Package ‘wrMisc’

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Title Analyze Experimental High-Throughput (Omics) Data

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Description The efficient treatment and convenient analysis of experimental high-throughput (omics) data gets facilitated through this collection of diverse functions. Several functions address advanced object-conversions, like manipulating lists of lists or lists of arrays, reorganizing lists to arrays or into separate vectors, merging of multiple entries, etc.

Another set of functions provides speed-optimized calculation of standard deviation (sd), coefficient of variance (CV) or standard error of the mean (SEM) for data in matrixes or means per line with respect to additional grouping (eg n groups of replicates).

Other functions facilitate dealing with non-redundant information, by indexing unique, adding counters to redundant or eliminating lines with respect redundancy in a given reference-column, etc.

Help is provided to identify very closely matching numeric values to generate (partial) distance matrixes for very big data in a memory efficient manner or to reduce the complexity of large data-sets by combining very close values.

Many times large experimental datasets need some additional filtering, adequate functions are provided.

Batch reading (or writing) of sets of files and combining data to arrays is supported, too.

Convenient data normalization is supported in various different modes, parameter estimation via permutations or boot-strap as well as flexible testing of multiple pair-wise combinations using the framework of ‘limma’ is provided, too.

VignetteBuilder knitr

Depends R (>= 3.1.0)

Imports grDevices, graphics, MASS, stats, utils

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addLetterWoLast

Description

This function allows to add ‘addChr’ to all entries, without the last entry

Usage

addLetterWoLast(x, addChr)

Arguments

x (character) main input
addChr (character)
Value

This function returns a modified character vector

See Also

`paste`; used in `cutAtMultSites`

Examples

```r
.addLetterWoLast(c("abc","efgh"),"Z")
```

---

.allRatioMatr1to2

*Calculate ratios for each column to each column of reference-matrix*

Description

This function calculates ratio(s) for each column of matrix `x` versus all/each column(s) of matrix `y` (reference)

Usage

```r
.allRatioMatr1to2(x, y, asLog2 = TRUE, sumMeth = "mean", callFrom = NULL)
```

Arguments

- `x` (matrix or data.frame) main input1
- `y` (matrix or data.frame) main input2
- `asLog2` (logical)
- `sumMeth` (character) method
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a numeric vector or matrix in dimension of `x` (so far summarize all ratios from mult division from mult ref cols as mean or median)

See Also

`makeMAList`, `grep`

Examples

```r
.allRatioMatr1to2(matrix(11:14, ncol=2), matrix(21:24, ncol=2))
```
Description

This function extracts/cuts text-fragments out of txt following specific anchors defined by arguments cutFrom and cutTo.

Usage

`.allRatios(dat, ty = "log2", colNaSep = ")

Arguments

dat (matrix or data.frame) main input
ty (character) type of ratio (eg 'log2')
colNaSep (character) separator

Value

This function returns a numeric vector

See Also

`makeMAList, grep`

Examples

`.allRatios(matrix(11:14, ncol=2))`

Description

This function allows summarizing along columns of multiple arrays in list
Usage

.arrLstMean(
    arrLst,
    sumType = "mean",
    arrOutp = FALSE,
    signifDig = 3,
    formatCheck = FALSE,
    silent = FALSE,
    debug = FALSE,
    callFrom = NULL
)

Arguments

arrLst (list) main input
sumType (character)
arrOutp (logical)
signifDig (integer)
formatCheck (logical)
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

array (1st dim will be summary along cols, rows will be layers of 3rd array-dim

See Also

used in cutArrayInCluLike

Examples

.datSlope(c(3:6))

---

Summary of function .arrLstSEM

Description

This function allows summarizing along columns of mult arrays in list
Usage

.asDF2

Arguments

arrLst (list) main input
arrOutp (logical)
signifDig (integer)
formatCheck (logical)
silent (logical) suppress messages
depug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

array (1st dim will be summary along cols, rows will be layers of 3rd array-dim ie dim(arrLst[[1]])[3])

See Also

used in cutArrayInCluLike

Examples

.datSlope(c(3:6))

.asDF2 Convert anything to data.frame

Description

This function allows converting anything to data.frame

Usage

.asDF2(z)
Arguments

z  (numeric vector, factor, matrix or list) main input

Value
data.frame

See Also

as.data.frame

Examples

.asDF2(c(3:6))

---

.breakInSer  Get series of values after last discontinuity

Description

This function aims to get series of values after last discontinuity

Usage

.breakInSer(x, getFrom = "last")

Arguments

x  (numeric) main input

getFrom  (character)

Value

This function returns a numeric vector of reduced length

See Also

dist

Examples

.breakInSer(c(11:14,16:18))
.bringToCtr  

**Bring most extreme to center**

**Description**
This function aims to bring most extreme value to center

**Usage**
```r
.bringToCtr(aa, ctr, ctrFa = 0.75)
```

**Arguments**
- **aa** *(numeric)* main input
- **ctr** *(numeric)* 'control'
- **ctrFa** *(numeric <1)* modulate amplitude of effect

**Value**
This function returns an adjusted numeric vector

**See Also**
dist

**Examples**
```r
.bringToCtr(11:14, 9)
```

---

.checkArgNa  

**Check argument names**

**Description**
This function allows checking of argument names

**Usage**
```r
.checkArgNa(x, argNa, lazyEval = TRUE)
```

**Arguments**
- **x** *(character)* main input
- **argNa** *(character)* argument name
- **lazyEval** *(logical)* decide if argument should be evaluated with abbreviated names, too
This function returns a elongated character vector

See Also

chartr

Examples

.checkArgNa("Abc",c("ab","Ab","BCD"))

Description

This function allows to check list of arrays for consistent dimensions of all arrays

Usage

.checkedConsistentArrList(
  arrLst,
  arrNDim = 3,
  fxName = NULL,
  varName = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

arrLst (list) main input
arrNDim (integer) number of dimensions for arrays
fxName (character) this name will be given in message
varName (character)
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

list
### Description

This function allows converting 'dat' (may be list, data.frame etc) to simple vector, more elaborate than unlist()

### Usage

```
.checkConvt2Vect(dat, toNumeric = TRUE)
```

### Arguments

- `dat`  
  (list, data.frame) main input

- `toNumeric`  
  (logical)

### Value

character (or numeric) vector

### See Also

`unlist`; used in `equLenNumber`

### Examples

```r
aa <- matrix(1:14, ncol=2)
.checkConvt2Vect(aa)
```
Description

This function was designed to check a factor object

Usage

```r
.checkFactor(
  fac,
  facNa = NULL,
  minLev = 2,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `fac` (factor) main input
- `facNa` (character) level-names
- `minLev` (integer) minimum number of levels
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a corrected/adjusted factor

See Also

`factor`

Examples

```r
.checkFactor(gl(3,2))
```
Description

checkFileNameExtensions Function for checking file-names.

Usage

.checkFileNameExtensions(fileNa, ext)

Arguments

fileNa (character) file name to be checked
ext (character) file extension

Value

modified character vector

Examples

.checkFileNameExtensions("testFile.txt","txt")

Description

This function allows checking an argument for Location of legend, if value provided not found as valid, it returns 'defLoc

Usage

.checkLegendLoc(
  legLoc,
  defLoc = "topright",
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
.checkLmConfInt

Arguments

- legLoc (character) main input
- defLoc (character)
- silent (logical) suppress messages
- debug (logical) additional messages for debugging
- callFrom (character) allow easier tracking of messages produced

Value

This function returns a character vector designing the potential location of legend

See Also

- legend

Examples

  .checkLegendLoc("abc")

________________________________________________________________________________

.checkLmConfInt  Compare 'dat' to confidence interval of linear model 'lMod' (eg from lm())

________________________________________________________________________________

Description

This function allows to compare 'dat' to confidence interval of linear model 'lMod' (eg from lm())

Usage

  .checkLmConfInt(dat, lMod, level = 0.95)

Arguments

- dat matrix or data.frame, main input
- lMod linear model, only used to extract coefficients offset & slope
- level (numeric) alpha threshold for linear model

Value

This function returns a logical vector for each value in 2nd col of 'dat' if INSIDE confid interval

See Also

- searchLinesAtGivenSlope

Examples

  set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
.checkRegrArguments  Check regression arguments

Description
This function allows to check arguments for linear regression. Used as argument checking for regrBy1or2point and regrMultBy1or2point

Usage
.checkRegrArguments(inData, refList, regreTo, callFrom = NULL)

Arguments
- inData (numeric vector) main input
- refList (list)
- regreTo (numeric vector)
- callFrom (character) allow easier tracking of messages produced

Value
list

See Also
append; lrbind

Examples
.datSlope(c(3:6))

.chooseGrpCol  Automatic choice of colors

Description
This function allows to do automatic choice of colors: if single-> grey, if few -> RColorBrewer, if many : gradient green -> grey/red

Usage
.chooseGrpCol(nGrp, paired = FALSE, alph = 0.2)
The function `combineListAnnot` allows to combine information (annotation) from list of matrices (i.e., replace when NA), using always the columns specified in `useCol` (numeric).

**Usage**

```r
.combineListAnnot(
  lst,
  useCol = 1:2,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `lst` (list): main input
- `useCol` (numeric vector): which columns should be used
- `silent` (logical): suppress messages
- `debug` (logical): additional messages for debugging
- `callFrom` (character): allow easier tracking of messages produced

**Value**

This function returns a single matrix of combined (non-redundant) info.
.compareByDiff

See Also

used in cutArrayInCluLike

Examples

.datSlope(c(3:6))

---

.compareByDiff  Compare by distance/difference

Description

This function allows to compare by distance/difference

Usage

.compareByDiff(dat, limit, distVal = FALSE)

Arguments

dat  list of 2 numerical vectors
limit (numeric, length=1) threshold value for retaining values, used with distance-type specified in argument 'compTy'
distVal (logical) to toggle output as matrix of numeric (distance values above 'limit', others NA) or matrix of logical

Value

This function returns a list with close matches of 'x' to given 'y', the numeric value depends on 'sortMatch' (if FALSE then always value of 'y' otherwise of longest of x&y)

See Also

findCloseMatch, checkSimValueInSer, and also .compareByLogRatio, for convenient output countCloseToLimits

Examples

c <- list(aa=11:14, bb=c(13.1,11.5,14.3,20:21))
.compareByLogRatio  
\textit{Compare by log-ratio}

\textbf{Description}

This function allows to compare by log-ratio

\textbf{Usage}

\texttt{.compareByLogRatio(dat, limit, distVal = FALSE)}

\textbf{Arguments}

\begin{itemize}
\item \texttt{dat} \hspace{1cm} list of 2 numerical vectors
\item \texttt{limit} \hspace{1cm} (numeric, length=1) threshold value for retaining values, used with distance-type specified in argument ‘compTy’
\item \texttt{distVal} \hspace{1cm} (logical) to toggle output as matrix of numeric (distance values above 'limit', others NA) or matrix of logical
\end{itemize}

\textbf{Value}

This function returns a list with close matches of 'x' to given 'y', the numeric value depends on 'sortMatch' (if FALSE then always value of 'y' otherwise of longest of x&y)

\textbf{See Also}

\texttt{findCloseMatch, checkSimValueInSer}, and also \texttt{.compareByDiff, for convient output countCloseToLimits}

\textbf{Examples}

\begin{verbatim}
cc <- list(aa=11:14, bb=c(13.1,11.5,14.3,20:21))
.compareByLogRatio(cc, 1)
\end{verbatim}

--

.\texttt{compareByPPM}  
\textit{Compare by PPM}

\textbf{Description}

This function allows to compare by ppm

\textbf{Usage}

\texttt{.compareByPPM(dat, limit, distVal = FALSE)}
.complCols

Arguments

dat | list of 2 numerical vectors
limit | (numeric, length=1) threshold value for retaining values, used with distance-type specified in argument 'compTy'
distVal | (logical) to toggle output as matrix of numeric (distance values above 'limit', others NA) or matrix of logical

Value

This function returns a list with close matches of 'x' to given 'y', the numeric value depends on 'sortMatch' (if FALSE then always value of 'y' otherwise of longest of x&y)

See Also

findCloseMatch, checkSimValueInSer, and also .compareByDiff, for convenient output countCloseToLimits

Examples

cc <- list(aa=11:14, bb=c(13.1,11.5,14.3,20:21))
.complCols(.compareByPPM(cc, 1))

__Description__

This function was designed to complete the selection of columns of sparse matrix 'dat' with sets of 'nCombin' columns at complete 'coverage'. Context: In sparse matrix 'dat' search subsets of columns with some rows as complete (no NA).

__Usage__

.complCols(x, dat, nCombin)

__Arguments__

x | (integer, length=1) column number for with other columns to combine & give (some) complete non-NA lines are sought
dat | (matrix) .. init data, smay be parse matrix with numerous NA
nCombin | (integer) .. number of columns used to make complete subset

__Value__

This function returns a matrix of column-indexes complementing (nCombin rows)
See Also

rowNormalize

Examples

.composeCallName(3, dat=matrix(c(NA, 12:17, NA, 19), ncol=3), nCombin=3)

---

.composeCallName  Compose sequence of (function-)calls

Description

This function was designed for tracing the hierarchy of function-calls. It allows to remove any tailing space or ': ' from `callFrom` (character vector) and return with added 'newNa' (+ 'add2Tail')

Usage

.composeCallName(newNa, add2Head = "", add2Tail = " : ", callFrom = NULL)

Arguments

- `newNa`  (character vector) main input
- `add2Head`  (character)
- `add2Tail`  (character)
- `callFrom`  (character) may also contain multiple separate names (ie length >1), will be concatenated using '->'

Value

character vector (history of who called whom)

See Also

paste

Examples

.composeCallName("newFunction", callFrom="initFunction")
### .convertMatrToNum

*Convert numeric matrix to numeric*

**Description**

Take matrix and return vector

**Usage**

```r
.convertMatrToNum(matr, useCol = NULL)
```

**Arguments**

- `matr` (matrix) main input
- `useCol` (integer) design the columns to be used

**Value**

numeric vector

**See Also**

`matrix`

**Examples**

```r
.convertMatrToNum(matrix(1:6, ncol=2))
```

---

### .convertNa

*Convert/standardize names of 'query' to standard names from 'ref'*

**Description**

This function converts/standardizes names of 'query' to standard names from 'ref' (list of possible names (char vect) where names define standardized name). It takes 'query' as character vector and return character vector (same length as 'query') with 'converted/corrected' names

**Usage**

```r
.convertNa(query, ref, partMatch = TRUE)
```

**Arguments**

- `query` (matrix or data.frame, min 2 columns) main input
- `ref` (list) list of multiple possible names associated to given group, reference name for each group is name of list
- `partMatch` (logical) allows partial matching (ie name of 'ref' must be in head of 'query')
Value

This function returns a character vector

Examples

daPa <- matrix(c(1:5, 8, 2:6, 9), ncol=2)

d.corDuplItemsByIncrem(letters[1:6], letters[8:4])
.cutAtSearch

Search character-string and cut either before or after

Description

This function extracts/cuts text-fragments out of `txt` following specific anchors defined by arguments `cutFrom` and `cutTo`.

Usage

```r
.cutAtSearch(
  x,
  searchChar,
  after = TRUE,
  silent = TRUE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `x` character vector to be treated
- `searchChar` (character) text to look for
- `after` (logical)
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a modified character vector

See Also

grep

Examples

```r
.cutAtSearch("abcdefg","de")
```
.cutStr  
Cut string to get all variants from given start with min and max length

Description
This function allows truncating character vector to all variants from given start, with min and optional max length. Used to evaluate argument calls without giving full length of argument.

Usage
.cutStr(txt, startFr = 1, minLe = 1, maxLe = NULL, reverse = TRUE)

Arguments

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>txt</td>
<td>(character) main input, may be length &gt;1</td>
</tr>
<tr>
<td>startFr</td>
<td>(interger) where to start</td>
</tr>
<tr>
<td>minLe</td>
<td>(interger) minimum length of output</td>
</tr>
<tr>
<td>maxLe</td>
<td>(interger) maximum length of output</td>
</tr>
<tr>
<td>reverse</td>
<td>(logical) return longest text-fragments at beginning of vector</td>
</tr>
</tbody>
</table>

Value
This function returns a character vector.

See Also
used in pasteC; substr

Examples
.cutStr("abcdefg", minLe=2)

.datSlope  
Model linear regression and optional plot

Description
This function allows to model a linear regression and optionally to plot the results.
Usage

`.datSlope(
    dat,
    typeOfPlot = "sort",
    toNinX = FALSE,
    plotData = FALSE,
    silent = FALSE,
    debug = FALSE,
    callFrom = NULL
)
`}

Arguments

dat (vector or matrix) main input
typeOfPlot (character)
toNinX (logical)
plotData (logical)
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

numeric vector with intercept and slope, optional plot

See Also

`append; lrbind`

Examples

`.datSlope(c(3:6))`

Description

This function allows extracting NA-neighbour value

Usage

`.extrNAneighb(x, grp)`
Arguments

x    initial matrix to treat
grp  (factor) grouping of replicates

Value

numeric vector

See Also

unique, nonAmbiguousNum, faster than firstOfRepeated which gives more detail in output (lines/elements/indexes of omitted)

Examples

extrNAneighb(c(11:14,NA), rep(1,5))
.extrNumHeadingSepChar

Extract numbers before separator followed by alphabetic character

Description

This function aims to extract number(s) before separator followed by alphabetic character (return named numeric vector, NAs when no numeric part found)

Usage

.extrNumHeadingSepChar(x, sep = "_")

Arguments

x character vector to be treated
sep (character) separator

Value

This function returns a numeric vector

See Also

nchar

Examples

.extrNumHeadingSepChar(" 1B ")

.filterNetw

Filter nodes & edges for extracting networks (main) This function allows extracting and filtering network-data based on fixed threshold (limInt) and add sandwich-nodes (nodes inter-connecting initial nodes) out of node-based queries.

Description

Filter nodes & edges for extracting networks (main)

This function allows extracting and filtering network-data based on fixed threshold (limInt) and add sandwich-nodes (nodes inter-connecting initial nodes) out of node-based queries.
Usage

\[ \text{.filterSw(} \]
\[ \quad \text{lst,} \]
\[ \quad \text{remOrphans = TRUE,} \]
\[ \quad \text{reverseCheck = TRUE,} \]
\[ \quad \text{filtCol = 2,} \]
\[ \quad \text{callFrom = NULL,} \]
\[ \quad \text{silent = FALSE,} \]
\[ \quad \text{debug = FALSE} \]
\[ \text{)} \]

Arguments

\- **lst** (list, composed of multiple matrix or data.frames) main input (each list-element should have same number of columns)
\- **remOrphans** (logical) remove networks consisting only of 2 connected edges
\- **reverseCheck** (logical)
\- **filtCol** (integer, length=1) which column of lst should be used to filter using thresholds limInt and sandwLim
\- **callFrom** (character) allow easier tracking of message(s) produced
\- **silent** (logical) suppress messages
\- **debug** (logical) display additional messages for debugging

Value

This function returns a matrix or data.frame

See Also

filterNetw and other CRAN package dedicated to networks

Examples

\begin{verbatim}
ab <- 1:10
\end{verbatim}

Description

Filtering of matrix or array x (may be 3-dim array) according to fiTy and checkVa

Usage

\[ \text{.filterSw(x, fiTy, checkVa, indexRet = TRUE)} \]
Arguments

x  array (3-dim) of numeric data
fiTy (character) which type of testing to perform ('eq', 'inf', 'infeq', 'sup', 'supeq', '>', '<', '>=', '<=', '==')
checkVa (logical) s
indexRet (logical) if TRUE (default) rather return index numbers than filtered values

Value

This function returns either index (position within 'x') or concrete (filtered) result

See Also

filt3dimArr; filterList; filterLiColDeList;

Examples

arr1 <- array(11:34, dim=c(4,3,2), dimnames=list(c(LETTERS[1:4]), paste("col",1:3,sep=""),c("ch1","ch2")))
filt3dimArr(arr1, displCrit=c("col1","col2"), filtCrit="col2", filtVal=7)
.filterSw(arr1, fiTy="inf", checkVa=7)

Description

This function aims to filter for size

Usage

.filtSize(x, minSize = 5, maxSize = 36)

Arguments

x  main input
minSize (integer) minimum number of characters, if NULL set to 0
maxSize (integer) maximum number of characters

Value

list of filtered input

See Also

filtSizeUniq; correctToUnique, unique, duplicated
Examples

aa <- 1:10

Description

This function aims to find overlap instances among range of values in lines of 'x' (typically give just min & max)

Usage

.findBorderOverlaps(x, rmRedund = FALSE, callFrom = NULL)

Arguments

x (matrix of numeric values or all-numeric data.frame) main input
rmRedund (logical) report overlaps only in 1st instance (will show up twice otherwise)
callFrom (character) allow easier tracking of message(s) produced

Value

This function returns a matrix with line for each overlap found, cols 'refLi' (line no), 'targLi' (line no), 'targCol' (col no)

See Also

nchar

Examples

aa <- 11:15
.firstMin Get first minimum

Description
This function allows to find the first minimum of a numeric vector

Usage
.firstMin(x, positionOnly = FALSE)

Arguments
x (numeric vector) main input
definition (logical)

definition (logical)

Value
numeric vector

See Also
which.min

Examples
.firstMin(c(4,3:6))

.fuse2ArrBy2ndDim fuse 2 instances of 3dim arr as mult cols in 3dim array

Description
This function allows fusing 2 instances of 3dim arr as mult cols in 3dim array (ie fuse along 2nd
dim, increase cols)

Usage
.fuse2ArrBy2ndDim(arr1, arr2, silent = FALSE, debug = FALSE, callFrom = NULL)

Arguments
arr1 (array)
arr2 (array)
silent (logical) suppress messages
definition (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced
Value

This function returns a numeric vector with number of non-numeric characters (i.e., not '.' or 0-9))

See Also

array

Examples

aa <- 11:15

---

.getAmean Get A value for each group of replicates

Description

This function calculates the 'A' value (i.e., group mean) for each group of replicates (e.g., for MA-plot).

Usage

.getAmean(dat, grp)

Arguments

dat (matrix or data.frame) main input
grp (factor) grouping of replicates

Value

This function returns a numeric vector

See Also

makeMAList

Examples

.getAmean(matrix(11:18, ncol=4), gl(2,2))
.getAmean2  
*Get A value for each group of replicates based on comp*

**Description**

This function calculates the 'A' value (ie group mean) for each group of replicates (eg for MA-plot) comp is matrix telling which groups to use/compare, assuming that dat are already group-means

**Usage**

```
.getAmean2(dat, comp)
```

**Arguments**

- `dat` (matrix or data.frame) main input
- `comp` (matrix) tells which groups to use/compare, assuming that dat are already group-means

**Value**

This function returns a numeric vector

**See Also**

`makeMAList`

**Examples**

```
.getAmean(matrix(11:18, ncol=4), gl(2,2))
```

---

.getMvalue2  
*Get M value for each group of replicates based on comp*

**Description**

This function calculates the 'M' value (ie log-ratio) for each group of replicates based on comp (eg for MA-plot) comp is matrix telling which groups to use/compare, assuming that dat are already group-means

**Usage**

```
.getMvalue2(dat, comp)
```

**Examples**

```
.getMvalue2(matrix(11:18, ncol=4), gl(2,2))
```
.growTree

Arguments

- **dat** (matrix or data.frame) main input
- **comp** (matrix) tells which groups to use/compare, assuming that dat are already group-means

Value

This function returns a numeric vector

See Also

makeMAList

Examples

\texttt{.getAmean(matrix(11:18, ncol=4), gl(2,2))}

---

.growTree  \hspace{1cm} \textit{Grow tree}

Description

This function allows growing tree-like structures (data.tree objects)

Usage

\texttt{.growTree(tm, setX, addToObj = NULL)}

Arguments

- **tm** (list) main input, $\text{disDat}$ .. matrix with integer start & end sites for fragments; $\text{lo}$ (logical) which fragments may be grown; $\text{start}$ (integer) index for which line of $\text{disDat}$ to start; $\text{it}$ numeric version of $\text{lo}$; $\text{preN}$ for previous tree objects towards root; $\text{iter}$ for iterator (starting at 1)
- **setX** .. data.tree object (main obj from root)
- **addToObj** .. data.tree object (branch on which to add new branches/nodes)

Value

list

See Also

buildTree

Examples

\texttt{.datSlope(c(3:6))}
Segment (1-dim vector) 'dat' into clusters

This function allows segmenting (1-dim vector) 'dat' into clusters. If 'automClu=TRUE ..' first try automatic clustering, if too few clusters, run km with length(dat)^0.3 clusters. This function requires the package NbClust to be installed.

Usage

```r
.insp1dimByClustering(
  dat,
  automClu = TRUE,
  cluChar = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

dat matrix or data.frame, main input
automClu (logical) run automatic clustering
cluChar (logical) to display cluster characteristics
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This function returns clustering (class index) or (if 'cluChar' = TRUE) list with clustering and cluster-characteristics

See Also

`searchLinesAtGivenSlope`

Examples

```r
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
```
.inspectHeader

Inspect 'matr' and check if 1st line can be used/converted as header

Description

This function inspects 'matr' and check if 1st line can be used/converted as header. If colnames of 'matr' are either NULL or 'V1', etc the 1st row will be tested if it contains any of the elements (if not, 1st line won’t be used as new colnames) If 'numericCheck'=TRUE, all columns will be tested if they can be converted to numeric

Usage

```r
.inspectHeader(
  matr,
  headNames = c("Plate", "Well", "StainA"),
  numericCheck = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `matr` (matrix or data.frame) main input to be instected
- `headNames` (character) column-names t look for
- `numericCheck` (logical) allows reducing complexity by drawing for very long x or y
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a matrix vector or data.frame similar to input

See Also

`head` for looking at first few lines

Examples

```r
ma1 <- matrix(letters[1:6], ncol=3, dimnames=list(NULL,c("ab","Plate","Well"))
.inspectHeader(ma1)
```
Description

This function allows to refine/filter ‘dat1’ (1dim dataset, eg cluster) with aim of keeping center of data. It is done based on most freq class of histogram keep/filter data if 'core' (core = NULL, keepOnly = TRUE, displPlot = FALSE, silent = TRUE, debug = FALSE, callFrom = NULL)

Usage

.keepCenter1d(
  dat1,
  core = NULL,
  keepOnly = TRUE,
  displPlot = FALSE,
  silent = TRUE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dat1       simple numeric vector
core       numeric vector (betw 0 and 1) for fraction of data to keep; if null trimmed-Mean/max hist occurance will be used, limited within 30-70 perent; may also be 'high' or 'low' for forcing low (20-60percent) or high (75-99) percent of data to retain
keepOnly   (logical)
displPlot  (logical) show plot of hist & boundaries
silent     (logical) suppress messages
debug      (logical) additional messages for debugging
callFrom   (character) allow easier tracking of messages produced

Value

This function returns the index of values retained or if 'keepOnly' return list with 'keep' index and 'drop' index

See Also

.searchLinesAtGivenSlope

Examples

set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
.keepFiniteCol  Remove all columns where all data are not finite

Description
This function aims to remove all columns where all data are not finite

Usage
.keepFiniteCol(
  dat,
  msgStart = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments
dat (matrix or data.frame) main input
msgStart (character)
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allows easier tracking of messages produced

Value
This function returns a corrected matrix or data.frame

See Also
renameColumns; is.finite

Examples
ma1 <- matrix(c(1:5, Inf), ncol=2)
.keepFiniteCol(ma1)
**mayBeNum**

*Check if vector may be numeric content*

**Description**

This function allows to checking if a given vector may be numeric content

**Usage**

