor.

#### Value

The adjusted posterior probability.

#### See Also

Dal Pozzolo, Andrea, et al. "Calibrating probability with undersampling for unbalanced classification." Computational Intelligence, 2015 IEEE Symposium Series on. IEEE, 2015.

prepend<-

Prepend to a vector.

# Description

Prepends the 2nd argument to the 1st.

# Usage

```
prepend(x) \leftarrow value
```

#### **Arguments**

x A vector.

value The element to append.

```
x <- 1:5
prepend(x) <- 6
print(x)
[1] 6 1 2 3 4 5</pre>
```

reload 17

reload

Reload a package.

# Description

Unload and reload a package and sets the namespace search order.

# Usage

```
reload(package, pos = 2)
```

# Arguments

package pos Unquoted package name.

Namespace search position.

# **Examples**

\code{reload(trqwe)}

reloadtrqwe

Unload and reload trawe.

# Description

Unload and reload trqwe. Shortcut for reload(trqwe)

# Usage

reloadtrqwe()

se

Standard error.

# Description

Calculates the standard error of a sampling distribution.

# Usage

se(x)

# Arguments

Χ

A vector.

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#### Value

The standard error of x.

#### **Examples**

```
x <- rnorm(1e3)
se(x)
[1] 0.03192027</pre>
```

sigmoid

Sigmoid Function.

# Description

Calculates the results of the sigmoid function.

#### Usage

```
sigmoid(x)
```

#### **Arguments**

probs

x Input to the function.

#### Value

Sigmoid(x)

statsCallback

Variable information.

# Description

Automatically stores basic information of variables in the previous command. This function adds a callback which reports information on previous variables and stores this information in .stats.

# Usage

```
statsCallback()
```

```
statsCallback()
my_data <- VADeaths
   .stats
> [1] "dim: 5 4, length: 20, class: matrix, typeof: double"
```

tablec 19

tablec

*Fast C++ tabulation.* 

## **Description**

Takes in a character, integer or factor vector and tabulates the number of times each element appears.

#### Usage

```
tablec(x, sort = F)
```

#### **Arguments**

x A character, integer or factor vector. NAs are allowed.

sort TRUE if the result names should be sorted alphanumerically.

#### Value

A integer vector of counts of each element.

```
x <- factor(sample(1e5, 1e8, replace=T))</pre>
microbenchmark(table(x), tablec(x), times=3)
Unit: milliseconds
     expr
                            lq
                                             median
               min
                                                                      max
                                     mean
                                                            uq
 table(x) 9777.6457 10479.0949 10717.3382 11180.544 11187.1844 11193.8246
tablec(x) 678.0364
                      685.9467 713.1181
                                            693.857
                                                      730.6589
                                                                 767.4608
x <- sample(letters, 1e8, replace=T)</pre>
microbenchmark(table(x), tablec(x), tablec(x,sort=T), times=3)
Unit: seconds
                         min
                                   lq
                                                 median
               expr
                                          mean
                                                              uq
           table(x) 5.778514 5.829125 5.855560 5.879737 5.894083 5.908430
          tablec(x) 1.589360 1.589381 1.589516 1.589402 1.589594 1.589786
tablec(x, sort = T) 1.589386 1.590520 1.591824 1.591655 1.593044 1.594432
```

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tail2

Lower-left corner of matrix.

#### **Description**

Shortcut function for previewing the bottom of a matrix or data.frame by displaying the lower-left corner. Similar to tail.

#### Usage

```
tail2(x, n = 10, ncols = 10)
```

#### **Arguments**

x A wide matrix or data.frame.

n Number of lines to display. Default 10.ncols Number of columns to display. Default 10.

#### Value

n by ncols subset of the matrix taken from the lower-left corner.

TCGA\_barcode

Parse TCGA barcode.

#### **Description**

Taking in a full TCGA sample barcode, or any subset of the barcode, and return extracted values.

#### Usage

```
TCGA_barcode(x, what = "patient")
```

#### Arguments

x TCGA barcode, or a vector of barcodes.

what Which information to return.

#### Value

The specified information contained in the barcode of the same length as x.

```
TCGA_barcode(c("TCGA-02-0001-01C-01D-0182-01", "TCGA-02-0001-11C-01D-0182-01"), what="tissue")
[1] "01" "11"
```

timePrompt 21

# Description

Reports time in seconds to the R prompt of the previous command. This function adds a callback which saves the running of individual commands and reports the time in seconds on the next line.

#### Usage

```
timePrompt()
```

#### **Examples**

```
> timePrompt()
0.000s> x <- sample(1:10, size=1e8, replace=T)
1.240s>
```

topn

Highest elements in a vector.

#### **Description**

Finds the top elements in a vector very quickly. Equivalent ot -sort(-x, partial=1:n)

#### Usage

```
topn(x, n = 100, value = F, lowest = F)
```

# Arguments

x A numeric vector.

n The number of top elements to return.

value If TRUE, returns the values of the top elements. If FALSE, returns the indices.

lowest If TRUE, returns the lowest elements instead of the highest.

#### Value

A vector containing the indices or the values of the top elements.

#### See Also

http://stackoverflow.com/questions/18450778/

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#### **Examples**

varSizes

Size of R objects.

#### **Description**

Prints out the size of all R objects in the environment.

#### Usage

```
varSizes(env = globalenv(), units = "KB")
```

#### **Arguments**

env The environment to search (default global environment).

units Units to print out for each variable.

#### Value

A data.frame containing the size of each object.

ww\_test

Wald-Wolfowitz Runs Tests for Randomness

#### **Description**

This is the k-category asymptotic Z Test with continuity correction. Imagine rolling a die multiple times to obtain a sequence of rolls. This statistic tests whether there exists a "run" within the sequence where one particular number comes up more times in a row than expected randomly. If the test is significant, it can be concluded that the die rolls are not independent.

#### Usage

```
ww_test(x)
```

ww\_test 23

# Arguments

x A vector of items, coerced into a factor.

#### Value

A p-value for the statistical test.

#### See Also

Reference https://ncss-wpengine.netdna-ssl.com/wp-content/themes/ncss/pdf/Procedures/NCSS/Analysis\_of\_Runs.pdf

```
set.seed(1)
ww_test(sample(2, 100, replace=T))
ww_test(c(sample(6, 90, replace=T), rep(1,10)))
```

# **Index**

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