```r
.maybeNum(x, pattern = NULL)
```

**Arguments**

- **x** (numeric vector) main input
- **pattern** (character) custom pattern to check

**Value**

This function returns a logical/boolean vector for each of the elements of 'x'

**See Also**

numeric; convMatr2df

**Examples**

```r
.maybeNum(c(3:6))
```

---

**medianSpecGrp**

*Rescale respective to specific group*

**Description**

This function allows to rescale data 'x' so that specific group 'grpNum' gets normalized to predefined value 'grpVal'. In normal case x will be multiplied by 'grpVal' and devided by value obtained from 'grpNum'. If summary of 'grpNum-positions' or 'grpVal' is 0, then grpVal will be attained by subtraction of summary & adding grpVal

**Usage**

```r
.medianSpecGrp(x, grpNum, grpVal, sumMeth = "median", callFrom = NULL)
```
**Arguments**

- **x** (numeric vector) main input
- **grpNum** (numeric)
- **grpVal** (numeric)
- **sumMeth** (character) method for summarizing
- **callFrom** (character) allow easier tracking of messages produced

**Value**

numeric vector

**See Also**

- `which.min`

**Examples**

```
.firstMin(c(4,3:6))
```

---

### `.mergeMatrices`

**Merge Multiple Matrices (main)**

**Description**

This function allows merging of multiple matrix-like objects from an initial list.

**Usage**

```
.mergeMatrices(  
  inpL,  
  mode = "intersect",  
  useColumn = 1,  
  extrRowNames = FALSE,  
  na.rm = TRUE,  
  argL = NULL,  
  silent = FALSE,  
  debug = FALSE,  
  callFrom = NULL  
)
```
Arguments

inpL (list containing matrices or data.frames) main input (multiple matrix or data.frame objects)

mode (character) allows choosing restricting to all common elements (mode='intersect') or union (mode='union')

useColumn (integer, character or list) the column(s) to consider, may be 'all' to use all, integer to select specific indexes or list of indexes or colnames for custom-selection per matrix

eextrRowNames (logical) decide whether columns with all values different (ie no replicates or max divergency) should be excluded

na.rm (logical) suppress NAs

argL (list of arguments)

silent (logical) suppress messages

debug (logical) additional messages for debugging

callFrom (character) allow easier tracking of messages produced

Value

This function returns a matrix containing all selected columns of the input matrices to fuse

See Also

mergeMatrixList, merge, mergeMatrices for separate entries

Examples

mat1 <- matrix(11:18, ncol=2, dimnames=list(letters[3:6],LETTERS[1:2]))

.minDif

Description

This function aims to find closest neighbour to numeric vector

Usage

.minDif(z, initOrder = TRUE, rat = TRUE)

Arguments

z (numeric) vector to search minimum difference

initOrder (logical) return matrix so that 'x' matches exactly 2nd col of output

rat (logical) express result as ratio
Description

This function returns distances between sorted points of 2-column matrix `x`.

Usage

`.neigbDis(x, asSum = TRUE)`

Arguments

- **x**: (matrix or data.frame, min 2 columns) main input
- **asSum**: (logical) if `TRUE` (default) the sum of all distances will be returned, otherwise the individual distances

Value

This function returns a numeric vector with distances

Examples

daPa <- matrix(c(1:5,8,2:6,9), ncol=2)
`.neigbDis(daPa)`
.normalize  

Main Normalization function

Description

This function aims to normalize a matrix or data.frame by columns. It assumes all checks have been done before calling this function.

Usage

.normalize(
  dat,
  meth,
  mode,
  param,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dat  matrix or data.frame of data to get normalized
meth  (character) may be "mean", "median", "NULL", "none", "trimMean", "rowNormalize", "slope", "exponent", "slope2Sections", "vsn"; When NULL or 'none' is chosen the input will be returned
mode  (character) may be "proportional", "additive"; decide if normalization factors will be applies as multiplicative (proportional) or additive; for log2-omics data mode="additive" is suggested
param  (list) additional parameters
silent  (logical) suppress messages
debug  (logical) additional messages for debugging
callFrom  (character) allows easier tracking of messages produced

Value

This function returns a numeric vector

See Also

normalizeThis

Examples

aa <- matrix(1:12, ncol=3)
.normalize(aa,"median",mode="proportional",param=NULL)
Normalize columns of 2dim matrix to common linear regression fit within range of `useQuant`

Usage

```r
.normConstSlope(
  mat,
  useQuant = c(0.2, 0.8),
  refLines = NULL,
  diagPlot = TRUE,
  plotLog = "",
  datName = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `mat`: matrix or data.frame of data to get normalized
- `useQuant`: (numeric) quantiles to use
- `refLines`: (NULL or numeric) allows to consider only specific lines of 'dat' when determining normalization factors (all data will be normalized)
- `diagPlot`: (logical) draw diagnostic plot
- `plotLog`: (character) indicate which axis should be displayed on log-scale, may be 'x', 'xy' or 'y'
- `datName`: (character) use as title in diag plot
- `silent`: (logical) suppress messages
- `debug`: (logical) additional messages for debugging
- `callFrom`: (character) allows easier tracking of messages produced

Value

This function returns a numeric vector

See Also

- `normalizeThis`

Examples

```r
aa <- matrix(1:12, ncol=3)
```
# offCenter

Return position of `di` (numeric vector) which is most eccentric (distant to 0), starts with NAs as most eccentric

## Description
This function aims to return position of `di` (numeric vector) which is most eccentric (distant to 0), starts with NAs as most eccentric. It is used for identifying/removing (potential) outliers. Note: this fx doesn't consider reference distributions, even with "perfect data"! Most points will be tagged!

## Usage

```r
.offCenter(di, nMost = 1)
```

## Arguments

- `di` (numeric) main input
- `nMost` (integer)

## Value
This function returns a integer/numeric vector (indicating index)

## See Also
use in `presenceFilt; diff`

## Examples
```
.offCenter(11:14)
```

---

# pasteCols

Paste-concatenate all columns of matrix

## Description
This function allows paste columns

## Usage

```r
.pasteCols(mat, sep = "")
```

## Arguments

- `mat` initial matrix
- `sep` (character) separator
Value

simplified/non-redundant vector/matrix (ie fewer lines for matrix), or respective index

See Also

unique, nonAmbiguousNum, faster than firstOfRepeated which gives more detail in output (lines/elements/indexes of omitted)

Examples

.plotCountPie(matrix(11:16, ncol=2), sep="_")

Description

This function allows to inspect results of table or uniqCountReport on a pie-plot Note: fairly slow for long vectors!!

Usage

.plotCountPie(
  count,
  tit = NULL,
  col = NULL,
  radius = 0.9,
  sizeTo = NULL,
  clockwise = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

count (integer vector) counting result
tit (character) optional title in plot
col (character) custom colors in pie
radius (numeric) radius passed to pie
sizeTo (numeric or character) optional reference group for size-population relative adjusting overall surface of pie
clockwise (logical) argument passed to pie
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced
Value

vector with counts of n (total), nUnique (wo any repeated), nHasRepeated (first of repeated), nRedundant, optional figure

See Also

uniqCountReport, correctToUnique, unique

Examples

.plotCountPie(table(c(1:5,4:2)))

---

.plusLowerCaps  Add lower caps to character vector

Description

This function allows adding all content as lower caps to/of character vector

Usage

.plusLowerCaps(x)

Arguments

x (character) main input

Value

This function returns a elongated character vector

See Also

chartr

Examples

.plusLowerCaps(c("Abc","BCD"))
**.predRes**

*Calculate residues of (2-dim) linear model 'lMod'-prediction of/for 'dat'*

**Description**

This function calculates residues of (2-dim) linear model 'lMod'-prediction of/for 'dat' (using 2nd col of 'useCol') (indexing in 'dat', matrix or data.frame with min 2 cols), using 1st col of 'useCol' as 'x'. It may be used for comparing/identifying data close to regression (e.g., re-finding data on autoregression line in FT-ICR)

**Usage**

```
.predRes(dat, lMod, regTy = "lin", useCol = 1:2)
```

**Arguments**

- `dat` (matrix or data.frame, main input)
- `lMod` (linear model, only used to extract coefficients offset & slope)
- `regTy` (character) type of regression model
- `useCol` (integer) columns to use

**Value**

This function returns a numeric vector of residues (for each line of dat)

**See Also**

*searchLinesAtGivenSlope*

**Examples**

```
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
```

---

**.raiseColLowest**

*Raise all values close to lowest value*

**Description**

This function aims to raise all values close to lowest value to end up as at value of 'raiseTo'. This is done independently for each col of mat. This function sets all data to common raiseTo (which is min among all cols)
Usage

.removeCol(mat, 
raiseTo = NULL, 
minFa = 0.1, 
silent = FALSE, 
debug = FALSE, 
callFrom = NULL 
)

Arguments

mat  (matrix of numeric values) main input
raiseTo  (numeric)
minFa  (numeric) minimum factor
silent  (logical) suppress messages
debug  (logical) display additional messages for debugging
callFrom  (character) allow easier tracking of message(s) produced

Value

This function returns a numeric vector with number of non-numeric characters (i.e. not '.' or 0-9))

See Also

nchar

Examples

aa <- 11:15

removeCol

Remove columns indicated by col-number

Description

This function aims to remove columns indicated by col-number

Usage

.removeCol(matr, rmCol)

Arguments

matr  (matrix or data.frame) main input
rmCol  (integer) column index for removing
This function returns a matrix or data.frame

See Also
dist

Examples

```r
aa <- matrix(1:6, ncol=3)
.removeCol(aa, 2)
```

Description

This function aims to search for (empty) columns containing only entries defined in `searchFields` and remove such columns. If `fromBackOnly = TRUE` only tailing empty columns will be removed (other columns with "empty" entries in middle will be kept). If `""=TRUE` columns containing all NAs will be excluded as well. This function will also remove columns containing (exclusively) mixtures of the various `searchFields`.

Usage

```r
.removeEmptyCol(
  dat,
  fromBackOnly = TRUE,
  searchFields = c("", "", "NA.", NA),
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

dat (matrix or data.frame) main input
fromBackOnly (logical)
searchFields (character)
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allows easier tracking of messages produced

Value

This function returns a corrected matrix or data.frame
.replSpecChar

See Also

renameColumns; is.finite

Examples

ma1 <- matrix(c(1:5, NA), ncol=2)
.removeEmptyCol(ma1)

Examples

.replSpecChar(c("jhjh(ab)","abc"))
.retain1stPart

Trim character string: keep only text before 'sep'

Description
Trim character string: keep only text before 'sep' (length=1 !)

Usage
.retain1stPart(chr, sep = " = ", offSet = 1)

Arguments

<table>
<thead>
<tr>
<th>chr</th>
<th>character vector to be treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>sep</td>
<td>(character) separator</td>
</tr>
<tr>
<td>offSet</td>
<td>(integer) off-set</td>
</tr>
</tbody>
</table>

Value
This function returns a modified character vector

See Also
substr

Examples
.retain1stPart("abc = def")

.rowGrpCV

row group CV (main)

Description
This function calculates CVs for matrix with multiple groups of data, ie one CV for each group of data.

Usage
.rowGrpCV(x, grp, means)

Arguments

| x | numeric matrix where replicates are organized into separate columns |
| grp | (factor) defining which columns should be grouped (considered as replicates) |
| means | (numeric) alternative values instead of means by .rowGrpMeans() |
.rowGrpMeans

Value

This function returns a matrix of CV values

See Also

rowGrpCV, rowCVs, arrayCV, replPlateCV

Examples

```
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
grp1 <- gl(4,3,labels=LETTERS[1:4])[2:11]
head(.rowGrpCV(dat1, grp1, .rowGrpMeans(dat1, grp1)))
```

Description

This function calculates CVs for matrix with multiple groups of data, ie one CV for each group of data.

Usage

`.rowGrpMeans(x, grp, na.replVa = NULL, na.rm = TRUE)`

Arguments

- `x` numeric matrix where replicates are organized into separate columns
- `grp` (factor) defining which columns should be grouped (considered as replicates)
- `na.replVa` (numeric) value to replace NA values
- `na.rm` (logical) remove all NA values

Value

This function returns a matrix of mean values per row and group of replicates

See Also

rowGrpCV, rowCVs, arrayCV, replPlateCV

Examples

```
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
grp1 <- gl(4,3,labels=LETTERS[1:4])[2:11]
head(.rowGrpMeans(dat1, grp1))
```
Description

This function calculates sd for matrix with multiple groups of data, i.e. one sd for each group of data.

Usage

.rowGrpSds(x, grp)

Arguments

x numeric matrix where replicates are organized into separate columns
grp (factor) defining which columns should be grouped (considered as replicates)

Value

This function returns a matrix of sd values per row and group of replicates

See Also

.rowGrpCV, rowCVs, arrayCV, replPlateCV

Examples

set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
grp1 <- gl(4,3,labels=LETTERS[1:4])[2:11]
head(.rowGrpSds(dat1, grp1))

Description

This function calculates row-sums for matrix with multiple groups of data, with multiple groups of data, i.e. one sd for each group of data.

Usage

.rowGrpSums(x, grp, na.replVa = NULL, na.rm = TRUE)
.rowNorm

Arguments

x       numeric matrix where replicates are organized into separate columns
grp     (factor) defining which columns should be grouped (considered as replicates)
na.replVa (numeric) value to replace NA values
na.rm   (logical) remove all NA values

Value

This function returns a matrix of row-sums for matrix with multiple groups of data

See Also

rowGrpCV, rowCVs, arrayCV, replPlateCV

Examples

set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
grp1 <- gl(4,3,labels=LETTERS[1:4])[2:11]
head(.rowGrpSums(dat1, grp1))
Arguments

dat (matrix) .. init data, smay be parse matrix with numerous NA
refLi (NULL or numeric) allows to consider only specific lines of ’dat’ when determining normalization factors (all data will be normalized)
method (character) may be "mean","median" (plus "NULL","none"); When NULL or ’none’ is chosen the input will be returned as is
proportMode (logical) decide if normalization should be done by multiplicative or additive factor
maxFact (numeric, length=2) max normalization factor
fact0val (integer)
retFact (logical)
callFrom (character) This function allows easier tracking of messages produced
debug (logical) additional messages for debugging
silent (logical) suppress messages

Value

This function returns a matrix of normalized data same dimensions as ’dat’

See Also

rowNormalize

Examples

.rowNorm(matrix(11:31, ncol=3), refLi=1, method="mean", proportMode=TRUE)

Description

This function was designed to obtain normalization factors.

Usage

.rowNormFact(
  dat,
  combOfN,
  comUse,
  method = "median",
  refLi = NULL,
  refGrp = NULL,
  proportMode = TRUE,
Arguments

- **dat** (matrix): init data, may be parse matrix with numerous NA
- **combOfN** (matrix): # matrix of index for all sub-groups (assumed as sorted)
- **comUse** (list): index of complete lines for each col of combOfN
- **method** (character): may be "mean","median" (plus "NULL","none"); When NULL or 'none' is chosen the input will be returned as is
- **refLi** (NULL or numeric): allows to consider only specific lines of 'dat' when determining normalization factors (all data will be normalized)
- **refGrp** (integer): only the columns indicated will be used as reference, default all columns (integer or colnames)
- **proportMode** (logical): decide if normalization should be done by multiplicative or additive factor
- **minQuant** (numeric): optional filter to set all values below given value as NA
- **maxFact** (numeric, length=2): max normalization factor
- **omitNonAlignable** (logical): allow omitting all columns which can't get aligned due to sparseness
- **silent** (logical): suppress messages
- **debug** (logical): additional messages for debugging
- **callFrom** (character): This function allows easier tracking of messages produced

Value

This function returns a matrix of column-indexes complementing (nCombin rows)

See Also

- `rowNormalize`

Examples

```r
ma1 <- matrix(11:41, ncol=3)
```
### .scale01

**Scale between 0 and 1 (main)**

**Description**

This function rescales between 0 and 1

**Usage**

`.scale01(x)`

**Arguments**

- `x` numeric vector to be re-scaled

**Value**

This function returns a numeric vector of same length with re-scaled values

**See Also**

`scaleXY`, `scale`

**Examples**

`.scale01(11:15)`

---

### .scaleSpecGrp

**Rescale respective to specific group**

**Description**

This function allows to rescale data 'x' so that 2 specific groups get normalized to predefined values (and all other values follow proportionally) 'grp1Num' and 'grp2Num' should be either numeric for positions in 'x' or character for names of 'x'; if 'grp1Num' and/or 'grp2Num' design multiple locations: perform median or mean summarization, according to 'sumMeth'

**Usage**

`.scaleSpecGrp(  
  x,  
  grp1Num,  
  grp1Val,  
  grp2Num = NULL,  
  grp2Val = NULL,  
  sumMeth = "mean",  
  callFrom = NULL  
)`
Arguments

- **x** (numeric vector) main input
- **grp1Num** (numeric)
- **grp1Val** (numeric)
- **grp2Num** (numeric)
- **grp2Val** (numeric)
- **sumMeth** (character) method for summarizing
- **callFrom** (character) allow easier tracking of messages produced

Value

numeric vector

See Also

- **which.min**

Examples

```
.firstMin(c(4,3:6))
```

### .scaleXY

*Scale between min and max value (main)*

Description

This function rescales between user-defined min and max values

Usage

```
.scaleXY(x, minim = 2, maxim = 3)
```

Arguments

- **x** numeric vector to be re-scaled
- **minim** (numeric) minimum value for resultant vector
- **maxim** (numeric) minimum value for resultant vector

Value

This function returns a matrix of CV values

See Also

- **scaleXY**, **scale**
Examples

.seqCutStr(seqCutStr(11:15, min=1, max=100))

Description

This function is depreciated, please use /cutStr instead! This function allows truncating character vector to all variants from given start, with min and optional max length Used to evaluate argument calls without giving full length of argument

Usage

.seqCutStr(txt, startFr = 1, minLe = 1, reverse = TRUE)

Arguments

- **txt** (character) main input, may be length >1
- **startFr** (integer) where to start
- **minLe** (integer) minimum length of output
- **reverse** (logical) return longest text-fragments at beginning of vector

Value

This function returns a character vector

See Also

pasteC; substr

Examples

.seqCutStr("abcdefg", minLe=2)
### .setLowestTo

*Set lowest value to given value*

#### Description
This function aims to set lowest value of x to value 'setTo'

#### Usage
```
.setLowestTo(x, setTo)
```

#### Arguments
- **x** (numeric) main vector to be treated
- **setTo** (numeric) replacement value

#### Value
This function returns a numeric vector

#### See Also
- nchar

#### Examples
```
.setLowestTo(9:4, 6)
```

---

### .sortMid

*Choose most frequent or middle of sorted vector*

#### Description
This function chooses the (first) most frequent or middle of sorted vector

#### Usage
```
.sortMid(x, retVal = TRUE)
```

#### Arguments
- **x** (numeric) main input
- **retVal** (logical) return value of most frequent, if FALSE return index of (1st) 'x' for most frequent
Value
This function returns a numeric vector.

See Also
simple/partial functionality in `summarizeCols`, `checkSimValueInSer`

Examples
```r
.stackArray(11:14)
```

```
.stackArray
Reorganize array by reducing dimension 'byDim' (similar to `stack() for data-frames)
```

Description
This function aims to reorganize an array by reducing dimension 'byDim' (similar to `stack()` for data-frames). It returns an array/matrix of 1 dimension less than 'arr', 1st dim has more lines (names as paste with '_.')

Usage
```
.stackArray(arr, byDim = 3)
```

Arguments
```
arr (array) main input
byDim (integer)
```

Value
This function returns an array/matrix of 1 dimension less than 'arr', 1st dim has more lines (names as paste with '_.')

See Also
dist

Examples
```
(arr1 <- array(11:37, dim=c(3,3,3)))
.stackArray(arr1, 3)
```
.summarizeCols

Summarize columns of matrix (or data.frame) 'x' using apply (main)

Description

This function summarizes columns of matrix (or data.frame) 'x' using apply. Note, it cannot handle character entries! (all results will be NA)

Usage

.summarizeCols(
  x,
  me = c("median", "medianComplete", "mean", "meanComplete", "aver", "average", "min", "max", "maxOfRef", "minOfRef", "maxAbsOfRef", "lastLi", "last", "firstComplete", "first", "firstLi", "summary"),
  vectAs1row = TRUE
)

Arguments

x  data.frame matrix of data to be summarized by column
me  (character, length=1) summarization method (eg 'maxLast', 'minLast', 'maxLast', 'maxAbsLast', 'minLast', 'medianComplete' or 'meanComplete')
vectAs1row  (logical) if TRUE will interpret non-matrix 'x' as matrix with 1 row (correct effect of automatic conversion when extracting 1 line)

Value

vector with summary for each column

See Also

summarizeCols

Examples

t1 <- matrix(round(runif(30,1,9)), nc=3); rownames(t1) <- letters[c(1:5,3:4,6:4)]
Description

This function allows trimming/removing redundant text-fragments from end.

Usage

`.trimFromEnd(x, ..., callFrom = NULL, debug = FALSE, silent = TRUE)`

Arguments

- `x` character vector to be treated
- `...` more vectors to be treated
- `callFrom` (character) allow easier tracking of messages produced
- `debug` (logical) display additional messages for debugging
- `silent` (logical) suppress messages

Value

This function returns a modified character vector.

See Also

- `trimRedundText`; Inverse : Find/keep common text `keepCommonText`; you may also look for related functions in package `stringr`

Examples

```r
txt1 <- c("abcd_ccc","bcd_ccc","cde_ccc")
.trimFromEnd(txt1)
```

Description

This function allows trimming/removing redundant text-fragments from start.
Usage
.trimFromStart(
  x,
  ..., 
  minNchar = 1,
  silent = TRUE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

x          character vector to be treated
...        more vectors to be treated
minNchar   (integer) minimum number of characters that must remain
silent     (logical) suppress messages
debug      (logical) display additional messages for debugging
callFrom   (character) allow easier tracking of messages produced

Value

This function returns a modified character vector

See Also

.trimRedundText; Inverse : Find/keep common text keepCommonText; you may also look for related functions in package stringr

Examples

txt1 <- c("abcd_ccc","bcd_ccc","cde_ccc")
.trimFromStart(txt1)

.trimLeft
.Trim from Left

Description

This function allows trimming/removing redundant text-fragments from left side

Usage

.trimLeft(x, minNchar = 1, silent = TRUE, debug = FALSE, callFrom = NULL)
Arguments

- **x**: character vector to be treated
- **minNchar**: (integer) minimum number of characters that must remain
- **silent**: (logical) suppress messages
- **debug**: (logical) display additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

Value

This function returns a modified character vector

See Also

- `trimRedundText`: Inverse: Find/keep common text
- `keepCommonText`: you may also look for related functions in package `stringr`

Examples

```r
txt1 <- c("abcd_ccc","bcd_ccc","cde_ccc")
.trimLeft(txt1)
```

Description

This function allows trimming/removing redundant text-fragments from right side

Usage

```r
.trimRight(x, minNchar = 1, silent = TRUE, debug = FALSE, callFrom = NULL)
```

Arguments

- **x**: character vector to be treated
- **minNchar**: (integer) minimum number of characters that must remain
- **silent**: (logical) suppress messages
- **debug**: (logical) display additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

Value

This function returns a modified character vector
See Also

trimRedundText: Inverse : Find/keep common text keepCommonText: you may also look for related functions in package stringr

Examples

txt1 <- c("abcd_ccc","bcd_ccc","cde_ccc")
.trimRight(txt1)

-------------------------------------------------------------------------------
.uniqueWName   Check regression arguments
-------------------------------------------------------------------------------

Description

This function is an enhanced version of unique, names of elements are maintained

Usage

.uniqueWName(
  x,
  splitSameName = TRUE,
  silent = TRUE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

  x        (numeric or character vector) main input
  splitSameName  (logical)
  silent      (logical) suppress messages
  debug       (logical) additional messages for debugging
  callFrom    (character) allow easier tracking of messages produced

Value

  vector like input

See Also

unique

Examples

  aa <- c(a=11, b=12,a=11,d=14, c=11)
  .uniqueWName(aa)
  .uniqueWName(aa[-1]) # value repeated but different name
.vector2Matr  
*Convert numeric vector to matrix*

**Description**

Take (numeric) vector and return matrix, if 'colNa' given will be used as colname

**Usage**

```
.vector2Matr(x, colNa = NULL, rowsKeep = TRUE)
```

**Arguments**

- `x` (numeric or character) main input
- `colNa` (integer) design the column-name to be given
- `rowsKeep` (logical) is TRUE make matrix of 1 column, otherwise of 1 row

**Value**

matrix

**See Also**

`matrix`

**Examples**

```
.vector2Matr(c(3:6))
```

---

addBeforFileExtension  
*Add text before file-extension*

**Description**

This function helps changing character strings like file-names and allows adding the character vector 'add' (length 1) before the extension (defined by last '.') of the input string 'x'. Used for easily creating variants/additional filenames but keeping current extension.

**Usage**

```
addBeforFileExtension(
  x,
  add,
  sep = "_",
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)
```
adjBy2ptReg

Arguments

- **x**: main character vector
- **add**: character vector to be added
- **sep**: (character) separator between 'x' & 'add' (character, length 1)
- **silent**: (logical) suppress messages
- **callFrom**: (character) allow easier tracking of messages produced
- **debug**: (logical) additional messages for debugging

Value

modified character vector

Examples

```r
addBeforFileExtension(c("abd.txt","ghg.ijij.txt","kjh"),"new")
```

---

adjBy2ptReg  Linear rescaling of numeric vector or matrix

Description

adjBy2ptReg takes data within window defined by 'lims' and determines linear transformation so that these points get the regression characteristics 'regrTo', all other points (ie beyond the limits) will follow the same transformation. In other words, this function performs 'linear rescaling', by adjusting (normalizing) the vector 'dat' by linear regression so that points falling in 'lims' (list with upper & lower boundaries) will end up as 'regrTo'.

Usage

```r
adjBy2ptReg(
  dat,
  lims,
  regrTo = c(0.1, 0.9),
  refLines = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- **dat**: numeric vector, matrix or data.frame
- **lims**: (list, length=2) should be list giving limits (list(lo=c(min,max),hi=c(min,max))) in data allowing identifying which points will be used for determining slope & offset
adjustUnitPrefix

regrTo (numeric, length=2) to which characteristics data should be regressed
refLines (NULL or integer) optional subselection of lines of dat (will be used internal as refDat)
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This function returns a matrix (of same dimensions as input matrix) with normalized values

See Also

normalizeThis

Examples

set.seed(2016); dat1 <- round(runif(50,0,100),1)
## extreme values will be further away :
adjBy2ptReg(dat1,lims=list(c(5,9), c(60,90)))
plot(dat1, adjBy2ptReg(dat1, lims=list(c(5,9),c(60,90))))

adjustUnitPrefix  Adjust Value With Different Decimal Prefixes To Single Prefix Plus Unit

Description

This function provides help converting values with with different unit-prefixes to a single prefix-unit type. This can be used to convert a vector of mixed prefixes like 'p' and 'n'. Any text to the right of the unit will be ignored.

Usage

adjustUnitPrefix(
  x,
  pref = c("z", "a", "f", "p", "n", "u", "m"),
  unit = "sec",
  sep = c(".", " ", " "),
  headingTxt = " ",
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
appendNR

Arguments

x  
(character) vector containing digit uunit-prefix and unit terms

pref  
(character) multiplicative unit-prefixes, assumes as increasing factors of 1000

unit  
(character) unit name, the numeric part may be separated by one space-character

sep  
(character) separator characters that may appear between integer numeric value and unit description

headingTxt  
(character) additional text preceding the numeric part of `x` to be ignored/removed

silent  
(logical) suppress messages

debug  
(logical) additional messages for debugging

callFrom  
(character) allow easier tracking of messages produced

Details

Please note that the current version recognizes and converts only integer values, decimals or scientific writing won’t work. The resultant numeric vector expresses all values as lowest prefix unit level. In case of invalid entries NAs will be returned.

Please note the ‘u’ is used for ‘micro’.

Value

This function returns a numeric vector with quantities extracted and adjusted to a single type of unit (without the unit description)

See Also

convToNum

Examples

adjustUnitPrefix(c("10.psec abc","2 fsec etc"), unit="sec")

appendNR  
Append vectors or lists, without duplicating common elements

Description

This function allows combining two vectors or lists without duplicating common content (defined by name of list-elements).

Usage

appendNR(x, y, rmDuplicate = TRUE, silent = FALSE, callFrom = NULL)
Arguments

- **x** (vector or list) must have names to allow checking for duplicate names in `y`
- **y** (vector or list) must have names to allow checking for duplicate names in `x`
- **rmDuplicate** (logical) avoid duplicating list-elements present in both `x` and `y` (based on names of list-elements)
- **silent** (logical) suppress messages
- **callFrom** (character) allow easier tracking of message(s) produced

Details

When setting the argument `rmDuplicate=FALSE` the function will behave like `append`.

Value

If both `x` and `y` are vectors, the output will be a vector, otherwise it will be a list

See Also

- `append`
- `lrbind`

Examples

```r
li1 <- list(a=1, b=2, c=3)
li2 <- list(A=11, B=12, c=3)
appendNR(li1, li2)
append(li1, li2)
```

---

**arrayCV**  

CV of array

Description

*arrayCV* gets CVs for replicates in 2 or 3 dim array and returns CVs as matrix. This function may be used to calculate CVs from replicate microtiter plates (eg 8x12) where replicates are typically done as multiple plates, ie initial matrixes that are the organized into arrays.

Usage

```r
arrayCV(arr, byDim = 3, silent = TRUE, callFrom = NULL)
```

Arguments

- **arr** (3-dim) array of numeric data like where replicates are along one dimension of the array
- **byDim** (integer) over which dimension replicates are found
- **silent** (logical) suppress messages
- **callFrom** (character) allow easier tracking of message produced
asSepList

Value

matrix of CV values

See Also

cvs, rowGrpCV, replPlateCV

Examples

set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
head(arrayCV(dat1,byDim=2))

asSepList

Organize Data as Separate List-Entries

Description

asSepList allows reorganizing most types of input into a list with separate numeric vectors. For example, matrixes or data.frames will be split into separate columns (different to partUnlist which maintains the original structure). This function also works with lists of lists. This function may be helpful for reorganizing data for plots.

Usage

asSepList(
  y,
  minLen = 4,
  asNumeric = TRUE,
  exclElem = NULL,
  sep = "_",
  fillNames = TRUE,
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

y list to be separated/split in vectors
minLen (integer) min length (or number of rows), as add’l element to eliminate arguments given without names when asSepList is called in vioplot2
asNumeric (logical) to transform all list-elements in simple numeric vectors (won’t work if some entries are character)
exclElem (character) optional names to exclude if any (lazy matching) matches (to exclude other arguments be misinterpreted as data)
sep (character) separator when combining name of list-element to colames
fillNames   (logical) add names for list-elements/ series when not given
silent     (logical) suppress messages
callFrom   (character) allow easier tracking of messages produced
debug      (logical) display additional messages for debugging

Value

This function returns a list, partially unlisted to vectors

See Also

partUnlist, unlist

Examples

bb <- list(fa=gl(2,2), c=31:33, L2=matrix(21:28,nc=2),
   li=list(li1=11:14, li2=data.frame(41:44)))
asSepList(bb)
## multi data-frame examples
cb <- data.frame(a=11:15, b=21:25, c=31:35)
cb <- data.frame(a=51:53, b=61:63)
cc <- list(gl(3,2), ca, cb, 91:94, short=81:82, letters[1:5])
asSepList(cc)
cc <- list(gl(3,2), e1=gl(3,2), e2=ca, e3=cb, e4=91:94, short=81:82, e6=letters[1:5])
asSepList(cd)

buildTree

Connect edges to from tree and extract all possible branches

Description

It is assumed that multiple fragments from a common ancestor bay be charcterized by the their start-
and end-sites by integer values. For example, If 'abcdefg' is the ancestor, the fragments 'bcd' (from
position 2 to 4) to and 'efg' may then be assembled. To do so, all fragments must be presented
as matix specifying all start- and end-sites (and fragment-names). buildTree searches contiguous
fragments from columns 'posCo' (start/end) from 'disDat' to build tree & extract path information
starting with line 'startFr'. Made for telling if dissociated fragments contribute to long assemblies.
This function uses various functions of package data.tree which must be installed, too.

Usage

buildTree(
   disDat,
   startFr = NULL,
   posCo = c("beg", "end"),
   silent = FALSE,
   callFrom = NULL
)
cbindNR

Arguments

- **disDat** (matrix or data.frame) integer values with 1st column, 2nd column as end of fragments, rownames as unique IDs (node-names)
- **startFr** (integer) index for 1st node (typically =1 if 'disDat' sorted by "beg"), should point to a terminal node for consecutive growing of branches
- **posCo** (character) colnames specifying the begin & start sites in 'disDat', if NULL 1st & 2nd col will be used
- **silent** (logical) suppress messages
- **callFrom** (character) allow easier tracking of message(s) produced

Value

This function returns a list with $paths (branches as matrix with columns 'sumLen' & 'n'), $usedNodes (character vector of all names used to build tree) and $tree (object from data.tree)

See Also

package data.tree original function used Node; in this package : for exploiting edge/tree related issues simpleFragFig, countSameStartEnd and contribToContigPerFrag.

Examples

```r
frag2 <- cbind(beg=c(2,3,7,13,13,7,5,7,3),end=c(6,12,8,18,20,20,19,12,12,4))
rownames(frag2) <- c("A","B","C","D","E","F","G","H","I","J","K","L","M","N")
buildTree(frag2)
countSameStartEnd(frag2)
```

Description

cbindNR combines all matrixes given as arguments to non-redundant column names (by ADDING the number of 'duplicated' columns !). Thus, this function works similar to cbind, but allows combining multiple matrix-objects containing redundant column-names. Of course, all input-matrixes must have the same number of rows ! By default, the output gets sorted by column-names. Note, due to the use of '...' arguments must be given by their full argument-names, lazy evaluation might not recognize properly argument names.

Usage

cbindNR(
  ..., convertDfToMatr = TRUE, sortOutput = TRUE, summarizeAs = "sum",
)
Arguments

- all matrixes to get combined in cbind way
- convertDFtoMatr: (logical) decide if output should be converted to matrix
- sortOutput: (logical) optional sorting by column-names
- summarizeAs: (character) decide of combined values should get summed (default, 'sum') or averaged ('mean')
- silent: (logical) suppress messages
- callFrom: (character) allow easier tracking of messages produced

Value

This function returns a matrix or data.frame (as cbind would return)

See Also

- cbind, nonAmbiguousNum, firstOfRepLines

Examples

```r
ma1 <- matrix(1:6, ncol=3, dimnames=list(1:2, LETTERS[3:1]))
ma2 <- matrix(11:16, ncol=3, dimnames=list(1:2, LETTERS[3:5]))
cbindNR(ma1, ma2)
cbindNR(ma1, ma2, summarizeAs="mean")
```

---

checkAvSd

Check how multiple groups of data separate or overlap based on mean +/- sd

Description

checkAvSd compares if/how neighbour groups separate/overlap via the 'engineering approach' (+/- 2 standard-deviations is similar to a=0.05 t.test). This approach may be used as less elegant alternative to (multi-group) logistic regression. The function uses 'daAv' as matrix of means (rows are tested for up/down character/progression) which get compared with boundaries taken from daSd (for Sd values of each mean in 'daAv').
checkAvSd

Usage

checkAvSd(
  daAv,
  daSd,
  nByGr = NULL,
  multSd = 2,
  codeConst = "const",
  extSearch = FALSE,
  outAsLogical = TRUE,
  silent = FALSE,
  callFrom = NULL
)

Arguments

daAv  matrix or data.frame
daSd  matrix or data.frame
nByGr  optional specifying number of Elements per group, allows rather using SEM (adopt to variable n of different groups)
multSd  (numeric) the factor specifying how many sd values should be used as margin
codeConst  (character) which term/word to use when specifying 'constant'
extSearch  (logical) if TRUE, extend search to one group further (will call result 'nearUp' or 'nearDw')
outAsLogical  to switch between 2col-output (separate col for 'up' and 'down') or simple categorical vector ('const','okDw','okUp')
silent  (logical) suppress messages
callFrom  (character) allow easier tracking of message(s) produced

Value

vector describing character as 'const' or 'okUp','okDw' (or if extSearch=TRUE 'nearUp','nearDw')

See Also

rowGrpMeans

Examples

mat1 <- matrix(rep(11:24,3)[1:40],byrow=TRUE,ncol=8)
checkGrpOrderSEM(mat1,grp=gl(3,3)[-1])
checkAvSd(rowGrpMeans(mat1,gl(3,3)[-1]),rowGrpSds(mat1,gl(3,3)[-1]) )
# consider variable n :
checkAvSd(rowGrpMeans(mat1,gl(3,3)[-1]),rowGrpSds(mat1,gl(3,3)[-1]),nByGr=c(2,3,3))
checkFilePath  

**Check If File Is Available For Reading**

**Description**

This function allows testing if a given file-name corresponds to an existing file (e.g., for reading later). Indications to the path and file-extensions may be given separately. If no files do match .gz compressed versions may be searched, too.

**Usage**

```r
checkFilePath(
  fileName,  # (character) name of file to be tested; may also include an absolute or relative path
  path,      # (character, length=1) optional separate entry for path of fileName
  expectExt = "",  # (character) file extension (will not be considered if "")
  compressedOption = TRUE,  # (logical) also look for .gz compressed files
  strictExtension = FALSE,  # (logical) decide if extension (expectExt) - if given - should be considered obligatory
  stopIfNothing = FALSE,  # (logical) decide if function should give error or warning if no files found
  silent = FALSE,  # (logical) suppress messages
  debug = FALSE,  # (logical) additional messages for debugging
  callFrom = NULL
)
```

**Arguments**

- `fileName` (character) name of file to be tested; may also include an absolute or relative path
- `path` (character, length=1) optional separate entry for path of `fileName`
- `expectExt` (character) file extension (will not be considered if "")
- `compressedOption` (logical) also look for .gz compressed files
- `strictExtension` (logical) decide if extension (`expectExt`) - if given - should be considered obligatory
- `stopIfNothing` (logical) decide if function should give error or warning if no files found
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

**Details**

When the filename given by the user exists but its file-extension is not matching `expectExt` the argument `strictExtension` allows to decide if the filename will still be returned or not.

When `expectExt` is given, initial search will look for perfect matches. However, if nothing is found and `strictExtension=FALSE`, a more relaxed and non-case-sensitive search will be performed.
checkGrpOrder

Value
This function returns a character vector with verified path and file-name(s), returns NULL if nothing

See Also
file.exists

Examples
(RhomeFi <- list.files(R.home()))
file.exists(file.path(R.home(), "bin"))
checkFilePath(c("xxx","unins000"), R.home(), expectExt="dat")

Description
checkGrpOrder tests each line of 'x' if expected order appears. Used for comparing groups of measures with expected profile (simply by matching expected order)

Usage
checkGrpOrder(
  x,
  rankExp = NULL,
  revRank = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments
  x      matrix or data.frame
  rankExp  (numeric) expected order for values in columns, default 'rankExp' = 1:ncol(x)
  revRank  (logical) if 'revRank'=TRUE, the initial ranks & reversed ranks will be tested
  silent   (logical) suppress messages
  debug    (logical) display additional messages for debugging
  callFrom (character) allow easier tracking of messages produced

Value
 vector of logical values
checkGrpOrderSEM

See Also

checkGrpOrderSEM

Examples

set.seed(2005); mat1 <- rbind(matrix(round(runif(40),1),nc=4), rep(1,4))
checkGrpOrder(mat1)
checkGrpOrder(mat1,c(1,4,3,2))

checkGrpOrderSEM  
\textit{Check order of multiple groups including non-overlapping SEM-margins}

Description

checkGrpOrderSEM tests each line of ’x’ if expected order of (replicate-) groups (defined in ’grp’) appears intact, while including SEM of groups (replicates) via a proportional weight ’sdFact’ as (avGr1-gr1SEM) < (avGr1+gr1SEM) < (avGr2-gr2SEM) < (avGr2+gr2SEM). Used for comparing groups of measures with expected profile (by matching expected order) to check if data in ’x’ represting groups (’grp’) as lines follow. Groups of size=1: The sd (and SEM) can’t be estimated directly without any replicates, however, an estimate can be given by shrinking if ’shrink1sampSd’=TRUE under the hypothesis that the overall mechanisms determining the variances is constant across all samples.

Usage

checkGrpOrderSEM(
  x, 
  grp, 
  sdFact = 1, 
  revRank = TRUE, 
  shrink1sampSd = TRUE, 
  silent = FALSE, 
  callFrom = NULL
)

Arguments

x matrix or data.frame
grp (factor) to organize replicate columns of (x)
sdFact (numeric) is proportional factor how many units of SEM will be used for defining lower & upper bounds of each group
revRank (logical) optionally revert ranks
shrink1sampSd (logical)
silent (logical) suppress messages
callFrom (character) allow easier tracking of message(s) produced
checkSimValueInSer

Value

logical vector if order correct (as expected based on ranks)

See Also

takes only 10

Examples

mat1 <- matrix(rep(11:24,3)[1:40],byrow=TRUE,ncol=8)
checkGrpOrderSEM(mat1,grp=gl(3,3)[3])

Description

This function checks all values of 'x' for similar neighbour values within (relative) range of 'ppm'
(i.e. parts per million as measure of distance). By default values will be sorted internally, so if a
given value of x has anywhere in x another value close enough, this will be detected. However,
if sortX=FALSE only the values next to left and right will be considered. Return logical vector :
FALSE for each entry of 'x' if value inside of ppm range to neighbour (of sorted values)

Usage

checkSimValueInSer(
  x,
  ppm = 5,
  sortX = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

x numeric vector
ppm (numeric, length=1) ppm-range for considering as similar
sortX (logical) allows speeding up function when set to FALSE, for large data that are
already sorted
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced
checkStrictOrder

checkStrictOrder is a function that tests each line of a matrix or data frame for strict order (ascending, descending, or constant), relative to the line on its left. Each column of data is tested against its left neighbor.

**Arguments**
- `dat`: matrix or data frame
- `invertCount`: (logical) inverse counting (ie return 0 for all elements in order)
- `silent`: (logical) suppress messages
- `debug`: (logical) display additional messages for debugging
- `callFrom`: (character) allow easier tracking of messages produced

**Value**
- Matrix with counts of up pairs, down pairs, equal-pairs. If `invertCount`=TRUE all non-events are counted, ie a resulting 0 means all columns are following the described characteristics (with variable col-numbers easier to count).
checkVectLength

See Also

order, checkGrpOrder

Examples

set.seed(2005); mat1 <- rbind(matrix(round(runif(40),1),nc=4), rep(1,4))
checkStrictOrder(mat1); mat1[which(checkStrictOrder(mat1)[,2]==0),]

checkVectLength

Check length of vector

Description

checkVectLength checks argument 'x' for expected length 'expeL' and return either message or
error when expectation not met. May be used for parameter ('sanity') checking in other user front-
end functions.

Usage

checkVectLength(
  x,
  expeL = 1,
  stopOnProblem = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

x (numeric or character vector) input to check length
expeL (numeric) expected length
stopOnProblem (logical) continue on problems with message or stop (as error message)
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of message(s) produced

Value

This function returns NULL; it produces either error-message if length is not OK or optional message
if length is OK

Examples

aa <- 1:5; checkVectLength(aa, exp=3)
**cleanReplicates  **

*Replace Most Distant Values by NA*

**Description**

This procedure aims to straighten (clean) the most extreme values of noisy replicates by identifying the most distant points (among a set of replicates). The input 'x' (matrix or data.frame) is supposed to come from multiple different measures taken in replicates (e.g., weight of different individuals as rows taken as multiple replicate measures in subsequent columns).

**Usage**

```r
cleanReplicates(
  x,
  centrMeth = "median",
  nOutl = 2,
  retOffPos = FALSE,
  silent = FALSE,
  callFrom = NULL
)
```

**Arguments**

- **x**: matrix (or data.frame)
- **centrMeth**: (character) method to summarize (mean or median)
- **nOutl**: (integer) determines how many points per line will be set to NA (with n=1 the worst row of replicates will be 'cleaned')
- **retOffPos**: (logical) if TRUE, replace the most extreme outlier only
- **silent**: (logical) suppress messages
- **callFrom**: (character) allow easier tracking of messages produced

**Details**

With the argument nOutl the user chooses the total number of most extreme values to replace by NA. How many of the most extreme replicates of the whole dataset will be replaced by NA, i.e., with nOutl=1 only the single most extreme outlier will be replaced by NA. Outlier points are determined as point(s) with highest distance to (row) center (median and mean choice via argument 'centrMeth'). Thus function returns input data with 'removed' points set to NA, or if retOffPos=TRUE the most extreme/outlier positions.

**Value**

This function returns a matrix of same dimensions as input x, data-points which were tagged/removed are set to NA, or if retOffPos=TRUE the most extreme/outlier positions.
**closeMatchMatrix**

Reorganize results of search for close (similar) values in matrix-view

**Description**

`closeMatchMatrix` reorganizes/refines results from simple search of similar values of 2 sets of data by `findCloseMatch` (as list for one-to many relations) to more human friendly/readable matrix. This function returns results combining two sets of data which were initially compared (eg measured and theoretical values) as matrix-view using output of `findCloseMatch` and both original datasets. Additional information (covariables, annotation, ...) may be included as optional columns for either 'predMatr' or 'measMatr'. Note: It is important to run `findCloseMatch` with `sortMatch=FALSE`! Note: Results presented based on view of 'predMatr', so if multiple 'measMatr' are at within tolerated distance, lines of 'measMatr' will be repeated; Note: Distances 'disToMeas' and 'ppmToPred' are oriented: neg value if measured is lower than predicted (and pos values if higher than predicted); Note: Returns NULL when nothing within given limits of comparison.

**Usage**

```r
closeMatchMatrix(
  closeMatch,
  predMatr,
  measMatr,
  prefMatch = c("^x", "^y"),
  colPred = 1,
  colMeas = 1,
  limitToBest = TRUE,
  asDataFrame = FALSE,
  origNa = TRUE,
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)
```

**Arguments**

- `closeMatch` (list) output from `findCloseMatch`, ie list with hits for each 'x' (1st argument): named vectors of value & x index in name; run with 'sortMatch'=F
- `predMatr` (vector or matrix) predicted values, the column 'colPred' indicates which column is used for matching from `findCloseMatch`; if column 'id' present this column will be used as identifier for matching
- `measMatr` (vector or matrix) measured values, the column 'colMeas' indicates which column is used for matching from `findCloseMatch`; if column 'id' present this column will be used as identifier for matching

**Examples**

```r
mat3 <- matrix(c(19,20,30, 18,19,28, 16,14,35),ncol=3)
cleanReplicates(mat3, nOutl=1)
```
closeMatchMatrix

prefMatch (character, length=2) prefixes ('^x' and/or '^y') that may have been added by findCloseMatch

colPred (integer or text, length=1) column of 'predMatr' with main values of comparison

colMeas (integer or text, length=1) column of 'measMatr' with main measures of comparison

limitToBest (integer) column of 'measMatr' with main measures of comparison

asDataFrame (logical) convert results to data.frame if non-numeric matrix produced (may slightly slow down big results)

origNa (logical) will try to use original names of objects 'predMatr', 'measMatr', if they are not multi-column and not conflicting other output-names (otherwise 'predMatr', 'measMatr' will appear)

silent (logical) suppress messages

callFrom (character) allows easier tracking of message(s) produced

debug (logical) for bug-tracking: more/enhanced messages

Value

results as matrix-view based on initial results from findCloseMatch, including optional columns of suppelemental data for both sets of data for comparison. Returns NULL when nothing within limits

See Also

findCloseMatch, checkSimValueInSer

Examples

aA <- c(11:17); bB <- c(12.001,13.999); cC <- c(16.2,8,9,12.5,15.9,13.5,15.7,14.1,5)
(cloMa <- findCloseMatch(aA,cC,com="diff",lim=0.5,sor=FALSE))
# all matches (of 2d arg) to/within limit for each of 1st arg ('x'); 'y' ..to 2nd arg = cC
(maAa <- closeMatchMatrix(cloMa,aA,cC,lim=TRUE))
(maAa <- closeMatchMatrix(cloMa,aA,cC,lim=FALSE,origN=TRUE))
(maAa <- closeMatchMatrix(cloMa,cbind(valA=81:87,aA),cbind(valC=91:99,cC),colM=2,
colP=2,lim=FALSE))
(maAa <- closeMatchMatrix(cloMa,cbind(aA,val=81:87),cbind(valC=91:99,cC),colM=2,
colP=2,lim=FALSE,deb=TRUE))
a2 <- aA; names(a2) <- letters[1:length(a2)]; c2 <- cC; names(c2) <- letters[10+1:length(c2)]
(cloM2 <- findCloseMatch(x=a2,y=c2,com="diff",lim=0.5,sor=FALSE))
(maA2 <- closeMatchMatrix(cloM2,predM=cbind(valA=81:87,aA),measM=cbind(valC=91:99,cC),
colM=2,colP=2,lim=FALSE,asData=TRUE))
(maA2 <- closeMatchMatrix(cloM2,cbind(id=names(a2),valA=81:87,aA),cbind(id=names(c2),
valC=91:99,c2),colM=3,colP=3,lim=FALSE,deb=FALSE))
coinPermTest

Compare means of two vectors by permutation test

Description
Run coin-flipping like permutation tests (to compare difference of 2 means: 'x1' and 'x2') without any distribution-assumptions. This function uses the package coin, if not installed, the function will return NULL and give a warning.

Usage
coinPermTest(
  x1,
  x2,
  orient = "two.sided",
  nPerm = 5000,
  silent = FALSE,
  callFrom = NULL
)

Arguments
x1 numeric vector (to be compared with vector 'x2')
x2 numeric vector (to be compared with vector 'x1')
orient (character) may be "two.sided", "greater" or "less"
nPerm (integer) number of permutations
silent (logical) suppress messages
callFrom (character) allow easier tracking of messages produced

Value
This function returns an object of "MCp" class numeric output with p-values

See Also
oneway_test in LocationTests

Examples
coinPermTest(2, 3, nPerm=200)
**colMedSds**  
*Standard error of median for each column by bootstrap*

**Description**

Determine standard error (sd) of median by bootstrapping for multiple sets of data (rows in input matrix 'dat'). Note: The package `boot` must be installed from CRAN.

**Usage**

```r
colMedSds(dat, nBoot = 99, silent = FALSE, callFrom = NULL)
```

**Arguments**

- `dat` (numeric) matrix
- `nBoot` (integer) number if iterations
- `silent` (logical) suppress messages
- `callFrom` (character) allow easier tracking of messages produced

**Value**

This function returns a (numeric) vector with estimated standard errors

**See Also**

`boot`

**Examples**

```r
set.seed(2016); dat1 <- matrix(c(runif(200) + rep(1:10,20)), ncol=10)
colMedSds(dat1)
```

**colorAccording2**  
*Transform numeric values to color-gradient*

**Description**

This function helps making color-gradients for plotting a numerical variable. It requires the package 'RColorBrewer' being installed from CRAN. Note: RColorBrewer palettes were not integrated here, since they are not continuous.
Usage

colorAccording2(
  x,
  gradTy = "rainbow",
  nStartOmit = NULL,
  nEndOmit = NULL,
  revCol = FALSE,
  alpha = 1,
  callFrom = NULL
)

Arguments

x (character) color input

gradTy (character) type of gradeint may be 'rainbow', 'heat.colors', 'terrain.colors', 'topo.colors', 'cm.colors', 'hcl.colors', 'grey.colors', 'gray.colorsW' or 'logGray'

nStartOmit (integer) omit n steps from beginning of gradient range

nEndOmit (integer or "sep") omit n steps from end of gradient range, if nEndOmit="sep" 20 percent of initial grades will be removed to obtain 'separate' ie non-closing color-circles/gradients eg with rainbow

revCol (logical) reverse order

alpha (numeric) optional transparency value (1 for no transparency, 0 for complete opaqueness)

callFrom (character) allow easier tracking of message(s) produced

Value

This function returns a character vector (of same length as x) with color encoding

See Also

cut

Examples

set.seed(2015); dat1 <- round(runif(15),2)
plot(1:15,dat1,pch=16,cex=2,col=colorAccording2(dat1))
plot(1:15,dat1,pch=16,cex=2,col=colorAccording2(dat1,nStartO=0,nEndO=4,revCol=TRUE))
plot(1:9,pch=3)
points(1:9,1:9,col=transpGraySca(st=0,en=0.8,nSt=9,trans=0.3),cex=42,pch=16)
Description

colSds is a speed optimized sd for matrix or data.frames. It and treats each line as an independent set of data for calculating the sd (equiv to apply(dat,1,sd)). NAs are ignored from data.

Usage

colSds(dat)

Arguments

dat matrix (or data.frame) with numeric values (may contain NAs)

Value

numeric vector of sd values

See Also

sd

Examples

set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),nc=10)
colSds(dat1)

combinatIntTable

Planing for making all multiplicative combinations

Description

Provide all combinations for each of n elements of vector ’nMax’ (positive integer, eg number of max multiplicative value). For example, imagine, we have 3 cities and the (maximum) voting participants per city. Results must be read vertically and allow to see all total possible compositons.

Usage

combinatIntTable(
  nMax,
  include0 = TRUE,
  asList = FALSE,
  callFrom = NULL,
  silent = TRUE
)
**Arguments**

- `nMax` (positive integer) could be max number of voting participants form different cities, eg Paris max 2 persons, Lyon max 1 person ...
- `include0` (logical) include 0 occurrences, ie provide all combinations starting from 0 or from 1 up to nMax
- `asList` (logical) return result as list or as array
- `callFrom` (character) allow easier tracking of messages produced
- `silent` (logical) suppress messages

**Value**

list or array (as 2- or 3 dim) with possible number of occurrences for each of the 3 elements in nMax. Read results vertical: `out[[1]]` or `out[,1]`.. (multiplicative) table for 1st element of nMax; `out[,2]` .. for 2nd

**See Also**

`combn`

**Examples**

```r
combinatIntTable(c(1,1,1,2), include0=TRUE, asList=FALSE, silent=TRUE)
## Imagine we have 3 cities and the (maximum) voting participants per city :
## Imagine we have 3 cities and the (maximum) voting participants per city :
## Imagine we have 3 cities and the (maximum) voting participants per city :
## Imagine we have 3 cities and the (maximum) voting participants per city :
## Imagine we have 3 cities and the (maximum) voting participants per city : nMa <- c(Paris=2, Lyon=1, Strasbourg=1) combinatIntTable(nMa, include0=TRUE, asList=TRUE, silent=TRUE)
```

---

**Description**

The aim of this function is to choose a fixed number (`nCombin`) of list-elements from `lst` and count the number of common values/words. Furthermore, one can define levels to fine-tune the types of combinations to examine. In case multiple combinations for a given level are possible, some basic summary statistics are provided, too.

**Usage**

```r
combineAsN(
  lst,
  lev = NULL,
  nCombin = 3,
  remDouble = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```
**Arguments**

- **lst**: (list of character or integer vectors) main input
- **lev**: (character) define groups of lst
- **nCombin**: (integer) number of list-elements to combine from lst
- **remDouble**: (logical) remove intra-duplicates (defaults to TRUE)
- **silent**: (logical) suppress messages
- **debug**: (logical) additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

**Details**

Note of caution: With very long lists and/or high numbers of repeats of given levels, however, the computational effort increases very much (like it does when using `table`). Thus, when exploring all different combinations of large data-sets may easily result in queries consuming many resources (RAM and processing time)! It is recommended to start testing with test smaller sub-groups.

The main idea of this function is to count frequency of terms when combining different drawings. For example, you ask students from different cities which are their preferred hobbies, they may have different preference depending on the city (defined by `lev`). Now, if you want to make groups of 3 students, possibly with one from each city (A, B and C), you want to count (estimate) the frequency of different combinations possible. Thus, using this function all combinations of the students from city A with the students from city B and C will be made when counting the number of common hobbies (by `nCombin` students). Then, all counting results will be summarized to the average count for the various categories (which hobbies were seen once, twice or 3 times...), `sem` (standard error of the mean) and `CI` (95% confidence interval).

Of course, the number of potential combinations may quickly get very large. Using the argument `remDouble=TRUE` you can limit the search to either finding all students giving the same answer plus all student giving different answers. In this case, when a given level appears multiple times, all possible combinations using one of the respective entries will be be made with the other levels.

**Value**

This function returns an array with 3 dimensions: i) the combinations of `nCombin` list-elements, ii) the number of counts (`n`), `sem` (standard error of the mean), `CI` (confidence interval) and `sd`

**See Also**

`table`, `replicateStructure`

**Examples**

```r
## all list-elements are considered equal
tm1 <- list(a1=LETTERS[1:17], a2=LETTERS[3:19], a3=LETTERS[6:20], a4=LETTERS[8:22])
combineAsN(tm1, lev=gl(1,4))[,1,]

## different levels/groups in list-elements
tm4 <- list(a1=LETTERS[1:15], a2=LETTERS[3:16], a3=LETTERS[6:17], a4=LETTERS[8:19],
            b1=LETTERS[5:19], b2=LETTERS[7:20], b3=LETTERS[11:24], b4=LETTERS[13:25], c1=LETTERS[17:26],
```
combineByEitherFactor

Create factor-like column regrouping data regrouping simultaneously by two factors

Description

This function aims to address the situation when two somehow different groupings (of the same data) exist and need to be joined. It is not necessary that both alternative groupings use the same labels, neither. combineByEitherFactor adds new (last) column named 'grp' to input matrix representing the combined factor relative to 2 specified columns from input matrix mat (via 'refC1','refC2'). Optionally, the output may be sorted and a column giving n per factor-level may be added. The function treats selected columns of mat as pairwise combination of 2 elements (that may occur multiple times over all lines of mat) and sorts/organizes all instances of such combined elements (ie from both selected columns) as repeats of a given group, who's class number is given in output column 'grp', the (total) number of repeats may be displayed in column 'nGrp' (nByGrp=TRUE). If groups are overlapping (after re-ordering), an iterative process of max 3x2 passes will be launched after initial matching. Works on numeric as well as character input.

Usage

```r
combineByEitherFactor("mat", refC1, refC2, nByGrp = FALSE, convergeMax = TRUE, callFrom = NULL, debug = FALSE, silent = FALSE)
```

Arguments

- **mat**: main input matrix
- **refC1**: (integer) column-number of 'mat' to use as 1st set
- **refC2**: (integer) column-number of 'mat' to use as 2nd set
- **nByGrp**: (logical) add last col with n by group
- **convergeMax**: (logical) if TRUE, run 2 add’l iterative steps to search convergence to stable result
- **callFrom**: (character) allow easier tracking of message(s) produced
- **debug**: (logical) display additional messages for debugging
- **silent**: (logical) suppress messages
Value
This function returns a matrix containing both selected columns plus additional column(s) indicating group-number of the pair-wise combination (and optional the total n by group)

Examples
```
n <- rep(c("a","e","b","c","d","g","f"),c(3,1,2,2,1,2,1))
qq <- rep(c("m","n","p","o","q"),c(2,1,1,4,4))
nq <- cbind(nn,qq)[c(4,2,9,11,6,10,7,3,5,1,12,8),]
combineByEitherFactor(nq,1,2,nBy=TRUE); combineByEitherFactor(nq,1,2,nBy=FALSE)
combineByEitherFactor(nq,1,2,conv=FALSE); combineByEitherFactor(nq,1,2,conv=TRUE)
##
mm <- rep(c("a","b","c","d","e"),c(3,4,2,3,1)); pp <- rep(c("m","n","o","p","q"),c(2,2,2,2,5))
combineByEitherFactor(cbind(mm,pp), 1, 2, con=FALSE, nBy=TRUE)
combineByEitherFactor(cbind(mm,pp), 1, 2, con=TRUE, nBy=TRUE)
```

Description
Search points in x,y space that are located very close and thus likely to overlap. In case of points close enough, various options for joining names (and shortening longer descriptions) are available.

Usage
```
combineOverlapInfo(
  dat,
  suplInfo = NULL,
  disThr = 0.01,
  addNsimil = TRUE,
  txtSepChar = ",",
  combSym = "+",
  maxOverl = 50,
  callFrom = NULL,
  debug = FALSE,
  silent = FALSE
)
```

Arguments
- **dat** (matrix) matrix or data.frame with 2 cols (used ONLY 1st & 2nd column !), used as x & y coordinates
- **suplInfo** (NULL or character) when points are considered overlapping the text from 'suplInfo' will be reduced to fragment before 'txtSepChar' and combined (with others from overlapping text) using 'combSym', if NULL $combInf will appear with row-numbers
**combineRedBasedOnCol**

Combine/reduce redundant lines based on specified column

**Description**

This function works similar to `unique`, but it takes a matrix as input and considers one specified column to find unique instances. It identifies 'repeated' lines of the input-matrix (or data.frame) 'mat' based on (repeated) elements in/of column with name 'colNa' (or column-number). Redundant lines (ie repeated lines) will disappear in output. Eg used with extracted annotation where 1 gene has many lines for different GO annotation.

**Usage**

```
combineRedBasedOnCol(mat, colNa, sep = " ", silent = FALSE, callFrom = NULL)
```
combineRedundLinesInList

Arguments

- **mat** 
  input matrix or data.frame
- **colNa** 
  character vector (length 1) matching 1 column name (if mult only 1st will be used), in case of mult matches only 1st used
- **sep** 
  (character) separator (default=";")
- **silent** 
  (logical) suppress messages
- **callFrom** 
  (character) allow easier tracking of messages produced

Value

matrix containing the input matrix without lines considered repeated (unique-like)

See Also

findRepeated, firstOfRepLines, organizeAsListOfRepl, combineRedundLinesInList

Examples

```r
matr <- matrix(c(letters[1:6], "h","h","f","e", LETTERS[1:5]), ncol=3,
               dimnames=list(letters[11:15], c("xA", "xB", "xC")))
combineRedBasedOnCol(matr, colN="xB")
combineRedBasedOnCol(rbind(matr[1,], matr), colN="xB")

combineRedundLinesInList

Combine Redundant Lines In List

Description

This function provides help for combining/summarizing lines of numeric data which may be summarized according to reference vector or matrix of annotation (part of the same input-list). The data and reference will be aligned and data corresponding to redundant information be combined/summarized.

Usage

```r
combineRedundLinesInList(
  lst,
  refNa = "ref",
  datNa = "quant",
  refColNa = "GeneName",
  supRefColNa = NULL,
  summarizeType = "av",
  NA.rm = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```
Arguments

**lst**
(list) main input, containing matrix or data.frame of numeric data (see datNa and annotation (see refNa) and possibly unrelated stuff

**refNa**
(character) name of list-element containing annotation

**datNa**
(character) name(s) of list-element(s) containing numeric/quantitation data

**refColNa**
(character) in case the list-element to be used as reference is matrix or data.frame, the column to be used must be specified here

**supRefColNa**
(character) in case the lst$refNa has no rownames, the content of column lst$supRefColNa will be used instead

**summarizeType**
(character) the summarization method gets specified here; so far 'sum','av','med','first' and 'last' are implemented

**NA.rm**
(logical) pass to summarizing functions order to omit NAs, defaults to TRUE

**silent**
(logical) suppress messages

**debug**
(logical) additional messages for debugging

**callFrom**
(character) allow easier tracking of messages produced

Details

All input data should be in a list, ie one or multipl matrix or data.frame for numeric data (see argument datNa), as well as the reference (see argument refNa). The reference may be a named character vector or a matrix for which the column to be used should be specified using the argument refColNa. In case the annotation is a matrix, the rownames will be used as unique/independent identifiers to adjust potentially different order of numeric data and annotation. In absence of rownames, an additional column supRefColNa of the annotation may be designed for adjusting the order of annotation and numeric data.

The numeric list may contain multiple matrixes or data.frames which will all be summarized by the same procedure as long as they have the same initial dimensions and are specified by refNa.

Please note that all other list elements from input not specified by refNa (or datNa) will be maintained in the output just as they are.

Value

This function returns a list of same length as input

See Also

`findRepeated`, `firstOfRepLines`, `organizeAsListOfRepl`, `combineRedBasedOnCol`

Examples

```r
x1 <- list(quant=matrix(11:34, ncol=3, dimnames=list(letters[8:1], LETTERS[11:13])),
           annot=matrix(paste0(LETTERS[c(1:4,6,3:5)],LETTERS[c(1:4,6,3:5)]), ncol=1,
                        dimnames=list(paste(letters[1:8]),"xx"))
combineRedundLinesInList(lst=x1, refNa="annot", datNa="quant", refColNa="xx")
```
combineRedundLinesInListAcRef

Combine Redundant Lines In List, Deprecated

Description

The function combineRedundLinesInListAcRef() has been deprecated and replaced by combineRedundLinesInList() from the same package.

Usage

```r
combineRedundLinesInListAcRef(
  lst,
  listNa = c("ref", "quant"),
  refColNa = "xx",
  summarizeType = "av",
  NA.rm = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `lst` (list) main input
- `listNa` (character) names of list-elements containing quantitation data (1st position) and protein/line annotation (2nd position)
- `refColNa` (character) in case the list-element to be used as reference is matrix or data.frame, the column to be used must be specified here
- `summarizeType` (character) the summarization method gets specified here; so far 'sum','av','med','first' and 'last' are implemented
- `NA.rm` (logical) pass to summarizing functions order to omit NAs, defaults to TRUE
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a list of same length as input

See Also

- `combineRedundLinesInList`
Examples

```r
x1 <- list(quant=matrix(11:34, ncol=3, dimnames=letters[8:1], LETTERS[11:13])),
    annot=matrix(paste0(LETTERS[c(1:4,6,3:5)],LETTERS[c(1:4,6,3:5)]), ncol=1,
        dimnames=list(paste(letters[1:8]),"xx"))
## please use combineRedundLinesInList()
combineRedundLinesInList(lst=x1, refNa="annot", datNa="quant", refColNa="xx")
```

---

**Description**

Suppose multiple measures (like multiple channels) are taken for subjects and these measures are organized as groups in a list, like multiple parameters (= channels) or types of measurements (typically many parameters are recorded when screening compounds in microtiter plates). Within one parameter/channel all replicate-data from separate list-entries ('lst') will get combined according to names of list-elements. The function will trim any redundant text in names of list-elements, try to isolate separator (may vary among replicate-groups, but should be 1 character long). eg names "hct116 1.1.xlsx" & "hct116 1.2.xlsx" will be combined as replicates, "hct116 2.1.xlsx" will be considered as new group.

**Usage**

```r
combineReplFromListToMatr(lst, callFrom = NULL)
```

**Arguments**

- **lst** (list) list of arrays (typically multi-parameter measures of microtiterplate data)
- **callFrom** (character) allows easier tracking of message(s) produced

**Value**

list of arrays now with same dimension of arrays (but shorter, since replicate-arrays were combined)

**See Also**

`extr1chan`, `organizeAsListOfRepl`

**Examples**

```r
lst2 <- list(aa_1x=matrix(1:12,nrow=4,byrow=TRUE),ab_2x=matrix(24:13,nrow=4,byrow=TRUE))
combineReplFromListToMatr(lst2)
```
**combineSingleT**  
*Get all combinations with TRUE from each column*

**Description**

This function addresses the case when multiple alternatives ways exit to combine two elements. combineSingleT makes combinatory choices: if multiple TRUE in given column of ‘mat’ make all multiple selections with always one TRUE from each column. The resultant output contains index for first and second input columns elements to be combined.

**Usage**

```
combineSingleT(mat)
```

**Arguments**

- `mat` 2-column matrix of logical values

**Value**

matrix with indexes of combinations of TRUE

**Examples**

```r
## Example: Fist column indicates which boys want to dance and second column
## which girls want to dance. So if several boys want to dance each of the girls
## will have the chance to dance with each of them.
matr <- matrix(c(TRUE,FALSE,TRUE,FALSE,TRUE,FALSE),ncol=2)
combineSingleT(matr)
```

---

**completeArrLst**  
*Complete list of arrays for same dimensions*

**Description**

This functions aims to inspect repeating structures of data given as list of arrays and will try to complete arrays with fewer lines or columns (as this may appear eg with the very last set of high-throughput screening data if fewer measures remain in the last set). Thus, the dimensions of the arrays are compared and cases with fewer (lost) columns (eg fewer experimental replicates) will be adjust/complete by adding column(s) of NA. Used eg when at reading microtiterplate data the last set is not complete.

**Usage**

```
completeArrLst(arrLst, silent = FALSE, callFrom = NULL)
```
Arguments

arrLst  
(list) list of arrays (typically 1st and 2nd dim for specific genes/objects, 3rd for different measures associated with)

silent  
(logical) suppress messages

callFrom  
(character) allows easier tracking of message(s) produced

Value

list of arrays, now with same dimension of arrays

See Also

organizeAsListOfRepl, extr1chan

Examples

arr1 <- array(1:24,dim=c(4,3,2),dimnames=list(c(LETTERS[1:4]),
paste("col",1:3,sep=""),c("ch1","ch2")))
arr3 <- array(81:96,dim=c(4,2,2),dimnames=list(c(LETTERS[1:4]),
paste("col",1:2,sep=""),c("ch1","ch2")))
arrL3 <- list(pl1=arr1,pl3=arr3)
completeArrLst(arrL3)
Arguments

- **x** (vector) the values to be matched
- **table** (vector) the values to be matched against (ie reference)
- **sep** (character) separator character in case concatenation of entries is tested
- **sepPattern** (character or NULL) optional custom pattern for splitting concatenations of x) and table (in case NULL) is not sufficient
- **globalPat** (character) pattern for additional trimming of search-terms. If `globalPat="digitExtension"` all terminal digits will not be considered when matching
- **nomatch** (vector) similar to `match` the value to be returned in the case when no match is found
- **incomparables** (vector) similar to `match`, a vector of values that cannot be matched. Any value in x matching a value in this vector is assigned the nomatch value.
- **extensPat** (logical) similar to `match` the value to be returned in the case when no match is found
- **silent** (logical) suppress messages
- **debug** (logical) additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced

Details

The main motivation to create this function was to be able to treat concatenated entries and to look if any of the concatenated values match to 'x'. This function offers additional options for trimming values before running the main comparison.

Of course, the concatenation strategy must be known and only a single concatenation separator (which may be multiple characters long) may be used for both x and match. Thus result will only indicate that at least one of the concatenated terms had a match, but not which one. Finally, both vectors x and table may contain concatenated terms. In this case this function will require much more computational resources due to the increased combinatorics when comparing larger vectors.

Please note, that in case of multiple to multiple matches, only the first hit gets reported.

The argument `globalPat="digitExtension"` allows eg reducing 'A1234-4' to 'A1234'.

Value

This function returns a character vector with verified path and file-name(s), returns NULL if nothing

See Also

- `match` (for two simple vectors without concatenated terms), `grep`

Examples

```r
x1 <- c("ZZ","YY","AA","BB-2","DD","CCdef","Dxy")  # modif of single ID (no concat)
concatMatch(x1, tab2)
```

confInt

Confidence Interval To Given Alpha

Description

This little function returns the confidence interval associated to a given significance level \( \alpha \) under the hypothesis of the Normal distribution is valid.

Usage

\[
\text{confInt}(x, \ alpha = 0.05, \ \text{distrib} = \text{"Normal"}, \ \text{silent} = \text{FALSE}, \ \text{callFrom} = \text{NULL})
\]

Arguments

- \( x \) (numeric) main input
- \( \alpha \) (numeric) significance level, accepted type I error
- \( \text{distrib} \) (character) distribution, so far only Normal is implemented
- \( \text{silent} \) (logical) suppress messages
- \( \text{callFrom} \) (character) allow easier tracking of message(s) produced

Value

This function returns the confidence interval to a given \( \alpha \) under the hypothesis of the Normal distribution.

See Also

TDist; confint

Examples

\[
\text{confInt(c(5,2:6))}
\]
contribToContigPerFrag

Characterize individual contribution of single edges in tree-structures

Description

This function helps investigating tree-like structures with the aim of indicating how much individual tree components contribute to compose long stretches. contribToContigPerFrag characterizes individual (isolated) contribution of single edges in tree-structures. Typically used to process/exploit summarized trees (as matrix) made by buildTree which makes use of the package data.tree. For example if A,B and C can be joined as well and B +D, this function will check if A+B+C is longer and if A contributes to the longest tree.

Usage

contribToContigPerFrag(joinMat, fullLength = NULL, nDig = 3)

Arguments

joinMat (matrix) matrix with concatenated edges as rownames (separated by slashes), column sumLen for total length and column n for number of edges

fullLength (integer) custom total length (useful if the concatenated edges do not cover 100 percent of the original precursor whose fragments are studied)

nDig (integer) rounding: number of digits for 3rd column len.rat in output

Value

matrix of 3 columns: with length of longest tree-branches where given edge participates (column sumLen), the (total) number of edges therein (col n.frag) and a relative value (len.rat)

See Also

to build tree buildTree

Examples

path1 <- matrix(c(17,19,18,17, 4,4,2,3),ncol=2, dimnames=list(c("A/B/C/D","A/B/G/D","A/H","A/H/I"),c("sumLen","n")))
contribToContigPerFrag(path1)
**conv01toColNa**  
*Convert matrix of integer to matrix of x-times repeated column-names*

**Description**

`conv01toColNa` transforms matrix of integers (eg 0 and 1) to repeated & concatenated text from argument `colNa`, the character string for 0 occurrences of argument `zeroTex` may be customized. Used eg when specifying (and concatenating) various counted elements (eg properties) along a vector like variable peptide modifications in proteomics.

**Usage**

```r
conv01toColNa(mat, colNa = NULL, zeroTex = "", pasteCol = FALSE)
```

**Arguments**

- `mat`: input matrix (with integer values)
- `colNa`: alternative (column-)names to the ones from 'mat' (default colnames of 'mat')
- `zeroTex`: text to display if 0 (default ")"
- `pasteCol`: (logical) allows to collapse all columns to single chain of characters in output

**Value**

character vector

**Examples**

```r
(ma1 <- matrix(sample(0:3,40,repl=TRUE), ncol=4, dimnames=list(NULL, letters[11:14])))
conv01toColNa(ma1)
conv01toColNa(ma1, colNa=LETTERS[1:4], ze=".")
conv01toColNa(ma1, colNa=LETTERS[1:4], pasteCol=TRUE)
```

---

**convColorToTransp**  
*Assign new transparency to given colors*

**Description**

This function allows (re-)defining a new transparency. A color encoding vector will be transformed to the same color(s) but with new transparency (alpha).

**Usage**

```r
convColorToTransp(color, alph = 1)
```
convMatr2df

Arguments

color  (character) color input
alphp  (numeric) transparency value (1 for no transparency, 0 for complete opaqueness), values <1 will be treated as percent-values

Value

character vector (of same length as input) with color encoding for new transparency

See Also

rgb, par

Examples

col0 <- c("#998FCC","#5AC3BA","#CBD34E","#FF7D73")
col1 <- convColorToTransp(col0,alph=0.7)
layout(1:2)
pie(rep(1,length(col0)),col=col0)
pie(rep(1,length(col1)),col=col1,main="new transparency")

convMatr2df  Convert matrix (eg with redundant) row-names to data.frame

Description

This function provides flexible converting of matrix to data.frame. For example repeated/redundant rownames are not allowed in data.frame(), thus the corresponding column-names have to be renamed using a counter-suffix. In case of non-redundant rownames, a new column 'addIniNa' will be introduced at beginning to document the initial (redundant) rownames, non-redundant rownames will be created. Finally, this functions converts the corrected matrix to data.frame and checks/converts columns for transforming character to numeric if possible. If the input is a data.frame containing factors, they will be converted to character before potential conversion. Note: for simpler version (only text to numeric) see from this package .convertMatrToNum.

Usage

convMatr2df(
  mat, 
  addIniNa = TRUE, 
  duplTxtSep = "_", 
  silent = FALSE, 
  callFrom = NULL
)
**convToNum**

**Convert vector to numeric**

**Description**

This function checks if input vector/character string contains numbers (with or without comma) and attempts converting to numeric. This functions was designed for extracting the numeric part of character-vectors (or matrix) containing both numbers and character-elements. Depending on the parameters convert and remove text-entries can be converted to NA (in resulting numeric objects) or removed (the number of elements/lines gets reduced, in consequence). Note: if `x` is a matrix, its matrix-dimensions & -names will be preserved. Note: so far Inf and -Inf do not get recognized as numeric.

**Arguments**

- **mat**: matrix (or data.frame) to be converted
- **addIniNa**: (logical) if TRUE an additional column ('ID') with rownames will be added at beginning
- **duplTxtSep**: (character) separator for enumerating replicated names
- **silent**: (logical) suppres messages
- **callFrom**: (character) allow easier tracking of message(s) produced

**Value**

This functions returns a data.frame equivalent to the input matrix, an additional column named 'ID' will be added for initial rownames

**See Also**

- `numeric`, for simpler version (only text to numeric) see from this package .convertMatrToNum

**Examples**

```r
dat1 <- matrix(1:10, ncol=2)
rownames(dat1) <- letters[c(1:3,2,5)]
## as.data.frame(dat1) ... would result in an error
convMatr2df(dat1)

df1 <- data.frame(a=as.character((1:3)/2), b=LETTERS[1:3], c=1:3)
str(convMatr2df(df1))

df2 <- df1; df2$b <- as.factor(df2$b)
str(convMatr2df(df2))
```
Usage

convToNum(
  x,
  autoConv = TRUE,
  spaceRemove = TRUE,
  convert = c(NA, "sparseChar"),
  remove = NULL,
  euroStyle = TRUE,
  sciIncl = TRUE,
  callFrom = NULL,
  silent = TRUE
)

Arguments

x           vector to be converted
autoConv    (logical) simple automatic conversion based on as.numeric; if TRUE all other
            arguments except spaceRemove will not be considered
spaceRemove (logical) to remove all heading and trailing (white) space (until first non-space
            character)
convert     (character) define which type of non-conform entries to convert to NAs. Note,
            if remove is selected to eliminate character-entries they cannot be converted
            any more. Use 'allChar' for all character-entries; 'sparseChar' sparse (ie rare)
            character entries; NA for converting 'Na' or 'na' to NA; if 'none' or NULL no
            conversions at all.
remove      (character) define which type of non-conform entries to remove, removed items
            cannot be converted to NA any more. Use 'allChar' for removing all character
            entries; NA for removing all instances of NA (except thise created by converting
            text); all elements will be kept if 'none' or NULL.
euroStyle   (logical) if TRUE will convert all ',' (eg used as European decimal-separator) to
            '.' (as internally used by R as decimal-separator), thus allowing converting the
            European decimal format.
sciIncl     (logical) include recognizing scientific notation (eg 2e-4)
callFrom    (character) allow easier tracking of messages produced
silent      (logical) suppress messages

Details

This function may be used in two modes, depending if argument autoConv is TRUE or FALSE. The
first options allows accessing an automatic mode based on as.numeric, while the second options
investigates all characters if they may belong to numeric expressions and allows removing specific
text-elements.

Value

This function returns a numeric vector (or matrix (if 'x' is matrix))
coordOfFilt

See Also

table and as.numeric (on same help-page)

Examples

x1 <- c("+4"," + 5","6","bb","Na","-7")
convToNum(x1)
convToNum(x1, autoConv=FALSE, convert=c("allChar"))
convToNum(x1, autoConv=FALSE)  # too many non-numeric instances for 'sparseChar'

x2 <- c("+4"," + 5","6","-7"," - 8","1e6","+ 2.3e4","-3E4","- 4E5")
convToNum(x2)
convToNum(x2, autoConv=FALSE, convert=NA, remove=c("allChar",NA))
convToNum(x2, autoConv=FALSE, convert=NA, remove=c("allChar",NA), sciIncl=FALSE)

doFnFilt

grodd coordinates of values/points in matrix according to filtering condition

Description

Get coordinates of values/points in matrix according to filtering condition

Usage

doFnFilt(mat, cond, sortByRows = FALSE, silent = FALSE, callFrom = NULL)

Arguments

mat (matrix or data.frame) matrix or data.frame
cond (logical or integer) condition/test to see which values of mat fulfill test, or integer of index passing
sortByRows (logical) optional sorting of results by row-index
silent (logical) suppress messages
callFrom (character) allow easier tracking of message(s) produced

Value

matrix columns 'row' and 'col'

See Also

which
Examples

```r
set.seed(2021); ma1 <- matrix(sample.int(n=40, size=27, replace=TRUE), ncol=9)
## let's test which values are >37
which(ma1 > 37)  # doesn't tell which row & col
coordOfFilt(ma1, ma1 > 37)
```

`correctToUnique` checks 'x' for unique entries, while maintaining the original length. If necessary a counter will be added to non-unique entries.

**Description**

`correctToUnique` checks 'x' for unique entries, while maintaining the original length. If necessary a counter will be added to non-unique entries.

**Usage**

```r
correctToUnique(
  x,
  sep = "_",
  atEnd = TRUE,
  maxIter = 4,
  NAenum = TRUE,
  silent = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `x` (input character vector)
- `sep` (character) separator used when adding counter
- `atEnd` (logical) decide location of placing the counter (at end or at beginning of initial text)
- `maxIter` (numeric) max number of iterations
- `NAenum` (logical) if TRUE all NAs will be enumerated (NA_1, NA_2, ...)
- `silent` (logical) suppress messages
- `callFrom` (character) for better tracking of use of functions

**Value**

This function returns a character vector

**See Also**

- `unique` will simply remove repeated elements, i.e., length of 'x' won't remain constant, `filtSizeUniq` is more complex and slower, `treatTxtDuplicates`
**correctWinPath**

**Examples**

```
correctToUnique(c("li0","n",NA,NA,rep(c("li2","li3"),2),rep("n",4)))
```

```
correctWinPath(  
x,
  asHtml = FALSE,
  anyPlatf = FALSE,
  silent = TRUE,
  callFrom = NULL  
)
```

**Description**

This function corrects paths character strings for mixed slash and backslash in file path. In Windows the function `tempdir()` will use double backslashes as separator while `file.path()` uses regular slashes. So when combining these two one might encounter a mix of slashes and double backslashes which may cause trouble, unless this is straightened out to a single separator used. When pointing to given files inside html-files, paths need to have a prefix, this can be added using the argument `asHtml`.

**Usage**

```
correctWinPath(  
x,
  asHtml = FALSE,
  anyPlatf = FALSE,
  silent = TRUE,
  callFrom = NULL  
)
```

**Arguments**

- `x` (character) input path to test and correct
- `asHtml` (logical) option for use in html: add prefix "file:/"
- `anyPlatf` (logical) if TRUE, checking will only be performed in Windows environment
- `silent` (logical) suppress messages
- `callFrom` (character) allows easier tracking of message(s) produced

**Value**

character vector with corrected path

**See Also**

`tempfile`, `file.path`

**Examples**

```
path1 <- 'D:\temp\Rtmp6X8/working_dir\RtmpKC/example.txt'
(path1b <- correctWinPath(path1, anyPlatf=TRUE))
(path1h <- correctWinPath(path1, anyPlatf=TRUE, asHtml=TRUE))
```
**countCloseToLimits**

**Count from two vectors number of values close within given limits**

**Description**

This function summarizes the search of similar (or identical) numeric values from 2 initial vectors, it evaluates the result from initial search run by `findCloseMatch()`, whose output is a less convenient list. `countCloseToLimits` checks furthermore how many results within additional (more stringent) distance-limits may be found and returns the number of distance values within the limits tested. Designed for checking if threshold used with `findCloseMatch()` may be set more stringent, eg when searching reasonable FDR limits ...

**Usage**

```r
countCloseToLimits(closeMatch, limitIdent = 5, prefix = "lim_")
```

**Arguments**

- `closeMatch` (list) output from `findCloseMatch()`, ie list indicating which instances of 2 series of data have close matches
- `limitIdent` (numeric) max limit or panel of threshold values to test (if single value, in addition a panel with values below will be tested)
- `prefix` (character) prefix for names of output

**Value**

integer vector with counts for number of list-elements with at least one absolute value below threshold, names

**See Also**

`findCloseMatch`

**Examples**

```r
set.seed(2019); aa <- sample(12:15,20,repl=TRUE) + round(runif(20),2)-0.5
bb <- 11:18
match1 <- findCloseMatch(aa,bb,com="diff",lim=0.65)
head(match1)
(tmp3 <- countCloseToLimits(match1,lim=c(0.5,0.35,0.2)))
(tmp4 <- countCloseToLimits(match1,lim=0.7))
```
countSameStartEnd  

Count same start- and end- sites of edges (or fragments)

Description

Suppose a parent sequence/string 'ABCDE' gets cut in various fragments (eg 'ABC','AB' ...). countSameStartEnd counts how many (ie re-occuring) start- and end- sites of edges do occur in the input-data. The input is presented as matrix of/indicating start- and end-sites of edges. The function is used to characterize partially redundant edges and accumulation of cutting/breakage sites.

Usage

countSameStartEnd(frag, minFreq = 2, nDig = 4)

Arguments

- **frag**  
  (matrix) 1st column *beg* start-sites, 2nd column *end* end-sites of edges, row-names to precise fragment identities are recommended

- **minFreq**  
  (integer) min number of accumulated sites for taking into account (allows filtering with large datasets)

- **nDig**  
  (integer) rounding: number of digits for columns *beg.rat* and *end.rat* in output

Value

matrix of 6 columns: input (beg and end), beg.n, beg.rat, end.n, end.rat

See Also

to build initial tree **buildTree, contribToContigPerFrag, simpleFragFig**

Examples

```r
frag1 <- cbind(beg=c(2,3,7,13,13,15,7,9,7, 3,3,5), end=c(6,12,18,20,20,19,12,12, 12,4,5,7))
rownames(frag1) <- letters[1:nrow(frag1)]
countSameStartEnd(frag1)
simpleFragFig(frag1)
```
cutArrayInCluLike  
Cut 3-dim array in list of matrixes (or arrays) similar to organizing into clusters

Description

cutArrayInCluLike cuts 'dat' (matrix, data.frame or 3-dim array) in list (of appended lines) according to 'cluOrg', which serves as instruction which line of 'dat' should be placed in which list-element (like sorting according to cluster-numbers).

Usage

cutArrayInCluLike(dat, cluOrg, silent = FALSE, debug = FALSE, callFrom = NULL)

Arguments

dat array (3 dim)
cluOrg (factor) organization of lines to clusters
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of message(s) produced

Value

This function returns a list of matrixes (or arrays)

Examples

mat1 <- matrix(1:30,nc=3,dimnames=list(letters[1:10],1:3))
cutArrayInCluLike(mat1,cluOrg=factor(c(2,rep(1:4,2),5)))

cutAtMultSites  
Cut character-vector at multiple sites

Description

This function cuts character vector after 'cutAt' (ie keep the search substring 'cutAt', different to strsplit). Used for theoretical enzymatic digestion (eg in proteomics)

Usage

cutAtMultSites(y, cutAt)
Arguments

  y character vector (better if of length=1, otherwise one won’t know which fragment stems from which input)
  cutAt (character) search sub sting, ie 'cutting rule'

Value

  modified (ie cut) character vector

See Also

  strsplit, nFragments0, nFragments

Examples

tmp <- "MSVSTMEDSCEDLVYYVTERIIAVSFSTANEENFRSNLREVAQMLSKHGNYLLFLNLSERRPDITKLHAKVLEFGPDLDHTPALEKI"
cutAtMultSites(c(tmp,"ojioRij"),c("R","K"))

cutToNgrp

Cut numeric vector to n groups (ie convert to factor)

Description

cutToNgrp is a more elaborate version of cut for cutting a the content of a numeric vector 'x' into a given number of groups, taken from the length of 'lev'. Besides, this function provides the group borders/limits for convention use with legends.

Usage

cutToNgrp(x, lev, NAuse = FALSE, callFrom = NULL)

Arguments

  x numeric vector
  lev (character or numeric), the length of this argument tells the number of groups to be used for cutting
  NAuse (logical) include NAs as separate group
  callFrom (character) for better tracking of use of functions

Value

  list with $grouped telling which element of 'x' goes in which group and $legTxt with gourp-borders for convenient use with legends

See Also

  cut
Examples

```r
set.seed(2019); dat <- runif(30) + (1:30)/2
cutToNgrp(dat, 1:5)
plot(dat, col = (1:5)[as.numeric(cutToNgrp(dat, 1:5)$grouped)])
```

diffCombin

Compute matrix of differences for all pairwise combinations of numeric vector

Description

diffCombin returns matrix of differences (eg resulting from substitution) for all pairwise combinations of numeric vector `x`.

Usage

diffCombin(x, diagAsNA = FALSE, prefix = TRUE, silent = FALSE, callFrom = NULL)

Arguments

- `x`: numeric vector to compute differences for all combinations
- `diagAsNA`: (logical) return all self-self combinations as NA (otherwise 0)
- `prefix`: (logical) if TRUE, dimnames of output will specify orientation (prefix='from.' and 'to.')</n- `silent`: (logical) suppress messages
- `callFrom`: (character) allow easier tracking of message(s) produced

Value

numeric matrix of all pairwise differences

See Also

diff for simple differences

Examples

diffCombin(c(10, 11.1, 13.3, 16.6))
**Description**

This is a `diff()`-like function to return difference in ppm between subsequent values. Result is oriented, i.e., negative ppm value means decrease (from higher to lower value). Note that if the absolute difference remains the same, the difference in ppm will not remain the same. Any difference to NA is returned as NA, thus a single NA will result in two NAs in output (unless NA is 1st or last).

**Usage**

```r
diffPPM(dat, toPrev = FALSE, silent = FALSE, callFrom = NULL)
```

**Arguments**

- `dat` (numeric) vector for calculating difference to preceding/following value in ppm
- `toPrev` (logical) determine orientation
- `silent` (logical) suppress messages
- `callFrom` (character) allows easier tracking of messages produced

**Value**

This function returns a list with close matches of 'x' to given 'y', the numeric value depends on 'sortMatch' (if FALSE then always value of 'y' otherwise of longest of x&y)

**See Also**

- `checkSimValueInSer` and (from this package) `.compareByDiff, diff`

**Examples**

```r
aa <- c(1000.01, 1000.02, 1000.05, 1000.08, 1000.09, 1000.08)
.compareByPPM(list(aa, aa), 30, TRUE) # tabular 'long' version
diffPPM(aa)
```
elimCloseCoord

Eliminate close (overlapping) points (in x & y space)

Description

elimCloseCoord reduces number of rows in 'dat' by eliminating lines where x & y coordinates (columns of matrix 'dat' defined by 'useCol') are identical (overlay points) or very close. The stringency for 'close' values may be fine-tuned using nDig), this function uses internally firstOfRepeated.

Usage

elimCloseCoord(
  dat,
  useCol = 1:2,
  elimIdentOnly = FALSE,
  refine = 2,
  nDig = 3,
  callFrom = NULL,
  silent = FALSE
)

Arguments

dat      matrix (or data.frame) with main numeric input
useCol   (numeric) index for numeric columns of 'dat' to use/consider
elimIdentOnly   (logical) if TRUE, eliminate real duplicated points only (ie identical values only)
refine    (numeric) allows increasing stringency even further (higher 'refine' .. more lines considered equal)
nDig      (integer) number of significant digits used for rounding, if two 'similar' values are identical after this rounding the second will be eliminated.
callFrom  (character) allows easier tracking of message(s) produced
silent    (logical) suppress messages

Value

resultant matrix/data.frame

See Also

findCloseMatch, firstOfRepeated

Examples

da1 <- matrix(c(rep(0:4,5),0.01,1.1,2.04,3.07,4.5),nc=2); da1[,1] <- da1[,1]*99; head(da1)
elimCloseCoord(da1)
equLenNumber

Equal character-length number

Description

equLenNumber convert numeric entry 'x' to text, with all elements getting the same number of characters (ie by adding preceeding or tailing 0s, if needed). So far, the function cannot handle scientific annotations.

Usage

equLenNumber(x, silent = FALSE, callFrom = NULL, debug = FALSE)

Arguments

x (caracter) input vector
silent (logical) suppress messages
callFrom (character) allow easier tracking of messages produced
debug (logical) additional messages for debugging

Value

character vector formated as equal number of characters per value

See Also

sprintf

Examples

equLenNumber(c(12,-3,321))
equLenNumber(c(12,-3.3,321))

exclExtrValues

Exclude extreme values (based on distance to mean)

Description

This function aims to identify extreme values (values most distant to mean, thus potential outliers), mark them as NA or directly exclude them (depending on 'showNAs'). Note that every set of non-identical values will have at least one most extreme value. Extreme values are part of many distributions, they are not necessarily true outliers.
Usage

```r
exclExtrValues(
  dat,
  result = "val",
  CVlim = NULL,
  maxExcl = 1,
  showNA = FALSE,
  goodValues = TRUE,
  silent = FALSE,
  callFrom = NULL
)
```

Arguments

dat numeric vector, main input

result (character) may be 'val' for returning data without extreme values or 'pos' for returning position/index of extreme values

CVlim (NULL or numeric) allows to retain extreme values only if a certain CV (for all 'dat') is exceeded (to avoid calling extreme values form homogenous data-sets)

maxExcl (integer) max number of elements to exclude

showNA (logical) will display extreme values as NA

goodValues (logical) allows to display rather the good values instead of the extreme values

silent (logical) suppress messages

callFrom (character) allow easier tracking of message(s) produced

Value

numeric vector wo extremle values or index-position of extreme values

See Also

`firstOfRepLines`, `get1stOfRepeatedByCol` for treatment of matrix

Examples

```r
x <- c(rnorm(30), -6, 20)
exclExtrValues(x)
```
exponNormalize Normalize by adjusting exponent

Description
This function normalizes 'dat' by optimizing exponent function (ie dat ^exp) to fit best to 'ref' (default: average of each line of 'dat').

Usage
exponNormalize(
  dat,
  useExpon,
  dynExp = TRUE,
  nStep = 20,
  startExp = 1,
  simMeas = "cor",
  refDat = NULL,
  refGrp = NULL,
  refLines = NULL,
  rSquare = FALSE,
  silent = FALSE,
  callFrom = NULL
)

Arguments
dat matrix or data.frame of numeric data to be normalized
useExpon (numeric vector or matrix) exponent values to be tested
dynExp (logical) require 'useExpon' as 2 values (matrix), will gradually increase exponent from 1st to 2nd; may be matrix or data.frame for dynamic, in this case 1st line for exp for lowest data, 2nd line for highest
nStep (integer) number of exponent variations (steps) when testing range from-to
startExp (numeric)
simMeas (character) similarity metric to be used (so far only "cor"), if rSquare=TRUE, the r-squared will be returned
refDat (matrix or data.frame) if null average of each line from 'dat' will be used as reference in similarity measure
refGrp (factor) designing which col of 'ref' should be used with which col of 'dat' (length equal to number of cols in 'dat'). Note: 'refGrp' not yet coded optimally to extract numeric part of character vector, potential problems when all lines or cols of dat are NA
refLines (NULL or integer) optional subset of lines to be considered (only) when determining normalization factors
### extr1chan

Extract just one series, ie channel, of list of arrays

#### Description

This function was designed for handling measurements stored as list of multiple arrays, like eg compound-screens using microtiter-plates where multiple parameters ('channels') were recorded for each well (element). The elements (eg compounds screened) are typically stored in the 1st dimension of the arrays, the replicated in the secon dimension and different measure types/parameters in the 3rd channel. In order to keep the structure of individual microtiter-plates, typically each plate forms a separate array (of same dimensions) in a list. The this function allows extracting a single channel of the list of arrays (3rd dim of each array) and return row-appended matrix.

#### Usage

```
extr1chan(arrLst, cha, na.rm = TRUE, rowSep = "__")
```
**extractLast2numericParts**

**Arguments**

- `arrLst` (list) list of arrays (typically 1st and 2nd dim for specific genes/objects, 3rd for different measures associated with)
- `cha` (integer) channel number
- `na.rm` (logical) default =TRUE to remove NAs
- `rowSep` (character) separator for rows

**Value**

list with just single channel extracted

**See Also**

organizeAsListOfRep1

**Examples**

```r
arr1 <- array(1:24,dim=c(4,3,2),dimnames=list(c(LETTERS[1:4]),
paste("col",1:3,sep=""),c("ch1","ch2")))
arr2 <- array(74:51,dim=c(4,3,2),dimnames=list(c(LETTERS[1:4]),
paste("col",1:3,sep=""),c("ch1","ch2")))
arrL1 <- list(pl1=arr1,pl2=arr2)
extr1chan(arrL1,ch=2)
```

---

**extractLast2numericParts**

*Extract last two numeric parts from character vector*

**Description**

`extractLast2numericParts` extracts last 2 (integer) numeric parts between punctuations out of character vector 'x'. Runs faster than `gregexpr`. Note: won’t work correctly with decimals or exponential signs !! (such characters will be considered as punctuation, ie as separator)

**Usage**

```r
extractLast2numericParts(x, silent = FALSE, callFrom = NULL)
```

**Arguments**

- `x` main character input
- `silent` (logical) suppres messages
- `callFrom` (character) allow easier tracking of message(s) produced

**Value**

(numeric) matrix with 2 columns (eg from initial concatenated coordinates)
See Also
gregexpr from grep

Examples
extractLast2numericParts(c("M01.1-4","M001/2.5","M_0001_03-16","zyx","012","a1.b2.3-7,2"))

dFr <- data.frame(a=11:14, b=24:21, cc=LETTERS[1:4], dd=rep(c(TRUE,FALSE),2))
extrColsDeX(dFr, c("b","cc","notThere"))
extrColsDeX(dFr, c("b","cc","notThere"), doExtractCols=TRUE)
extrColsDeX(dFr, list(c("nn","b","a"), c("cc","a"),"notThere"))
extrNumericFromMatr  

*Extract numeric part of matrix or data.frame*

Description

extrNumericFromMatr extracts numeric part of matrix or data.frame, removing remaining non-numeric elements if trimToData is set to TRUE. Note, that cropping entire lines where a (single) text element appeared may quickly reduce the overall content of the input data.

Usage

```r
extrNumericFromMatr(dat, trimToData = TRUE, silent = FALSE, callFrom = NULL)
```

Arguments

- `dat`: matrix (or data.frame) for extracting numeric parts
- `trimToData`: (logical) default to remove (crop) lines and cols contributing to NA, non-numeric data is transformed to NA
- `silent`: (logical) suppress messages
- `callFrom`: (character) allow easier tracking of message(s) produced

Value

matrix of numeric data

Examples

```r
mat <- matrix(c(letters[1:7],14:16,LETTERS[1:6]),nrow=4,dimnames=list(1:4,letters[1:4]))
mat; extrNumericFromMatr(mat)
mat <- matrix(c(letters[1:4],1,"e",12:19,LETTERS[1:6]),nr=5,dimnames=list(11:15,letters[1:4]))
mat; extrNumericFromMatr(mat)
```

extrSpcText  

*Extract specific text*

Description

This function extracts/cuts text-fragments out of `txt` following specific anchors defined by arguments `cutFrom` and `cutTo`. 
Usage

extrSpcText(  
txt,  
cutFrom = " GN=",  
cutTo = " PE=" ,  
missingAs = NA,  
exclFromTag = TRUE,  
silent = FALSE,  
debug = FALSE,  
callFrom = NULL  }
)

Arguments

txt character vector to be treated

cutFrom (character) text where to start cutting

cutTo (character) text where to stop cutting

missingAs (character) specific content of output at line/location of 'exclLi'

exclFromTag (logical) to exclude text given in 'cutFrom' from result

silent (logical) suppress messages

debug (logical) additional messages for debugging

callFrom (character) allow easier tracking of messages produced

Details

In case cutFrom is not found missingAs will be returned. In case cutTo is not found, text gets extracted with chaMaxEl characters.

Value

This function returns a modified character vector

See Also

substr

Examples

eextrSpcText(c(" ghjg GN=thisText PE=001"," GN=_ PE=" , NA, "abcd"))
eextrSpcText(c("ABCDEF.3-6","05g","bc.4-5"), cutFr="\." , cutT="-")
**filt3dimArr**

Filter three-dimensional array of numeric data

**Description**
Filtering of matrix or (3-dim) array x: filter column according to `filtCrit` (eg 'inf') and threshold `filtVal`

**Usage**
```
filt3dimArr(
  x, filtVal, filtTy = ">", filtCrit = NULL, displCrit = NULL, silent = FALSE, debug = FALSE, callFrom = NULL
)
```

**Arguments**
- `x`: array (3-dim) of numeric data
- `filtVal`: (numeric, length=1) for testing inferior/superior/equal condition
- `filtTy`: (character, length=1) which type of testing to perform (may be 'eq', 'inf', 'infeq', 'sup', 'supeq', '>', '<', '>=', '<=', '==')
- `filtCrit`: (character, length=1) which column-name consider when filtering filter with 'filtVal' and 'filtTy'
- `displCrit`: (character) column-name(s) to display
- `silent`: (logical) suppress messages
- `debug`: (logical) additional messages for debugging
- `callFrom`: (character) allow easier tracking of messages produced

**Details**
and extract/display all col matching 'displCrit'.

**Value**
This function returns a list of filtered matrixes (by 3rd dim)

**See Also**
`filterList`; `filterLiColDeList`;
Examples

arr1 <- array(11:34, dim=c(4,3,2), dimnames=list(c(LETTERS[1:4]), paste("col",1:3,sep=""), c("ch1","ch2"))
filt3dimArr(arr1,disp1Crit=c("col1","col2"),filtCrit="col2",filtVal=7)

filterLiColDeList

Description

Filter all elements of list (or S3-object) according to criteria designed to one selected reference-element of the list. All simple vectors, matrix, data.frames and 3-dimensional arrays will be checked if matching number of rows and/or columns to decide if they should be filtered the same way. If the reference element has same number of rows and columns simple (1-dimensional) vectors won’t be filtered since it not clear if this should be done to lines or columns.

Usage

filterLiColDeList(
  lst,
  useLines,
  useCols = NULL,
  ref = 1,
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

lst (list or S3 object) main input
useLines (integer, logical or character) vector to assign lines to keep when filtering along lines; set to NULL for no filtering; if 'allNA' all lines composed uniquely of NA values will be removed.
useCols (integer, logical or character) vector for filtering columns; set to NULL for no filtering; if 'allNA' all columns uniquely NA values will be removed
ref (integer) index for designing the element of 'lst' to take as reference for checking which other list-elements have suitable number of rows or columns
silent (logical) suppress messages
callFrom (character) allow easier tracking of messages produced
debug (logical) additional messages for debugging

Details

This function is used eg in package wrProteo to simultaneously filter raw and transformed data.
filterList

Value
This function returns the correct(ed) input (object of same class, of same length)

See Also
moderTest2grp for single comparisons, lmFit

Examples

```r
lst1 <- list(m1=matrix(11:18,ncol=2), m2=matrix(21:30,ncol=2), indR=31:34,
m3=matrix(c(21:23,NA,25:27,NA),ncol=2))
## here $m2 has more lines than $m1, and thus will be ignored when ref=1
filterLiColDeList(lst1, useLines=2:3)
filterLiColDeList(lst1, useLines="allNA", ref=4)
```

filterList

Filter for unique elements

Description
This function aims to apply a given filter-criterium, a matrix or vector of FALSE/TRUE which is typically combined with a second layer which filters for a min content of filer-passing values per line for the first/main criterium. Then all lines concerned will be removed. This will be done for all list-elements (of appropriate size) of the input-list (while maintaining the list-structure in the output) not matching the filtering criteria.

Usage

```r
filterList(lst, filt, minLineRatio = 0.5, silent = FALSE, callFrom = NULL)
```

Arguments

- **lst** (list) main input, each vector, matrix or data.frame in this list will be filtered if its length or number of lines fits to filt
- **filt** (logical) vector of FALSE/TRUE to use for filtering. If this a matrix is given, the value of minLineRatio will be applied as threshod of min content of TRUE for each line of filt
- **minLineRatio** (numeric) in case filt is a matrix of FALSE/TRUE, this value will be used as threshold of min content of TRUE for each line of filt
- **silent** (logical) suppress messages
- **callFrom** (character) allow easier tracking of message(s) produced

Value
filtered list
See Also
correctToUnique, unique, duplicated, extrColsDeX

Examples

```r
set.seed(2020); dat1 <- round(runif(80),2)
list1 <- list(m1=matrix(dat1[1:40],ncol=8), m2=matrix(dat1[41:80],ncol=8), other=letters[1:8])
rownames(list1$m1) <- rownames(list1$m2) <- paste0("line",1:5)
filterList(list1, list1$m1[1,] >0.4)
filterList(list1, list1$m1 >0.4)
```

filterNetw

Filter nodes & edges for extracting networks This function allows extracting and filtering network-data based on fixed threshold (limInt) and add sandwich-nodes (nodes inter-connecting initial nodes) out of node-based queries.

Description

Filter nodes & edges for extracting networks

This function allows extracting and filtering network-data based on fixed threshold (limInt) and add sandwich-nodes (nodes inter-connecting initial nodes) out of node-based queries.

Usage

```r
filterNetw(
  lst,  
  filtCol = 3,  
  limInt = 5000,  
  sandwLim = 5000,  
  filterAsInf = TRUE,  
  outFormat = "matrix",  
  remOrphans = TRUE,  
  remRevPairs = TRUE,  
  elemNa = "genes",  
  silent = FALSE,  
  callFrom = NULL,  
  debug = FALSE  
)
```

Arguments

- `lst` (list, composed of multiple matrix or data.frames ) main input (each list-element should have same number of columns)
- `filtCol` (integer, length=1) which column of lst should be used to filter using thresholds limInt and sandwLim
filterNetw

limInt (numeric, length=1) filter main edge-scores according to filterAsInf
sandwLim (numeric, length=1) filter sandwich connection edge-scores according to filterAsInf
filterAsInf (logical) filter as 'inferior or equal' or 'superior or equal'
outFormat (character) may be 'matrix' for tabular output, 'all' as list with matrix and list of node-names
remOrphans (logical) remove networks consisting only of 2 connected edges
remRevPairs (logical) remove duplicate edges due to reverse massping (eg A - B and B - A); NOTE: use only when edges don’t have orientation!
elemNa (character) used only for messages
silent (logical) suppress messages
callFrom (character) allow easier tracking of message(s) produced
debug (logical) display additional messages for debugging

Value
This function returns a matrix or data.frame

See Also
in cbind

Examples

```r
lst2 <- list('121'=data.frame(ID=as.character(c(141,221,228,229,449)),11:15),
              '131'=data.frame(ID=as.character(c(228,331,332,333,339)),11:15),
              '141'=data.frame(ID=as.character(c(121,151,229,339,441,442,449)),c(11:17)),
              '151'=data.frame(ID=as.character(c(449,141,551,552)),11:14),
              '161'=data.frame(ID=as.character(171),11), '171'=data.frame(ID=as.character(161),11),
              '181'=data.frame(ID=as.character(881:882),11:12))

lst2 <- list('121'=data.frame(ID=as.character(c(141,221,228,229,449)),11:15,21:25),
              '131'=data.frame(ID=as.character(c(228,331,332,333,339)),11:15,21:25),
              '141'=data.frame(ID=as.character(c(121,151,229,339,441,442,449)),c(11:17),21:27),
              '151'=data.frame(ID=as.character(c(449,141,551,552)),11:14,21:24),
              '161'=data.frame(ID=as.character(171),11,21), '171'=data.frame(ID=as.character(161),11,21),
              '181'=data.frame(ID=as.character(881:882),11:12,21:22))

(te1 <- filterNetw(lst2, limInt=90, remOrphans=FALSE))
(te2 <- filterNetw(lst2, limInt=90, remOrphans=TRUE))
(te3 <- filterNetw(lst2, limInt=13, remOrphans=FALSE))
(te4 <- filterNetw(lst2, limInt=13, remOrphans=TRUE))
```
filtSizeUniq  
*Filter for unique elements*

**Description**

This function aims to identify and remove duplicated elements in a list and maintain the list-structure in the output. `filtSizeUniq` filters 'lst' (list of character-vectors or character-vector) for elements being unique (to 'ref' or if NULL to all 'lst') and of character length. In addition, the min- and max- character length may be filtered, too. Eg, in proteomics this helps removing peptide sequences which would not be measured/detected any way.

**Usage**

```
filtSizeUniq(
  lst, 
  ref = NULL, 
  minSize = 6, 
  maxSize = 36, 
  filtUnique = TRUE, 
  byProt = TRUE, 
  inclEmpty = TRUE, 
  silent = FALSE, 
  debug = FALSE, 
  callFrom = NULL
)
```

**Arguments**

- **lst**: list of character-vectors or character-vector
- **ref**: (character) optional alternative 'reference', if not NULL used in addition to 'lst' for considering elements of 'lst' as unique
- **minSize**: (integer) minimum number of characters, if NULL set to 0
- **maxSize**: (integer) maximum number of characters
- **filtUnique**: (logical) if TRUE return unique-only character-strings
- **byProt**: (logical) if TRUE organize output as list (by names of input, eg protein-names) - if 'lst' was named list
- **inclEmpty**: (logical) optional including empty list-elements when all elements have been filtered away - if 'lst' was named list
- **silent**: (logical) suppress messages
- **debug**: (logical) additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

**Value**

list of filtered input
findCloseMatch

See Also
correctToUnique, unique, duplicated

Examples

```r
filtSizeUniq(list(A="a",B=c("b","bb","c"),D=c("dd","d","ddd","c")),filtUn=TRUE,minSi=NULL)
# input: c and dd are repeated
filtSizeUniq(list(A="a",B=c("b","bb","c"),D=c("dd","d","ddd","c")),ref=c(letters[c(1:26,1:3)],
"dd","dd","bb","ddd"),filtUn=TRUE,minSi=NULL)  # a,b,c,dd repeated
```

Description

findCloseMatch finds close matches (similar values) between two numeric vectors ('x', 'y') based on method 'compTy' and threshold 'limit'. Return list with close matches of 'x' to given 'y', the numeric value depends on 'sortMatch' (if FALSE then always value of 'y' otherwise of longest of x&y). Note: Speed & memory improvement if 'sortMatch'=TRUE (but result might be inversed!): adopt search of x->y or y->x to searching matches of each longest to each shorter (ie flip x & y). Otherwise, if length of 'x' & 'y' are very different, it may be advantageous to use a long(er) 'x' and short(er) 'y' (with 'sortMatch'=FALSE). Note: Names of 'x' & 'y' or (if no names) prefix letters 'x' & 'y' are always added as names to results.

Usage

```r
findCloseMatch(
  x,
  y,
  compTy = "ppm",
  limit = 5,
  asIndex = FALSE,
  maxFitShort = 100,
  sortMatch = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- **x**: numeric vector for comparison
- **y**: numeric vector for comparison
- **compTy**: (character) may be 'diff' or 'ppm', will be used with threshold from argument 'limit'
- **limit**: (numeric) threshold value for retaining values, used with distance-type specified in argument 'compTy'
asIndex  (logical) optionally rather report index of retained values
maxFitShort  (numeric) limit output to max number of elements (avoid returning high number of results if filtering was not enough stringent)
sortMatch  (logical) if TRUE than matching will be preformed as 'match longer (of x & y) to closer', this may process slightly faster (eg 'x' longer: list for each 'y' all 'x' that are close, otherwise list of each 'x'),
silent  (logical) suppress messages
debug  (logical) display additional messages for debugging
callFrom  (character) allow easier tracking of message(s) produced

Value

This function returns a list with close matches of 'x' to given 'y', the numeric value dependes on 'sortMatch' (if FASLE then always value of 'y' otherwise of longest of x&y)

See Also

checkSimValueInSer and (from this package) .compareByDiff, for convient output countCloseToLimits

Examples

findCloseMatch(aa,bb,com="diff",lim=0.6)

findCloseMatch(c(a=5,b=11,c=12,d=18),c(G=2,H=11,I=12,J=13)+0.5, comp="diff", lim=2)
findCloseMatch(c(4,5,11,12,18),c(2,11,12,13,33)+0.5, comp="diff", lim=2)
findCloseMatch(c(4,5,11,12,18),c(2,11,12,13,33)+0.5, comp="diff", lim=2, sort=FALSE)

findCloseMatch(aa,bb,com="diff",lim=0.6)

findCloseMatch(c(a=5,b=11,c=12,d=18),c(G=2,H=11,I=12,J=13)+0.5, comp="diff", lim=2)
findCloseMatch(c(4,5,11,12,18),c(2,11,12,13,33)+0.5, comp="diff", lim=2)
findCloseMatch(c(4,5,11,12,18),c(2,11,12,13,33)+0.5, comp="diff", lim=2, sort=FALSE)

a2 <- c(11:20); names(a2) <- letters[11:20]
b2 <- c(25:5)+c(rep(0,5),(1:10)/50000,rep(0,6)); names(b2) <- LETTERS[25:5]
which(abs(b2-a2[8]) < a2[8]*1e-6*5) # find R=18 : no10
findCloseMatch(a2, b2, com="ppm", lim=5)  # find Q,R,S,T
findCloseMatch(a2, b2, com="ppm", lim=5,asI=TRUE)  # find Q,R,S,T
findCloseMatch(b2, a2, com="ppm", lim=5,asI=TRUE,sort=FALSE)
findCloseMatch(a2, b2, com="ratio", lim=1.000005) # find Q,R,S,T
findCloseMatch(a2, b2, com="diff", lim=0.000005)  # find S,T

findRepeated  Find repeated elements

Description

findRepeated gets index of repeated items/values in vector 'x' (will be treated as character). Return (named) list of indexes for each of the repeated values, or NULL if all values are unique. This approach is similar but more basic compared to get1stOfRepeatedByCol.
findSimilFrom2sets

Usage

findRepeated(x, nonRepeated = FALSE, silent = FALSE, callFrom = NULL)

Arguments

x character vector

nonRepeated (logical) if =TRUE, return list with elements $rep and $nonrep

silent (logical) suppress messages

callFrom (character) allow easier tracking of message(s) produced

Value

(named) list of indexes for each of the repeated values, or NULL if all values unique

See Also

similar approach but more basic than get1stOfRepeatedByCol

Examples

aa <- c(11:16,14:12,14); findRepeated(aa)

findSimilFrom2sets Find similar numeric values from two vectors/matrixes

Description

findSimilFrom2sets compares to vectors or matrixes and returns combined view including only all close (by findCloseMatch). Return matrix (predMatr) with add'l columns for index to and 'grp' (group of similar values (1-to-many)), 'nGrp' (n of grp), 'isBest' or 'nBest', 'disToMeas' (distance/difference between pair) & 'ppmToPred' (distance in ppm). Note: too wide 'limitComp' will result in large window and many 'good' hits will compete (and be mutually excluded) if selection 'bestOnly' is selected

Usage

findSimilFrom2sets(
    predMatr,
    measMatr,
    colMeas = 1,
    colPre = 1,
    compareTy = "diff",
    limitComp = 0.5,
    bestOnly = FALSE,
    silent = FALSE,
    callFrom = NULL,
    debug = FALSE
)
Arguments

- **predMatr**: (matrix or numeric vector) dataset number 1, referred to as 'predicted', the column specified in argument `colPre` points to the data to be used.
- **measMatr**: (matrix or numeric vector) dataset number 2, referred to as 'measured', the column specified in argument `colMeas` points to the data to be used.
- **colMeas**: (integer) which column number of 'measMatr' to consider.
- **colPre**: (integer) which column number of 'predMatr' to consider.
- **compareTy**: (character) 'diff' (difference) 'ppm' (relative difference).
- **limitComp**: (numeric) limit used by 'compareTy'.
- **bestOnly**: (logical) allows to filter only hits with min distance (defined by 'compareTy'), 3rd last col will be 'nBest' - otherwise 3rd last col 'isBest'.
- **silent**: (logical) suppress messages.
- **callFrom**: (character) allow easier tracking of messages produced.
- **debug**: (logical) for bug-tracking: more/enhanced messages.

Value

This function returns a matrix (predMatr) with add'l columns for index to and 'grp' (group of similar values (1-to-many)), 'nGrp' (n of grp), 'isBest' or 'nBest', 'disToMeas' (distance/difference between pair) & 'ppmToPred' (distance in ppm).

See Also

- checkSimValueInSer
- findCloseMatch
- closeMatchMatrix

Examples

```r
aA <- c(11:17); bB <- c(12.001,13.999); cC <- c(16.2,8,9,12.5,12.6,15.9,14.1)
aZ <- matrix(c(aA,aA+20),ncol=2,dimnames=list(letters[1:length(aA)],c("aaA","aZ"))
cZ <- matrix(c(cC,cC+20),ncol=2,dimnames=list(letters[1:length(cC)],c("ccC","cZ"))
findCloseMatch(cC,aA,com="diff",lim=0.5,sor=FALSE)
findSimilFrom2sets(aA,cC)
findSimilFrom2sets(cC,aA)
findSimilFrom2sets(aA,cC,best=FALSE)
findSimilFrom2sets(aA,cC,comp="ppm",lim=5e4,deb=TRUE)
findSimilFrom2sets(aA,cC,comp="ppm",lim=9e4,bestO=FALSE)
# below: find fewer 'best matches' since search window larger (ie more good hits compete !)
findSimilFrom2sets(aA,cC,comp="ppm",lim=9e4,bestO=TRUE)
```
findUsableGroupRange  Select groups within given range

Description
This function aims to help finding stretches/segments of data with a given maximum number of NA-instances. This function is used to inspect/filter each line of 'dat' for a subset with sufficient presence/absence of NA values (ie limit number of NAs per level of 'grp'). Note: optimal performance with n.lines > n.groups

Usage
findUsableGroupRange(dat, grp, maxNA = 1, callFrom = NULL)

Arguments
- dat: (matrix or data.frame) main input
- grp: (factor) information which column of 'dat' is replicate of whom
- maxNA: (integer) max number of tolerated NAs
- callFrom: (character) allow easier tracking of message(s) produced

Value
matrix with boundaries of 1st and last usable column (NA if there were no suitable groups found)

Examples
```r
dat1 <- matrix(1:56,nc=7)
dat1[c(2,3,4,5,6,10,12,18,19,20,22,23,26,27,28,30,31,34,38,39,50,54)] <- NA
rownames(dat1) <- letters[1:nrow(dat1)]
findUsableGroupRange(dat1,gl(3,3)[-c(3:4)])
```

firstLineOfDat  Filter matrix to keep only first of repeated lines

Description
This function aims to reduce the complexity of a matrix (or data.frame) in case column 'refCol' has multiple lines with same value. In this case, it reduces the input-data to 1st line of redundant entries and returns a matrix (or data.frame) without lines identified as redundant entries for 'refCol'). In sum, this functions works like using unique on a given column, and propagates the same treatment to all other columns.

Usage
firstLineOfDat(dat, refCol = 2, silent = FALSE, debug = FALSE, callFrom = NULL)
Arguments

- **dat** (matrix or data.frame) main input
- **refCol** (integer) column number of reference-column
- **silent** (logical) suppress messages
- **debug** (logical) additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced

Value

matrix (same number of columns as input)

See Also

- `firstOfRepeated`, `unique`, `duplicated`

Examples

```r
(mat1 <- matrix(c(1:6,rep(1:3,1:3)),ncol=2,dimnames=list(letters[1:6],LETTERS[1:2])))
firstLineOfDat(mat1)
```

---

### Description

This function works similar to `unique`, but provides additional information about which elements of original input 'x' are repeated by providing indexes realtoe to the input. `firstOfRepeated` makes list with 3 elements: $indRepeated.. index for first of repeated 'x', $indUniq.. index of all unique + first of repeated, $indRedund.. index of all redundant entries, ie non-unique (wo 1st). Used for reducing data to non-redundant status, however, for large numeric input the function `nonAmbiguousNum()` may perform better/faster. NAs won't be considered (NAs do not appear in reported index of results), see also `firstOfRepLines()`.

Usage

```r
firstOfRepeated(x, silent = FALSE, debug = FALSE, callFrom = NULL)
```

Arguments

- **x** (character or numeric) main input
- **silent** (logical) suppress messages
- **debug** (logical) display additional messages for debugging
- **callFrom** (character) allow easier tracking of message(s) produced
**Value**

list with indices: $\text{indRepeated}$, $\text{indUniq}$, $\text{indRedund}$

**See Also**

duplicated, nonAmbiguousNum, firstOfRepLines gives less detail in output (lines/elements/indexes of omitted not directly accessible) and works faster

**Examples**

```r
x <- c(letters[3,2:4,8,NA,3:1,NA,5:4]); names(x) <- 100+(1:length(x))
firstOfRepeated(x)
x[firstOfRepeated(x)$indUniq]  # only unique with names
```

---

**firstOfRepLines**

*Reduce to first occurrence of repeated lines*

**Description**

This function concatenates all columns of input-matrix and then searches like unique for unique elements, optionally the indexes of unique elements may get returned. Note: This function treats input as character (thus won’t understand `10==10.0`). Returns simplified/non-redundant vector/matrix (ie fewer lines), or respective index. faster than firstOfRepeated

**Usage**

```r
firstOfRepLines(
  mat,
  outTy = "ind",
  useCol = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- **mat** initial matrix to treat
- **outTy** for output type: 'ind'.. index to 1st occurance (non-red),'orig'..non-red lines of mat, 'conc'.. non-red concatenated values, 'num'.. index to which group/category the lines belong
- **useCol** (integer) custom choice of which columns to paste/concatenate
- **silent** (logical) suppress messages
- **debug** (logical) additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced
fuseAnnotMatr

Value
simplified/non-redundant vector/matrix (ie fewer lines for matrix), or respective index

See Also
unique, nonAmbiguousNum, faster than firstOfRepeated which gives more detail in output (lines/elements/indexes of omitted)

Examples
mat <- matrix(c("e","n","a","n","z","z","n","z","b","n","c","n","n","n","n","n","z"),ncol=2)
firstOfReplLines(mat,out="conc")

fuseAnnotMatr
Fuse annotation matrix to initial matrix

Description
In a number of instances experimental measurements and additional information (annotation) are provided by separate objects (matrixes) as they may not be generated the same time. The aim of this function is provide help when matching approriate lines for 2 sets of data (experimental measures in iniTab and annotation from annotTab) for fusing. fuseAnnotMatr adds suppelmental columns/annotation to an initial matrix iniTab : using column 'refIniT' as key (in iniTab) to compare with key 'refAnnotT' (from 'annotTab'). The columns to be added from annotTab must be chosen explicitely. Note: if non-unique IDs in iniTab : runs slow (but save) due to use of loop for each unique ID.

Usage
fuseAnnotMatr(
iniTab,
annotTab,
refIniT = "Uniprot",
refAnnotT = "combName",
addCol = c("ensembl_gene_id", "description", "geneName", "combName"),
dump = TRUE,
silent = FALSE,
callFrom = NULL
)

Arguments
iniTab (matrix), that may have lines with multiple (=repeated) key entries
annotTab (matrix) containing reference annotation
refIniT (character) type of reference (eg 'Uniprot')
refAnnotT (character) column name to use for reference-annotation
Fuse content of list-elements with redundant (duplicated) names

FuseCommonListElem fuses (character or numeric) elements of list re-occurring under same name, so that resultant list has unique names. Note: will not work with list of matrixes

Usage

FuseCommonListElem(
  lst,
  initOrd = TRUE,
  removeDuplicates = FALSE,
  callFrom = NULL
)

Arguments

lst (list) main input, list of numeric vectors
initOrd (logical) preserve initial order in output (if TRUE) or otherwise sort alphabetically

Examples

tab0 <- matrix(rep(letters[1:25], 8), ncol = 10)
tab1 <- cbind(Uniprot = paste(tab0[, 1], tab0[, 2]), col1 = paste(tab0[, 3],
  tab0[, 4], tab0[, 5], " ", tab0[, 7], tab0[, 6]))
tab2 <- cbind(combName = paste(tab0[, 1], tab0[, 2]), col2 = paste(tab0[, 8], tab0[, 9], tab0[, 10]))
fuseAnnotMatr(tab1, tab2[c(20:11, 2:5),], refIni = "Uniprot", refAnnotT = "combName", addCol = "col2")
fuseAnnotMatr(tab2[c(20:11, 2:5),], tab1, refAnnotT = "Uniprot", refIni = "combName", addCol = "col1")
removeDuplicates

(logical) allow to remove duplicate entries (if vector contains names, both the name and the value need to be identical to be removed; note: all names must have names with more than 0 characters to be considered as names)

callFrom

(character) allows easier tracking of message(s) produced

Value

fused list (same names as elements of input)

See Also

unlist

Examples

val1 <- 1:10:26
names(val1) <- letters
lst1 <- list(c=val1[3:6],a=val1[1:3],b=val1[2:3],a=val1[12],c=val1[13])
fuseCommonListElem(lst1)

Description

Fuse previously identified pairs to 'clusters', return vector with cluster-numbers.

Usage

fusePairs(
  datPair,
  refDatNames = NULL,
  inclRepLst = FALSE,
  maxFuse = NULL,
  debug = FALSE,
  silent = TRUE,
  callFrom = NULL
)

Arguments

datPair 2-column matrix where each line represents 1 pair

refDatNames (NULL or character) allows placing selected pairs in context of larger data-set (names to match those of 'datPair')

inclRepLst (logical) if TRUE, return list with 'clu' (clu-numbers, default output) and 'reflst' (list of clustered elements, only n>1)
maxFuse (integer, default NULL) maximal number of groups/clusters
debug (logical) display additional messages for debugging
silent (logical) suppress messages
callFrom (character) allow easier tracking of message(s) produced

Value
This function returns a vector with cluster-numbers

Examples

daPa <- matrix(c(1:5,8,2:6,9), ncol=2)
fusePairs(daPa, maxFuse=4)

get1stOfRepeatedByCol

Description
get1stOfRepeatedByCol sorts matrix 'mat' and extracts only 1st occurance of values in column 'sortBy'. Returns then non-redundant matrix (ie for column 'sortBy', if 'markIfAmbig' specifies existing col, mark ambig there). Note : problem when sortSupl or sortBy not present (or not intended for use)

Usage
get1stOfRepeatedByCol(
  mat,
  sortBy = "seq",
  sortSupl = "ty",
  asFirstLast = c("full", "inter"),
  markIfAmbig = c("ambig", "seqNa"),
  asList = FALSE,
  abmiPref = "_"
)

Arguments
mat (matrix or data.frame) numeric vector to be tested
sortBy column name for which elements should be made unique, numeric or character column; 'sortSupl' .. add'l colname to always select specific 1st)
sortSupl default="ty"
asFirstLast (character,length=2) to force specific strings from colunm 'sortSupl' as first and last when selecting 1st of repeated terms, default=c("full","inter")
markIfAmbig (character,length=2) 1st will be set to 'TRUE' if ambiguous/repeated, 2nd will get (heading) prefix, default=c("ambig","seqNa")
getValuesByUnique

asList (logical) to return list with non-redundant ('unique') and removed lines ('repeats')

abmiPref (character) prefix to note ambiguous entries/terms, default="_"

Value
depending on 'asList' either list with non-redundant ('unique') and removed lines ('repeats')

See Also

firstOfRepeated for (more basic) treatment of simple vector, nonAmbiguousNum for numeric use
(much faster !!!)

Examples

aa <- cbind(no=as.character(1:20),seq=sample(LETTERS[1:15],20,repl=TRUE),
ty=sample(c("full","Nter","inter"),20,repl=TRUE),ambig=rep(NA,20),seqNa=1:20)
get1stOfRepeatedByCol(aa)

description

When data have repeated elements (defined by names inside the vector), it may be advantageous to
run some operations only on a unique set of the initial data, or sometimes all repeated occurrences
need to be replaced by a common (summarizing) value. This function allows to re-introduce new
values from a second vector with unique names, to return a final vector of initial input-length and
order of names (elements) like initial, too. Normally the user would provide 'datUniq' (without
repeated names) containing new values which will be expanded to structure of 'dat', if 'datUniq' is
not provided a vector with unique names will be made using the first occurrence of repeated value(s).
For more complex cases the indexing relative to 'datUniq' can be returned (setting asIndex=TRUE).
Note: If not all names of 'dat' are found in 'datUniq' the missing spots will be returned as NA.

Usage

getValuesByUnique(
  dat,
  datUniq = NULL,
  asIndex = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

description

When data have repeated elements (defined by names inside the vector), it may be advantageous to
run some operations only on a unique set of the initial data, or sometimes all repeated occurrences
need to be replaced by a common (summarizing) value. This function allows to re-introduce new
values from a second vector with unique names, to return a final vector of initial input-length and
order of names (elements) like initial, too. Normally the user would provide 'datUniq' (without
repeated names) containing new values which will be expanded to structure of 'dat', if 'datUniq' is
not provided a vector with unique names will be made using the first occurrence of repeated value(s).
For more complex cases the indexing relative to 'datUniq' can be returned (setting asIndex=TRUE).
Note: If not all names of 'dat' are found in 'datUniq' the missing spots will be returned as NA.

Usage

getValuesByUnique(
  dat,
  datUniq = NULL,
  asIndex = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
Arguments

- **dat** (numeric or character) main long input, must have names
- **datUniq** (numeric or character) will be used to impose values on dat, must have names that should match names (at least partially) from dat
- **asIndex** (logical) if TRUE index values will be returned instead of replacing values
- **silent** (logical) suppress messages
- **debug** (logical) additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced

Value

vector of length dat with imposed values, or index values if asIndex=TRUE

See Also

unique, findRepeated, correctToUnique, treatTxtDuplicates

Examples

dat <- 11:19
names(dat) <- letters[c(6:3,2:4,8,3)]
## let's make a 'datUniq' with the mean of repeated values :
datUniq <- round(tapply(dat,names(dat),mean),1)
## now propagate the mean values to the full vector
getValuesByUnique(dat,datUniq)
cbind(ini=dat,firstOfRep=getValuesByUnique(dat,datUniq),
indexUniq=getValuesByUnique(dat,datUniq,asIn=TRUE))

Description

This functions converts a given urlName so that from data from git-hub can be read correctly that tabular data. Thus, this will remove '/blob/' and change starting characters to 'raw.githubusercontent.com'

Usage

gitDataUrl(
  urlName,
  replTxt = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
htmlSpecCharConv

Arguments

urlName (character) main url-address
replTxt (NULL or matrix) adjust/custom-modify search- and replacement items; should be matrix with 2 columns, the 1st column entries will be used as 'search-for' and the 2nd as 'replace by' for each row.
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

corrected urlName

See Also

sub;

Examples

gitDataUrl(url1)

htmlSpecCharConv

Description

Converts 'txt' so that (the most common) special characters (like 'beta','micro','square' etc) will be displayed correctly when used for display in html (eg at mouse-over). Note: The package stringi is required for the conversions (the input will get returned if stringi is not available). Currently only the 16 most common special characters are implemented.

Usage

htmlSpecCharConv(txt, silent = FALSE, callFrom = NULL, debug = FALSE)

Arguments

  txt character vector, including special characters
  silent (logical) suppress messages
  callFrom (character) allow easier tracking of messages produced
  debug (logical) additional messages for debugging
Value

This function returns a corrected character vector adopted for html display.

See Also


Examples

```r
## we'll use the package stringi to generate text including the 'micro'-symbol as input
x <- if(requireNamespace("stringi", quietly=TRUE)) {
  stringi::stri_unescape_unicode("\u00b5\u003d\u0061\u0062")
} else "x=axb"
htmlSpecCharConv(x)
```

Description

This function allows recovering the single longest common text-fragments (from center, head or tail) out of character vector `txt`. Only the first of all of the longest solutions will be returned.

Usage

```r
keepCommonText(  
  txt,  
  minNchar = 1,  
  side = "center",  
  hiResol = TRUE,  
  silent = TRUE,  
  callFrom = NULL,  
  debug = FALSE  
)
```

Arguments

- `txt` character vector to be treated
- `minNchar` (integer) minimum number of characters that must remain
- `side` (character) may be either 'center', 'any', 'terminal', 'left' or 'right'; only with side='center' or 'any' internal text-segments may be found
- `hiResol` (logical) find best solution, but at much higher computational cost (eg 3x slower, however hiResol=FALSE rather finds anchor which may need to get extended)
- `silent` (logical) suppress messages
- `callFrom` (character) allow easier tracking of messages produced
- `debug` (logical) display additional messages for debugging
Details

Please note, that finding common parts between chains of characters is not a completely trivial task. This topic still has ongoing research for the application of sequence-alignments, where chains of characters to be compared get very long. This function uses a k-mer inspired approach. The initial aim with this function was allowing to treat smaller chains of characters (and finding shorter stretches of common text), like eg with column-names.

Important : This function identifies only the first best hit, ie other shared/common character-chains of the same length will not be found!

Using the argument hiResol=FALSE it is possible to accelerate the search aprox 3x (with larger character-vectors), however, frequently the very best solution may not be found. This means, that in this case the result should rather be considered a 'seed', allowing check if further extension may improve the result, ie for identifying a (slightly) longer chain of common characters.

With longer vectors and longer character chains this may get demanding on computational resources, the argument hiResol=FALSE allows reducing this at the price of missing the best solution. With this argument single common/matching characters will not be searched if all text-elements are longer than 500 characters, an empty character vector will be returned.

When argument side is either left, right or terminal only terminal common text may be found (a potentially even longer internal text will be lost). Of course, choosing this option makes searches much faster.

This function does not return the position of the shared/common characters within the text, you may use gregexpr or regexec to locate them.

Value

This function returns a character vector of length=1, ie only one (normally the longest) common sequence of characters is identified. If nothing is found common/shared an empty character-vector is returned

See Also

Use gregexpr or regexec in grep for locating the identified common characters in the initial query.

Inverse : Trim redundant text (from either side) to keep only variable part using trimRedundText; you may also look for related functions in package stringr

Examples

```r
keepCommonText()
```

```r
txt1 <- c("abcd_abc_kj", "bcd_abc12", "cd_abc_po")
keepCommonText(txt1, side="center") # trim from right

txt2 <- c("ddd_abc", "ddd_bcd", "ddd_cde")
trimRedundText(txt2, side="left") #
keepCommonText(txt2, side="center") #
```
levIndex

Transform (factor) levels into index

Description
This function helps transforming a numeric or character vector into indexes of levels (of its original values). By default indexes are assigned by order of occurrence, i.e., the first value of \( x \) will be get the index of 1. Using the argument \( \text{byOccurance}=\text{FALSE} \) the resultant indexes will follow the sorted values.

Usage

```r
levIndex(
  dat,
  byOccurance = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `dat` (numeric or character vector or factor) main input
- `byOccurance` (logical) toggle if lowest index should be based on alphabetical order or on order of input
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

matrix with mean values

See Also

`rowSds`, `colSums`

Examples

```r
x1 <- letters[rep(c(5,2:3),1:3)]
levIndex(x1)
levIndex(x1, byOccurance=FALSE)
## with factor
fa1 <- factor(letters[rep(c(5,2:3),1:3)], levels=letters[1:6])
levIndex(fa1)
levIndex(fa1, byOccurance=FALSE)
```
Description

The aim of this function is to select the data suit ing set of levels of the main input data to construct a linear regression model. In real world measurements one may be confronted to the case of very low level analytes below the detection limit (LOD) and resulting read-outs fluctuate around around a common baseline (instead of NA). With such data it may be preferable to omit the read-outs for the lowest concentrations/levels of analytes if they are spread around a base-line value. This function allows trying to omit all starting levels designed in startLev, then the resulting p-values for the linear regression slopes will be checked and the best p-value chosen. The input may also be a MArrayLM-type object from package limma or from moderTestXgrp or moderTest2grp. In the graphical representation all points associated to levels omitted are shown in light green. For the graphical display additional information can be used: If the dat is list or MArrayLM-type object, the list-elements $raw (according to argument lisNa will be used to display points initially given as NA ad imputed later on in grey. Logarithmic (ie log-linear) data can be treated by setting argument logExpect=TRUE. Then the levels will be taken as exponent of 2 for the regression, while the original values will be displayed in the figure.

Usage

```r
linModelSelect(
  rowNa,
  dat,
  expect,
  logExpect = FALSE,
  startLev = NULL,
  lisNa = c(raw = "raw", annot = "annot", datImp = "datImp"),
  plotGraph = TRUE,
  tit = NULL,
  pch = c(1, 3),
  cexLeg = 0.95,
  cexSub = 0.85,
  xLab = NULL,
  yLab = NULL,
  cexXAxis = 0.85,
  cexYAxis = 0.9,
  xLabLas = 1,
  cexLab = 1.1,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```
linModelSelect 155

Arguments

rowNa  (character, length=1) rowname for line to be extracted from dat
dat   (matrix, list or MArrayLM-object from limma) main input of which columns should get re-ordered, may be output from moderTestXgrp or moderTest2grp.
expect (numeric of character) the expected levels; if character, constant unit-characters will be stripped away to extract the numeric content
logExpect  (logical) toggle to TRUE if the main data are logarithmic but expect is linear
startLev  (integer) specify all starting levels to test for omitting here (multiple start sites for modelling linear regression may be specified to finally pick the best model)
lisNa  (character) in case dat is list or MArrayLM-type object, the list-elements with these names will be used as $raw (for indicating initial NA-values, $datImp (the main quantitation data to use) and $annot for displaying the corresponding value from the "Accession"-column.
plotGraph  (logical) display figure
tit  (character) optional custom title
pch  (integer) symbols to use n optional plot; 1st for regular values, 2nd for values not used in regression
cexLeg  (numeric) size of text in legend
cexSub  (numeric) text-size for line (as subtitle) giving regression details of best linear model
xLab  (character) custom x-axis label
yLab  (character) custom y-axis label
cexXAxis  (character) cex-type for size of text for x-axis labels
cexYAxis  (character) cex-type for size of text for y-axis labels
xLabLas  (integer) las-type orientation of x-axis labels (set to 2 for vertical axix-labels)
cexLab  (numeric) cex-type for size of text in x & y axis labels (will be passed to cex.lab in plot())
silent  (logical) suppress messages
debug  (logical) additional messages for debugging
callFrom  (character) allow easier tracking of messages produced

Value

This function returns a list with $coef (coefficients), $name (as/from input rowNa), $startLev the best starting level)

See Also

moderTestXgrp for single comparisons, order
Examples

## Construct data
li1 <- rep(c(4,3,3:6),each=3) + round(runif(18)/5,2)
names(li1) <- paste0(rep(letters[1:5], each=3), rep(1:3,6))
li2 <- rep(c(6,3:7), each=3) + round(runif(18)/5, 2)
dat2 <- rbind(P1=li1, P2=li2)
exp2 <- rep(c(11:16), each=3)

## Check & plot for linear model
linModelSelect("P2", dat2, expect=exp2)

## Log-Linear data
## Suppose dat2 is result of measures in log2, but exp4 is not
exp4 <- rep(c(3,10,30,100,300,1000), each=3)
linModelSelect("P2", dat2, expect=exp4, logE=FALSE) # bad
linModelSelect("P2", dat2, expect=exp4, logE=TRUE)

linRegrParamAndPVal

Fit linear regression, return parameters and p-values

Description

This function fits a linear regression and returns the parameters, including p-values from Anova. Here the vector `y` (scalar response or dependent variable, i.e., the value that should get estimated) will be estimated according to `dep` (explanatory or independent variable). Alternatively, `dep` may me a matrix where 1st column will be used as `dep` and the 2nd column as `y`.

Usage

linRegrParamAndPVal(
  dep,
  y = NULL,
  asVect = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dep (numeric vector, matrix or data.frame) explanatory or dependent variable, if matrix or data.frame the 1st column will be used, if `y=NULL` the 2nd column will be used as `y`
y (numeric vector) independent variable (the value that should get estimated based on `dep`)
asVect (logical) return numeric vector (Intercept, slope, p.intercept, p.slope) or matrix or results
listBatchReplace replaces in list lst all entries with value searchValue by replaceBy

Usage

```r
listBatchReplace(
  lst,
  searchValue, replaceBy,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `lst` input-list to be used for replacing
- `searchValue` (character, length=1)
- `replaceBy` (character, length=1)
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a corrected list
listGroupsByNames

Organize values into list and sort by names

Description

Sort values of 'x' by its names and organize as list by common names, the names until 'sep' are used for (re)grouping. Note that typical separators occuring the initial names may need protection by '\' (this is automatically taken care of for the case of the dot ('.') separator).

Usage

listGroupsByNames(x, sep = ".", silent = FALSE, debug = FALSE, callFrom = NULL)

Arguments

x (list) main input
sep (character) separator (note that typical separators may need to be protected, only automatically added for ".")
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

matrix or data.frame

See Also

rbind in cbind

Examples

listGroupsByNames((1:10)/5)
ser1 <- 1:6; names(ser1) <- c("AA","BB","AA.1","CC","AA.b","BB.e")
listGroupsByNames(ser1)
**Description**

`lmSelClu` runs linear regression on data segmented previously (eg by clustering). This function offers various types of (2-coefficient) linear regression on 2 columns of 'dat' (matrix with 3rd col named 'clu' or 'cluID', numeric elements for cluster-number). If argument 'clu' is (default) 'max', the column 'clu' will be inspected to take most frequent value of 'clu', otherwise a numeric entry specifying the cluster to extract is expected. Note: this function was initially made for use with results from `diagCheck()` Note: this function lacks means of judging goodness of fit of the regression performed & means for plotting.

**Usage**

```r
lmSelClu(
  dat,
  useCol = 1:2,
  clu = "max",
  regTy = "lin",
  filt1 = NULL,
  filt2 = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `dat` (matrix or data.frame)
- `useCol` (integer or character) specify which 2 columns of `dat` to use for linear regression
- `clu` (character) name of cluster to be extracted and treated
- `regTy` (character) change type used for linear regression: 'lin' for 1st col ~ 2nd col, 'res' for residue ~ 2nd col, 'norRes' for residue/2nd col ~2nd col or 'sqNorRes', 'inv' for 1st col ~ 1/(2nd col), 'invRes' for residue ~ 1/(2nd col)
- `filt1` (logical or numerical) filter criteria for 1st of `useCol`, if numeric then select all lines of dat less than max of filt1
- `filt2` (logical or numerical) filter criteria for 2nd of `useCol`, if numeric then select all lines of dat less than max of filt2
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

**Value**

`lm` object (or NULL if no data left)
lrbind

**See Also**

lm

**Examples**

```r
set.seed(2016); ran1 <- runif(220)
mat1 <- round(rbind(matrix(c(1:100+ran1[1:100],rep(1,50)),ncol=3),
  matrix(c(1:60,68:9+ran1[101:160],rep(2,60)),nc=3)),1)
colnames(mat1) <- c("a","BB","clu")
lmSelClu(mat1)
plot(mat1[which(mat1[,3]=="2"),1:2],col=grey(0.6))
abline(lmSelClu(mat1),lty=2,lwd=2)
#
mat2 <- round(rbind(matrix(c(1:100+ran1[1:100],rep(1,50)),ncol=3),
  matrix(c(1:60,(2:61+ran1[101:160])**2,rep(2,60)),nc=3)),1)
colnames(mat2) <- c("a","BB","clu")
(reg2 <- lmSelClu(mat2,regTy="sqNor"))
plot(function(x) coef(reg2)[2]+ (coef(reg2)[2]*x^2),xlim=c(1,70))
points(mat2[which(mat2[,3]=="2"),1:2],col=2)
```

---

**Description**

lrbind-like function to append list-elements containing matrices (or data.frames) and return one long table. All list-elements must have same number of columns (and same types of classes in case of data.frames. Simple vectors (as list-elements) will be considered as single lines for attaching.

**Usage**

```r
lrbind(lst, silent = FALSE, debug = FALSE, callFrom = NULL)
```

**Arguments**

- `lst` (list, composed of multiple matrix or data.frames or simple vectors) main input (each list-element should have same number of columns, numeric vectors will be converted to number of columns of other columns/elements)
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

**Value**

This function returns (depending on input) a matrix or data.frame
**makeMAList**

Make MA-List object

**Description**

`makeMAList` extracts sets of data-pairs (like R & G series) and makes MA objects as MA-List object (eg for ratio oriented analysis). The grouping of columns as sets of replicate-measurements is done according to argument 'MAfac'. The output is fully compatible to functions of package `limma` (Bioconductor).

**Usage**

```r
makeMAList(
  mat,
  MAfac,
  useF = c("R", "G"),
  isLog = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `mat`: main input matrix
- `MAfac`: (factor) factor orgnaizing columns of `mat` (if `useF` contains the default 'R' and 'G', they should also be part of `MAfac`)
- `useF`: (character) two specific factor-leves of `MAfac` that will be used/extracted
- `isLog`: (logical) tell if data is already log2 (will be considered when computing M and A values)
- `silent`: (logical) suppress messages
- `debug`: (logical) additional messages for debugging
- `callFrom`: (character) allow easier tracking of messages produced

**Details**

This function requires Bioconductor package `limma` being installed.

**See Also**

`rbind` in `cbind`

**Examples**

```r
lst1 <- list(matrix(1:9, ncol=3, dimnames=list(letters[1:3],c("AA","BB","CC"))),
             11:13, matrix(51:56, ncol=3))
lrbind(lst1)
```
makeNRedMatr

Value

limma-type "MAList" containing M and A values

See Also

test2factLimma, for creating RG-lists within limma: MA.RG in normalizeWithinArrays

Examples

```r
set.seed(2017); t4 <- matrix(round(runif(40,1,9),2), ncol=4,
    dimnames=list(letters[c(1:5,3:4,6:4)], c("AA1","BB1","AA2","BB2")))
makeMAList(t4, gl(2,2,labels=c("R","G")))
```

Description

makeNRedMatr takes matrix or data.frame 'dat' to summarize redundant lines (column argument iniID) along method specified in summarizeRedAs to treat all lines with redundant iniID by same approach (ie for all columns the line where specified column is at eg max = 'maxOfRef'). If no name given, the function will take the last numeric (factors may be used - they will be read as levels).

Usage

```r
makeNRedMatr(
    dat,
    summarizeRedAs,
    iniID = "iniID",
    retDataFrame = TRUE,
    silent = FALSE,
    debug = FALSE,
    callFrom = NULL
)
```

Arguments

- `dat`: (matrix or data.frame) main input for making non-redundant
- `summarizeRedAs`: (character) summarization method(s), typical choices 'median','mean','min' or 'maxOfRef','maxAbsOfRef' for summarizing according to 1 specified column, may be single method for all or different method for each column (besides col 'iniID') or special method looking at column (if found, first of special methods used, everything else not considered).
- `iniID`: (character) column-name used as initial ID (default="iniID")
- `retDataFrame`: (logical) if TRUE, check if text-columns may be converted to data.frame with numeric
matchMatrixLinesToRef

Match All Lines of Matrix To Reference

Description

This function allows adjusting the order of lines of a matrix `mat` to a reference character-vector `ref`, even when initial direct matching of character-strings using `match` is not possible/successful. In this case, various variants of using `grep` will be used to see if unambiguous matching is possible of characteristic parts of the text. All columns of `mat` will be tested and the column giving the best results will be used.

Usage

```r
matchMatrixLinesToRef(
  mat,
  ref,
  addRef = TRUE,
  inclInfo = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

silent (logical) suppress messages
debug (logical) for bug-tracking: more/enhanced messages
callFrom (character) allows easier tracking of message(s) produced

Value

This function returns a (numeric) matrix or data.frame with summarized data and an additional column with the number of initial redundant lines.

See Also

simple/partial functionality in `summarizeCols`, `checkSimValueInSer`

Examples

```r
t3 <- data.frame(ref=rep(11:15,3), tx=letters[1:15],
  matrix(round(runif(30,-3,2),1),nc=2),stringsAsFactors=FALSE)
by(t3,t3[,1],function(x) x)
t(sapply(by(t3,t3[,1],function(x) x), summarizeCols, me="maxAbsOfRef"))
(xt3 <- makeNRedMatr(t3, summ="mean", iniID="ref"))
(xt3 <- makeNRedMatr(t3, summ=unlist(list(X1="maxAbsOfRef")), iniID="ref"))
```
Arguments

- **mat**: (matrix or data.frame) main input, all columns of mat will be tested for (partial) matching of ref
- **ref**: (character, length must match) reference for trying to match each of the columns of mat
- **addRef**: (logical), if TRUE the content of ref will be added to mat as additional column
- **inclInfo**: (logical) allows returning list with new matrix and additional information
- **silent**: (logical) suppress messages
- **debug**: (logical) additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

Details

This function tests all columns of mat to find perfect matching results to the reference ref. In case of multiple results the In case no direct matching is possible, grep will be used to find the best partial matching. The order of the rows of input mat will be adjusted according to the matching results.

If addRef=TRUE, the reference will be included as additional column to the results, too.

Value

This function returns the input matrix in an adjusted order (plus an optional additional column showing the reference) or if inclInfo=TRUE a list with $mat (adjusted matrix), $byColumn, $newOrder and $method; the reference can be added as additional last column if addRef=TRUE

See Also

match, grep, trimRedundText, replicateStructure

Examples

```r
## Note : columns b and e allow non-ambigous match, not all elements of e are present in a
mat0 <- cbind(a=c("mvvk","axxd","bxxd","vv"),b=c("iwwy","iyyu","kvvh","gxx"), c=rep(9,4),
   d=c("hgf","hgf","vxc","nvnn"), e=c("_vv_","_ww_","_xx_","_yy_"))
m1 <- matchMatrixLinesToRef(mat0[,1:4], ref=mat0[,5])
m2 <- matchMatrixLinesToRef(mat0[,1:4], ref=mat0[1:3,5], inclInfo=TRUE)
m3 <- matchMatrixLinesToRef(mat0[,-2], ref=mat0[,2], inclInfo=TRUE)  # needs 'reverse grep'
```
matchNamesWithReverseParts

**Value Matching with optional reversing of sub-parts of non-matching elements**

**Description**

This function provides a variant to `match`, where initially non-matching elements of `x` will be tested by decomposing non-matching elements, reversing the parts in front and after the separator `sep` and re-matching. If separator `sep` does not occur, a warning will be issued, if it occurs more than once, the parts before and after the first separator will be used and a warning issued.

**Usage**

```r
matchNamesWithReverseParts(
  x,
  y,
  sep = "-",
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `x` (character): first vector for match
- `y` (character): second vector for match
- `sep` (character): separator between elements
- `silent` (logical): suppress messages
- `debug` (logical): additional messages for debugging
- `callFrom` (character): allow easier tracking of messages produced

**Value**

index for matching (integer) x to y

**See Also**

`match`

**Examples**

```r
tx1 <- c("a-b","a-c","d-a","d-b","b-c","d-c")
tmp <- triCoord(4)
tx2 <- paste(letters[tmp[,1]],letters[tmp[,2]],sep="-")
## Some matches won't be found, since 'a-d' got reversed to 'd-a', etc...
match(tx1,tx1)
matchNamesWithReverseParts(tx1,tx2)
```
matchSampToPairw

Match names to concatenated pairs of names

Description

The column-names of multiple pairwise testing contain the names of the initial groups/conditions tested, plus there is a separator (eg ‘-’ in moderTestXgrp). Thus function allows to map back which groups/conditions were used by returning the index of the respective groups used in pair-wise sets.

Usage

matchSampToPairw(
  grpNa,
  pairwNa,
  sep = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

grpNa (character) the names of the groups of replicates (ie conditions) used to test
pairwNa (character) the names of pairwise-testing (ie ‘concatenated’ sampNa
sep (character) if not NULL the characters given will be used via stringsplit
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Details

There are two modes of operation: 1) Argument sep is set to NULL: The names of initial groups/conditions (grpNa) will be tested for exact pattern matching either at beginning or at end of pair-wise names (pairwNa). This approach has the advantage that it does not need to be known what character(s) were used as separator (or they may change), but the disadvantage that in case the perfect grpNa was not given, the longest best match of grpNa will be returned.

2) The separator sep is given and exact matches at both sides will be searched. However, if the character(s) from sep do appear inside grpNa no matches will be found.

If some grpNa are not found in pairwNa this will be marked as NA.

Value

matrix of 2 columns with indices of sampNa with pairwNa as rows
**matr2list**

*Transform columns of matrix to list of vectors*

**Description**

convert matrix to list of vectors: each column of 'mat' as vector of list

**Usage**

```r
matr2list(mat, concSym = ".", silent = FALSE, debug = TRUE, callFrom = NULL)
```

**Arguments**

- `mat` (matrix) main input
- `concSym` (character) symbol for concatenating: concatenation of named vectors in list names as colname(s)+'concSym'+rowname
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

**Value**

matrix or array (1st dim is intraplate-position, 2nd .. plate-group/type, 3rd .. channels)

**See Also**

- `convToNum`

**Examples**

```r
mat1 <- matrix(1:12,ncol=3,dimnames=list(letters[1:4],LETTERS[1:3]))
mat2 <- matrix(LETTERS[11:22],ncol=3,dimnames=list(letters[1:4],LETTERS[1:3]))
matr2list(mat1); matr2list(mat2)
```
**Description**

This function allows merging of multiple matrix-like objects. The matrix-rownames will be used to align common elements, either by returning all common elements mode='intersect' or containing all elements mode='union' (the result may contain additional NAs).

**Usage**

```r
mergeMatrices(
  ..., 
  mode = "intersect",
  useColumn = 1,
  na.rm = TRUE,
  extrRowNames = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `...` (matrix or data.frame) multiple matrix or data.frame objects may be entered
- `mode` (character) allows choosing restricting to all common elements (mode='intersect') or union (mode='union')
- `useColumn` (integer, character or list) the column(s) to consider, may be 'all' to use all, integer to select specific indexes or list of indexes or colnames for custom-selection per matrix
- `na.rm` (logical) suppress NAs
- `extrRowNames` (logical) decide whether columns with all values different (ie no replicates or max divergency) should be excluded
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

**Details**

Custom column-names can be given by entering matrices like named arguments (see examples below). The choice of columns to use may be adopted to each matrix entered, in this case the argument `useColumn` may be a list with matrix-names to use or a list of indexes (see examples below).

Note, that matrices may contain repeated rownames (see examples, mat3). In this case only the first of repeated rownames will be considered (and lines of repeated names ignored).
**mergeMatrixList**

**Value**

This function returns a matrix containing all selected columns of the input matrices to fuse.

**See Also**

merge, mergeMatrixList

**Examples**

```r
mat1 <- matrix(11:18, ncol=2, dimnames=list(letters[3:6], LETTERS[1:2]))
mat2 <- matrix(21:28, ncol=2, dimnames=list(letters[2:5], LETTERS[3:4]))
mat3 <- matrix(31:38, ncol=2, dimnames=list(letters[c(1,3:4,3)], LETTERS[4:5]))

mergeMatrices(mat1, mat2)
mergeMatrices(mat1, mat2, mat3, mode="union", useCol=2)
## custom names for matrix-origin
mergeMatrices(m1=mat1, m2=mat2, mat3, mode="union", useCol=2)
## flexible/custom selection of columns
mergeMatrices(m1=mat1, m2=mat2, mat3, mode="union", useCol=list(1,1:2,2))
```

---

**mergeMatrixList**  
**Merge Multiple Matrices from List**

**Description**

This function allows merging of multiple matrix-like objects from an initial list. The matrix-rownames will be used to align common elements, either be returning all common elements `mode="intersect"` or containing all elements `mode="union"` (the result may contains additional NAs).

**Usage**

```r
mergeMatrixList(
  matLst,
  mode = "intersect",
  useColumn = 1,
  na.rm = TRUE,
  extrRowNames = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `matLst` (list containing matrices or data.frames) main input (multiple matrix or data.frame objects)
- `mode` (character) allows choosing restricting to all common elements (mode='intersect') or union (mode='union')
mergeSelCol

**Description**

This function merges selected columns out of 2 matrix or data.frames. `selCols` will be used to define columns to be used; optionally may be different for `dat2` : define in `supCols2`. Output-cols will get additions specified in `newSuff` (default `.x` and `.y`).

**Usage**

```r
mergeSelCol(useColumn, na.rm = FALSE, extrRowNames = FALSE, silent = FALSE, debug = FALSE, callFrom = NULL)
```

**Arguments**

- `useColumn`: (integer, character or list) the column(s) to consider, may be 'all' to use all, integer to select specific indexes or list of indexes or colnames for cutom-selection per matrix
- `na.rm`: (logical) suppress NAs
- `extrRowNames`: (logical) decide whether columns with all values different (ie no replicates or max divergency) should be excluded
- `silent`: (logical) suppress messages
- `debug`: (logical) additional messages for debugging
- `callFrom`: (character) allow easier tracking of messages produced

**Details**

Custom column-names can be given by entering matrices like named arguments (see examples below). The choice of columns to use may be adopted to each matrix entered, in this case the argument `useColumn` may be a list with matrix-names to use or a list of indexes (see examples below).

Note, that matrices may contain repeated rownames (see examples, `mat3`). In this case only the first of repeated rownames will be considered (and lines of repeated names ignored).

**Value**

This function returns a matrix containing all selected columns of the input matrices to fuse

**See Also**

`merge`, `mergeMatrices` for separate entries

**Examples**

```r
mat1 <- matrix(11:18, ncol=2, dimnames=list(letters[3:6], LETTERS[1:2]))
mat2 <- matrix(21:28, ncol=2, dimnames=list(letters[2:5], LETTERS[3:4]))
mat3 <- matrix(31:38, ncol=2, dimnames=list(letters[c(1,3:4,3)], LETTERS[4:5]))
mergeMatrixList(list(mat1, mat2))
mergeMatrixList(list(m1=mat1, m2=mat2, mat3), mode="union", useCol=2)
```
mergeSelCol

Usage

mergeSelCol(
  dat1,
  dat2,
  selCols,
  supCols2 = NULL,
  byC = NULL,
  useAll = FALSE,
  setRownames = TRUE,
  newSuff = c(".x", ".y"),
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dat1 matrix or data.frame for fusing
dat2 matrix or data.frame for fusing
selCols will be used to define columns to be used; optionally may be different for 'dat2'
: define in 'supCols2'
supCols2 if additional column-names should be extracted from dat2
byC (character) 'by' value used in merge
useAll (logical) use all lines (will produce NAs when given identifier not found un 2nd
group of data)
setRownames (logical) if TRUE, will use values of col used as 'by' as rownames instead of
showing as add'1 col in output
newSuff (character) prefix (argument 'suffixes' in merge)
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This function returns a data.frame containing the merged columns

See Also

merge, merge 3 data.frames using mergeSelCol3

Examples

mat1 <- matrix(c(1:7,letters[1:7],11:17), ncol=3, dimnames=list(LETTERS[1:7],c("x1","x2","x3")))
mat2 <- matrix(c(1:6,c("b","a","e","f","g","k"), 31:36),
  ncol=3, dimnames=list(LETTERS[11:16],c("y1","x2","x3")))
mergeSelCol(mat1, mat2, selC=c("x2","x3"))
mergeSelCol3

Description

successive merge of selected columns out of 3 matrix or data.frames. ‘selCols’ will be used to define columns to be used; optionally may be different for ‘dat2’ : define in ‘supCols2’. Output-cols will get additions specified in newSuff (default ‘.x’ and ‘.y’)

Usage

mergeSelCol3(
  dat1,
  dat2,
  dat3,
  selCols,
  supCols2 = NULL,
  supCols3 = NULL,
  byC = NULL,
  useAll = FALSE,
  setRownames = TRUE,
  newSuff = c(".x", ".y", ".z"),
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dat1 matrix or data.frame for fusing
dat2 matrix or data.frame for fusing
dat3 matrix or data.frame for fusing
selCols will be used to define columns to be used; optionally may be different for ‘dat2’ : define in ‘supCols2’
supCols2 if additional column-names should be extracted form dat2
supCols3 if additional column-names should be extracted form dat3
byC (character) ‘by’ value used in merge
useAll (logical) use all lines (will produce NAs when given identifier not found un 2nd group of data)
setRownames if TRUE, will use values of col used as ‘by’ as rownames instead of showing as add’l col in output
newSuff (character) prefix (argument ‘suffixes’ in merge)
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of messages produced
**mergeVectors**

**Value**

This function returns a data.frame containing the merged columns

**See Also**

merge, mergeSelCol

**Examples**

```r
mat1 <- matrix(c(1:7, letters[1:7], 11:17), ncol=3, names=LETTERS[1:7], dimnames=list(LETTERS[1:7], c("x1", "x2", "x3")))
mat2 <- matrix(c(1:6, c("b", "a", "e", "f", "g", "k"), 31:36), ncol=3, names=LETTERS[11:16], dimnames=list(LETTERS[11:16], c("y1", "x2", "x3")))
mat3 <- matrix(c(1:6, c("c", "a", "e", "b", "g", "k"), 51:56), ncol=3, names=LETTERS[11:16], dimnames=list(LETTERS[11:16], c("z1", "x2", "x3")))
mergeSelCol(mat1, mat2, mat3, selC=c("x2", "x3"))
```

---

**mergeVectors**

**Merge Named Vectors**

**Description**

This function allows merging for multiple named vectors (each element needs to be named). Basically, all elements carrying the same name across different input-vectors will be aligned in the same column of the output (input-vectors appear as lines). If vectors are not given using a name (see first example below), they will be names 'x.1' etc (see argument `namePrefix`).

**Usage**

```r
mergeVectors(
  ..., 
  namePrefix = "x.", 
  NAto0 = FALSE, 
  silent = FALSE, 
  debug = FALSE, 
  callFrom = NULL
)
```

**Arguments**

- `...`: all vectors that need to be merged
- `namePrefix`: (character) prefix to numers used when vectors are not given with explicit names (second example)
- `NAto0`: (logical) optional replacement of NAs by 0
- `silent`: (logical) suppress messages
- `debug`: (logical) additional messages for debugging
- `callFrom`: (character) allow easier tracking of messages produced
Details

Note: The arguments 'namePrefix', 'NAto0', 'callFrom' and 'silent' must be given with full name to be recognized as such (and not get considered as vector for merging).

Value

This function returns a matrix of merged values

See Also

merge (for two data.frames)

Examples

x1 <- c(a=1, b=11, c=21)
x2 <- c(b=12, c=22, a=2)
x3 <- c(a=3, d=43)
mergeVectors(vect1=x1, vect2=x2, vect3=x3)
x4 <- 41:44  # no names - not conform for merging
mergeVectors(x1, x2, x3, x4)

mergeW2

Extended version of merge for multiple objects (even without row-names)

Description

mergeW2 provides flexible merging out of 'MArrayLM'-object (if found, won’t consider any other input-data) or of separate vectors or matrixes. The main idea was to have somthing not adding add’l lines as merge might do, but to stay within the frame of the 1st argument given, even when IDs are repeated, so the output follows the order of the 1st argument, non-redundant IDs are created (orig IDs as new column). If no 'MArrayLM'-object found: try to combine all elements of input '...', input-names must match predefined variants 'chInp'. IDs given in 1st argument and not found in later arguments will be displayed as NA in the output matrix of data.frame. Note: (non-data) arguments must be given with full name (so far no lazy evaluation, may conflict with names in 'inputNamesLst'). Note: special characters in colnames bound to give trouble. Note: when no names given, mergeW2 will presume order of elements (names) from 'inputNamesLst'. PROBLEM: error after xxMerg3 when several entries have matching (row)names but some entries match only partially (what to do: replace with NAs ??)

Usage

mergeW2(
  ..., nonRedundID = TRUE, convertDF = TRUE, selMerg = TRUE, inputNamesLst = NULL, ...)
Arguments

... all data (vectors, matrices or dataframes) intended for merge
nonRedundID (logical) if TRUE, always add 1st column with non-redundant IDs (add anyway if non-redundant IDs found)
convertDF (logical) allows converting output to data.frame, add new heading col with non-red rownames & check which cols should be numeric
selMerg (logical) if FALSE toggle to classic merge() (will give more rows in output in case of redundant names
inputNamesLst (list) named list with character vectors (should be unique), search these names in input for extracting/merging elements use for 'lazy matching' when checking names of input, default: 7 groups ('Mvalue', 'Avalue', 'p.value', 'mouseInfo', 'Lfdr', 'link', 'filt') with common short versions
noMatchPursue (logical) allows using entries where 0 names match (just as if no names given)
standColNa (logical) if TRUE return standard colnames as defined in 'inputNamesLst' (ie 'chInp'), otherwise colnames as initially provided
lastOfMultCols may specify input groups where only last col will be used/extracted
duplTxtSep (character) separator for counting/denomingating multiple occurrences of same name
silent (logical) suppress messages
debug (logical) for bug-tracking: more/enhanced messages and intermediate objects written in global name-space
callFrom (character) allows easier tracking of message(s) produced

Value
matrix or data.frame of fused data

See Also
merge

Examples
t1 <- 1:10; names(t1) <- letters[c(1:7,3:4,8)]
t2 <- 20:11; names(t2) <- letters[c(1:7,3:4,8)]
t3 <- 101:110; names(t3) <- letters[c(11:20)]
t4 <- matrix(100:81, ncol=2, dimnames=list(letters[1:10], c("col1","col2")))
t5 <- cbind(t1=t1,t52=t1+20,t53=t1+30)
t1; t2; t3; cbind(t1,t2)
mergeW2(Mval=t1,p.value=t2,debug=FALSE)

---

**minDiff**

*Minimum distance/difference between values*

### Description

This function aims to find the min distance (ie closest point) to any other x (numeric value), ie intra 'x' and returns matrix with 'index','value','dif','ppm','ncur','nbest','best'. At equal distance to lower & upper neighbour point, the upper (following) point is chosen (as single best). In case of multiple ex-aequo distance returns 1st of multiple, may be different at various repeats.

### Usage

```r
minDiff(
  x,
  digSig = 3,
  ppm = TRUE,
  initOrder = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

### Arguments

- **x** (numeric) vector to search minimum difference
- **digSig** number of significant digits, used for ratio or ppm column
- **ppm** (logical) display distance as ppm (1e6*diff/refValue, ie normalized difference eg as used in mass spectrometry), otherwise the ratio is given as : value(from 'x')/closestValue (from 'x')
- **initOrder** (logical) return matrix so that 'x' matches exactly 2nd col of output
- **silent** (logical) suppress messages
- **debug** (logical) additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced

### Value

This function returns a matrix

### See Also

`diff`
moderTest2grp

Examples

```r
set.seed(2017); aa <- 100*c(0.1 + round(runif(20),2),0.53,0.53)
minDiff(aa);
minDiff(aa,initO=TRUE,ppm=FALSE); .minDif(unique(aa))
```

moderTest2grp

**Moderated pair-wise t-test from limma**

**Description**

Runs moderated t-test from package 'limma' on each line of data. Note: This function requires the package *limma* from Bioconductor. The limma contrast-matrix has to be read by column, the lines in the contrast-matrix containing '+1' will be compared to the '-1' lines, eg grpA-grpB. Local false discovery rates (lfdr) estimations will be made using the CRAN-package *fdrtool* (if available).

**Usage**

```r
moderTest2grp(
  dat,
  grp,
  limmaOutput = TRUE,
  addResults = c("lfdr", "FDR", "Mval", "means"),
  testOrientation = "=",
  silent = FALSE,
  callFrom = NULL
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>dat</code></td>
<td>matrix or data.frame with rows for multiple (independent) tests, use ONLY with 2 groups; assumed as log2-data</td>
</tr>
<tr>
<td><code>grp</code></td>
<td>(factor) describes column-relationship of <code>dat</code> (1st factor is considered as reference -&gt; orientation of M-values !!)</td>
</tr>
<tr>
<td><code>limmaOutput</code></td>
<td>(logical) return full (or extended) MArrayLM-object from limma or <code>FALSE</code> for only the (uncorrected) p.values</td>
</tr>
<tr>
<td><code>addResults</code></td>
<td>(character) types of results to add besides basic limma-output, data are assumed to be log2 ! (eg &quot;lfdr&quot; using fdrtool-package, &quot;FDR&quot; or &quot;BH&quot; for BH-FDR, &quot;BY&quot; for BY-FDR, &quot;bonferroni&quot; for Bonferroni-correction, &quot;qValue&quot; for lfdr by qvalue, &quot;Mval&quot;, &quot;means&quot; or &quot;nonMod&quot; for non-moderated test and he equivaent all (other) multiple testing corrections chosen here)</td>
</tr>
<tr>
<td><code>testOrientation</code></td>
<td>(character) for one-sided test (&quot;&gt;&quot;,&quot;greater&quot; or &quot;&lt;&quot;,&quot;less&quot;), NOTE : 2nd grp is considered control/reference, '&lt;' will identify grp1 &lt; grp2</td>
</tr>
<tr>
<td><code>silent</code></td>
<td>(logical) suppress messages</td>
</tr>
<tr>
<td><code>callFrom</code></td>
<td>(character) allow easier tracking of message(s) produced</td>
</tr>
</tbody>
</table>
moderTestXgrp

Multiple moderated pair-wise t-tests from limma

Description

Runs all pair-wise combinations of moderated t-tests from package 'limma' on each line of data against 1st group from 'grp'. Note: This function requires the package limma from bioconductor. The limma contrast-matrix has to be read by column, the lines in the contrast-matrix containing '+1' will be compared to the '-1' lines, eg grpA-grpB.

Usage

moderTestXgrp(
  dat,
  grp,
  limmaOutput = TRUE,
  addResults = c("lfdr", "FDR", "Mval", "means"),
  testOrientation = "=",
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
moderTestXgrp

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Arguments
dat

matrix or data.frame with rows for multiple (independent) tests, use ONLY with
2 groups; assumed as log2-data !!!

grp

(factor) describes column-relationship of ’dat’ (1st factor is considered as reference -> orientation of M-values !!)

limmaOutput

(logical) return full (or extended) MArrayLM-object from limma or ’FAlSE’ for
only the (uncorrected) p.values

addResults

(character) types of results to add besides basic limma-output, data are assumed
to be log2 ! (eg "lfdr" using fdrtool-package, "FDR" or "BH" for BH-FDR,
"BY" for BY-FDR, "bonferroni" for Bonferroni-correction, "qValue" for lfdr by
qvalue, "Mval", "means" or "nonMod" for non-moderated test and he equivaent
all (other) multiple testing corrections chosen here)

testOrientation
(character) for one-sided test (">","greater" or "<","less"), NOTE : 2nd grp is
considered control/reference, ’<’ will identify grp1 < grp2
silent

(logical) suppress messages

debug

(logical) additional messages for debugging

callFrom

(character) allow easier tracking of message(s) produced

Value
This function returns a limma-type MA-object (list)
See Also
moderTest2grp for single comparisons, lmFit and the eBayes-family of functions in package
limma
Examples
grp <- factor(rep(LETTERS[c(3,1,4)],c(2,3,3)))
set.seed(2017); t8 <- matrix(round(rnorm(208*8,10,0.4),2), ncol=8,
dimnames=list(paste(letters[],rep(1:8,each=26),sep=""), paste(grp,c(1:2,1:3,1:3),sep="")))
# augment lines 3:6 (c-f)
t8[5:8,c(1:2,6:8)] <- t8[5:8,c(1:2,6:8)] -1.5
# lower lines
# augment lines
## expect to find C/A in c,d,g, (h)
## expect to find C/D in c,d,e,f
## expect to find A/D in f,g,(h)
test8 <- moderTestXgrp(t8, grp)
# If you have limma installed we can now see further
if("list" %in% mode(test8)) head(test8$p.value, n=8)


multiCharReplace

Multiple replacement of entire character elements in simple vector, matrix or data.frame

Description

This function allows multiple types of replacements of entire character elements in simple vector, matrix or data.frame. In addition, the result may be optionally directly transformed to logical or numeric.

Usage

multiCharReplace(
  mat, 
  repl, 
  convTo = NULL, 
  silent = FALSE, 
  debug = TRUE, 
  callFrom = NULL
)

Arguments

mat (character vector, matrix or data.frame) main data
repl (matrix or list) tells what to replace by what: If matrix the 1st column will be considered as 'old' and the 2nd as 'replaceBy'; if named list, the names of the list-elements will be considered as 'replaceBy'
convTo (character) optional conversion of content to 'numeric' or 'logical'
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This function returns an object of same dimension as input (with replaced content)

See Also
grep

Examples

x1 <- c("ab","bc","cd","efg","ghj")
multiCharReplace(x1, cbind(old=c("bc","efg"), new=c("BBCC","EF")))

x2 <- c("High","n/a","High","High","Low")
multiCharReplace(x2, cbind(old=c("n/a", "Low", "High"), new=c(NA, FALSE, TRUE)), convTo="logical")

# works also to replace numeric content:
x3 <- matrix(11:16, ncol=2)
multiCharReplace(x3, cbind(12:13, 112:113))

---

**multiMatch**

*Simple Multi-to-Multi Matching of (Concatenated) Terms*

**Description**

This function allows convenient matching of multi-to-multi relationships between two objects/vectors. It was designed for finding common elements in multiple to multiple matching situations (e.g., when comparing `c("aa; bb", "cc")` to `c("bb; ab", "dd")`, i.e., to find 'bb' as matching between both objects).

**Usage**

```r
multiMatch(
  x,
  y,
  sep = "; ",
  sep2 = NULL,
  method = "byX",
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- **x** (vector or list) first object to compare; if vector, the (partially) concatenated identifiers (will be split using separator `sep`), or list of items to be matched (i.e., already split)
- **y** (vector or list) second object to compare; if vector, the (partially) concatenated identifiers (will be split using separator `sep`), or list of items to be matched (i.e., already split)
- **sep** (character, length=1) separator used to split concatenated identifiers (if `x` or `y` is vector)
- **sep2** (character, length=1) optional separator used when `method="matched"` to concatenate all indexes of `y` for column `y.allInd`
- **method** (character) mode of operation: `asIndex` to return index of `y` (those who have matches) with names of `x` (which `x` are the corresponding match)
- **silent** (logical) suppress messages
- **debug** (logical) display additional messages for debugging
- **callFrom** (character) allow easier tracking of message(s) produced
Details

method='byX' .. returns data.frame with view oriented towards entries of x: character column x for entire content of x; integer column x.Ind for index of x; character column TagBest for most frequent matching isolated tag/ID; integer column y.IndBest index of most frequent matching y; character column y.IndAll index for all y matching any of the tags; character column y.Match for entire content of best matching y; character column y.Adj for y adjusted to best matching y for easier subsequent perfect matching.

method=c("byX","filter") .. combined argument to keep only lines with any matches

method='byTag' .. returns matrix (of integers) from view of isolated tags from x (a separate line for each tag from x matching to y);

method=c("byTag","filter") .. if combined as arguments, this will return a data.frame for all unique tags with any matches between x and y, with additional columns x.AllInd for all matching x-indexes, y.IndBest best matching y index; x.n for number of different x containing this tag; y.AllInd for all matching y-indexes

method='adjustXtoY' .. returns vector with x adjusted to y, ie those elements of x matching are replace by the exact corresponding term of y.

method=NULL .. If no term matching the options shown above is given, another version of 'asIndex' is returned, but indexes to y_after_spliting by sep. Again, this method can be filtered by using method="filter" to focus on the best matches to x.

Value

matrix, data.frame or list with matching results depending on method chosen

See Also

match; strsplit

Examples

aa <- c("m","k","j; aa","m; aa; bb; o; ee","n; dd; cc","aa","cc")
bb <- c("dd; r","aa","ee; bb; q; cc","p; cc")
(match1 <- multiMatch(aa, bb, method=NULL))  # match bb to aa
(match2 <- multiMatch(aa, bb, method="byX"))   # match bb to aa
(match3 <- multiMatch(aa, bb, method="byTag")) # match bb to aa
(match4 <- multiMatch(aa, bb, method=c("byTag","filter")))  # match bb to aa

naOmit

Fast na.omit

Description

naOmit removes NAs from input vector. This function has no slot for removed elements while na.omit does so. Resulting objects from naOmit are smaller in size and subsequent execution (on large vectors) is faster (in particular if many NAs get encountered). Note: Behaves differently to na.omit with input other than plain vectors. Will not work with data.frames!
Usage

naOmit(x)

Arguments

x (vector or matrix) input

Value

vector without NAs (matrix input will be transformed to vector). Returns NULL if input consists only of NAs.

See Also

na.fail, na.omit

Examples

aA <- c(11:13,NA,10,NA);
naOmit(aA)

nFragments

Number of fragments after cut at specific character(s) within size-range

Description

nFragments determines number of fragments /entry within range of 'sizeRa' (numeric,length=2) when cutting after 'cutAt'

Usage

nFragments(protSeq, cutAt, sizeRa)

Arguments

protSeq (character) text to be cut
cutAt (character) position to cut
sizeRa (numeric,length=2) min and max size to consider

Value

numeric vector with number of fragments for each entry 'protSeq' (names are 'protSeq')

See Also

cutAtMultSites, simple version {nFragments0} (no size-range)
Examples

tmp <- "MSVSREDSLVLVTERIIAVSFSTANEENFRSNLREVAQMLKSKHGNYLLFNSRDPIDTLHAKVLEFGWPDHITALKEK"
nFragments(c(tmp,"ojioRij"),c("R","K"),c(4,31))

---

nFragments0

**Number of fragments after cut at specific character(s)**

Description

nFragments0 tells the number of fragments/entry when cutting after 'cutAt’

Usage

nFragments0(protSeq, cutAt)

Arguments

- protSeq: (character) text to be cut
- cutAt: (integer) position to cut

Value

numeric vector with number of fragments for each entry `protSeq` (names are `protSeq’)

See Also

more elaborate {nFragments}; cutAtMultSites

Examples

tmp <- "MSVSRTMEDSCELVTIYVTERIIAVSFSTANEENFRSNLREVAQMLKSKHGNYLLFNSRDPIDTLHAKVLEFGWPDHITALKEK"
nFragments0(c(tmp,"ojioRij"),c("R","K"))

---

nNonNumChar

**Count number of non-numeric characters**

Description

nNonNumChar counts number of non-numeric characters. Made for positive non-scientific values (eg won’t count neg-sign, neither Euro comma ‘,’)

Usage

nNonNumChar(txt)
Arguments

txt character vector to be treated

Value

This function returns a numeric vector with number of non-numeric characters (ie not '.' or 0-9)

See Also

nchar

Examples

nNonNumChar("a1b "); sapply(c("aa","12ab","a1b2","12","0.5"), nNonNumChar)

nonAmbiguousMat

Transform matrix to non-ambiguous matrix (in respect to given column)

Description

nonAmbiguousMat makes values of matrix ‘mat’ in col ‘byCol’ unique.

Usage

nonAmbiguousMat(
  mat, byCol, uniqOnly = FALSE, asList = FALSE, nameMod = "amb_", callFrom = NULL)

Arguments

mat numeric or character matrix (or data.frame), column specified by ‘byCol’ must be/will be used as numeric, 1st column of ‘mat’ will be considered like index & used for adding prefix ‘nameMod’ (unless byCol=1, then 2nd col will be used)

byCol (character or integer-index) column by which ambiguity will be tested

uniqOnly (logical) if =TRUE return unique only, if =FALSE return unique and single representative of non-unique values (with “ added to name), selection of representative of repeated: first (of sorted) or middle if >2 instances

asList (logical) return result as list

nameMod (character) prefix added to 1st column of ‘mat’ (expect ‘by’) for indicating non-unique/ambiguous values

callFrom (character) allow easier tracking of message(s) produced
nonAmbiguousNum

Value

sorted non-ambiguous numeric vector (or list if 'asList'=TRUE and 'uniqOnly'=FALSE)

See Also

for non-numeric use firstOfRepeated - but 1000x much slower !; get1stOfRepeatedByCol

Examples

set.seed(2017); mat2 <- matrix(c(1:100,round(rnorm(200),2)),ncol=3,
dimnames=list(1:100,LETTERS[1:3]));
head(mat2U <- nonAmbiguousMat(mat2,by="B",na="_",uniqO=FALSE),n=15)
head(get1stOfRepeatedByCol(mat2,sortB="B",sortS="B"))

nonAmbiguousNum

make numeric vector non-ambiguous (ie unique)

Description

nonAmbiguousNum makes (named) values of numeric vector 'x' unique. Note: for non-numeric use firstOfRepeated - but 1000x slower ! Return sorted non-ambiguous numeric vector (or list if 'asList'=TRUE and 'uniqOnly'=FALSE)

Usage

nonAmbiguousNum(
  x,
  uniqOnly = FALSE,
  asList = FALSE,
  nameMod = "amb_",
  callFrom = NULL
)

Arguments

x (numeric) main input
uniqOnly (logical) if=TRUE return unique only, if =FALSE return unique and single representative of non-unique values (with " added to name), selection of representative of repeated: first (of sorted) or middle if >2 instances
asList (logical) return list
nameMod (character) text to add in case on ambiguous values, default="amb_"
callFrom (character) allow easier tracking of message(s) produced

Value

sorted non-ambiguous numeric vector (or list if 'asList'=TRUE and 'uniqOnly'=FALSE)
nonredDataFrame

**See Also**

- `firstOfRepeated` for non-numeric use (much slower !!!), `duplicated`

**Examples**

```r
set.seed(2017); aa <- round(rnorm(100),2); names(aa) <- 1:length(aa)
str(nonAmbiguousNum(aa))
str(nonAmbiguousNum(aa,uniq=FALSE,asLi=TRUE))
```

---

**nonredDataFrame**

*Filter for unique elements*

**Description**

`nonredDataFrame` filters 'x' (list of char-vectors or char-vector) for elements unique (to 'ref' or if NULL to all 'x') and of character length. May be used for different 'accession' for same pep sequence (same 'peptide_id'). Note : made for treating data.frames, may be slightly slower than matrix equivalent

**Usage**

```r
nonredDataFrame(
  dataFr, 
  useCol = c(pepID = "peptide_id", protID = "accession", seq = "sequence", mod = "modifications"),
  sepCollapse = "/\",
  callFrom = NULL
)
```

**Arguments**

- **dataFr** (data.frame) main input
- **useCol** (character,length=2) column names of 'dataFr' to use : 1st value designates where redundant values should be gathered; 2nd value designes column of which information should be concatenated
- **sepCollapse** (character) concatenation symbol
- **callFrom** (character) allow easier tracking of messages produced

**Value**

This function returns a data.frame of filtered (fewer lines) with additional 2 columns 'nSamePep' (number of redundant entries) and 'concID' (concatenated content)

**See Also**

- `combineRedBasedOnCol`, `correctToUnique`, `unique`
Examples

df1 <- data.frame(cbind(xA=letters[1:5], xB=c("h","h","f","e","f"), xC=LETTERS[1:5]))
nonRedDataFrame(df1, useCol=c("xB","xC"))

nonRedundLines <- function(dat, callFrom = NULL)
{
  # Function to remove non-redundant lines from a matrix or data frame
  # Usage:
  # nonRedundLines(dat, callFrom = NULL)

  # Arguments:
  # dat: (matrix or data.frame) main input
  # callFrom: (character) allow easier tracking of message(s) produced

  # Value:
  # matrix (or data.frame) without repeated lines (keep 1st occurrence)

  # Description:
  # nonRedundLines reduces complexity of matrix (or data.frame) if multiple consecutive (!) lines
  # with same values. Return matrix (or data.frame) without repeated lines (keep 1st occurrence)

  dat <- dat[apply(dat, 1, function(x) all(x == unique(x)))]

  return(dat)
}

Examples

mat2 <- matrix(rep(c(1,1:3,3,1),2),ncol=2,dimnames=list(letters[1:6],LETTERS[1:2]))
nonRedundLines(mat2)
**normalizeThis**

*Normalize data in various modes*

---

**Description**

Generic normalization of 'dat' (by columns), multiple methods may be applied. The choice of normalization procedures must be done with care, plotting the data before and after normalization may be critical to understanding the initial data structure and the effect of the procedure applied. Inappropriate methods chosen may render interpretation of (further) results incorrect.

**Usage**

```r
normalizeThis(
  dat,
  method = "mean",
  refLines = NULL,
  refGrp = NULL,
  mode = "proportional",
  trimFa = NULL,
  minQuant = NULL,
  sparseLim = 0.4,
  nCombin = 3,
  omitNonAlignable = FALSE,
  maxFact = 10,
  quantFa = NULL,
  expFa = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- **dat**: matrix or data.frame of data to get normalized
- **method**: (character) may be "mean", "median", "NULL", "none", "trimMean", "rowNormalize", "slope", "exponent", "slope2Sections", "vsn"; When NULL or 'none' is chosen the input will be returned
- **refLines**: (NULL or numeric) allows to consider only specific lines of 'dat' when determining normalization factors (all data will be normalized)
- **refGrp**: Only the columns indicated will be used as reference, default all columns (integer or colnames)
- **mode**: (character) may be "proportional", "additive"; decide if normalization factors will be applied as multiplicative (proportional) or additive; for log2-omics data mode="additive" is suggested
- **trimFa**: (numeric, length=1) additional parameters for trimmed mean
minQuant (numeric) only used with method='rowNormalize': optional filter to set all values below given value as NA; see also rowNormalize

sparseLim (integer) only used with method='rowNormalize': decide at which min content of NA values the function should go in sparse-mode; see also rowNormalize

nCombin (NULL or integer) only used with method='rowNormalize': used only in sparse-mode (ie if content of NAs higher than content of sparseLim): Number of groups of smaller matrices with this number of columns to be inspected initially; low values (small groups have higher chances of more common elements); see also rowNormalize

omitNonAlignable (logical) only used with method='rowNormalize': allow omitting all columns which can't get aligned due to sparseness; see also rowNormalize

maxFact (numeric, length=2) only used with method='rowNormalize': max normalization factor; see also rowNormalize

quantFa (numeric, length=2) additional parameters for quantiles to use with method='slope'

expFa (numeric, length=1) additional parameters for method='exponent'

silent (logical) suppress messages

debug (logical) additional messages for debugging

callFrom (character) allows easier tracking of messages produced

Details

In most cases of treating 'Omics'-data one works with the hypothesis that there are no global changes in the structure of all data/columns. Under this hypothesis it is very common to assume the the median (via the argument method) of all samples (ie columns) should remain constant. For examples samples/columns with less signal will be considered as having received 'accidentally' less material (eg due to the imprecision when transferring very small amounts of liquid samples). In consequence, a sample having received only 95 Thus, all measures will be multiplied by 1/0.95 (april 1.053) to compensate for supposed lack of starting material.

With the analysis of 'Omics'-data it is very common to work with data on log-scale. In this case the argument mode should be set to additive, since adding a constant factor to log-data corresponds to a multiplicative factor on regular scale. Please note that (at this point) the methods 'slope', 'exponent', 'slope2Sections' and 'vsn' don't distinguish between additive and proportional modes, but take the data 'as is' (you may look at the original documentation for more details, see exponNormalize, adjBy2ptReg, justvsn).

Normalization using method="rowNormalize" runs rowNormalize from this package. In this case, the working hypothesis is, that all values in each row are expected to be the same. This method could be applied when all series of values (ie columns) are replicate measurements of the same sample. There is also an option for treating sparse data (see argument sparseLim), which may, however, consume much more computational resources, in particular, when the value nCombin is low (compared to the number of samples/columns).

Normalization using method="vsn" runs justvsn from vsn (this requires a minimum of 42 rows of input-data and having the Bioconductor package vsn installed). Note: Depending on the procedure chosen, the normalized data may appear on a different scale.
numPairDeColNames

Value

This function returns a matrix of normalized data (same dimensions as input)

See Also

rowNormalize, exponNormalize, adjBy2ptReg, justvsn

Examples

set.seed(2015); rand1 <- round(runif(300)+rnorm(300,0,2),3)
dat1 <- cbind(ser1=round(100:1+rand1[1:100]), ser2=round(1.2*(100:1+rand1[101:200])-2),
    ser3=round((100:1 +rand1[201:300])^1.2-3))
dat1 <- cbind(dat1, ser4=round(dat1[,1]seq(2,5,length.out=100)+rand1[11:110],1))
dat1[dat1 <1] <- NA
summary(dat1)
    dat1[c(1:5,50:54,95:100),]
no1 <- normalizeThis(dat1, refGrp=1:3, meth="mean")
n02 <- normalizeThis(dat1, refGrp=1:3, meth="trimMean", trim=0.4)
n03 <- normalizeThis(dat1, refGrp=1:3, method="median")
n04 <- normalizeThis(dat1, refGrp=1:3, meth="slope", quantFa=c(0.2,0.8))
dat1[c(1:10,91:100),]
cor(dat1[,3],rowMeans(dat1[,1:2],na.rm=TRUE), use="complete.obs") # high
cor(dat1[,4],rowMeans(dat1[,1:2],na.rm=TRUE), use="complete.obs") # bad
cor(dat1[c(1:10,91:100),4],rowMeans(dat1[c(1:10,91:100),1:2],na.rm=TRUE),use="complete.obs")
cor(dat1[,3],rowMeans(dat1[,1:2],na.rm=TRUE)^ (1/seq(2,5,length.out=100),use="complete.obs")

numPairDeColNames

Extract pair of numeric values from vector or column-names

Description

This function extracts a pair of numeric values out of a vector or colnames (from a matrix). This is useful when pairwise comparisons are concatenated like '10c-100c', return matrix with 'index'=selComp, log2rat and both numeric. Additional white space or character text can be removed via the argument stripTxt. Of course, the separator sep needs to be specified and should not be included to 'stripTxt'.

Usage

numPairDeColNames(
    dat,
    selComp = NULL,
    stripTxt = NULL,
    sep = "-",
    columnLabel = "conc",
    sortByAbsRatio = FALSE,
    silent = FALSE,
    debug = FALSE,
    callFrom = NULL
)
orderMatrToRef

Order Lines of Matrix According to Reference (Character) Vector

Description

This function orders lines of matrix mat according to a (character) reference vector ref. To do so, all columns of mat will be considered to use the first column from left with the best (partial) matching results. This function first looks for unambiguous perfect matches, and if not found successive rounds of more elaborate partial matching will be engaged: In case of no perfect matches found, grep of ref on all columns of mat and/or grep of all columns of mat on ref (ie 'reverse grep') will be applied (finally a 'two way grep' approach). Until a perfect match is found each element of ref will be tested on mat and inversely (for each column) each element of mat will be tested on ref. The approach with the best number of (unique) matches will be chosen. In case of one-to-many matches, it will be tried to use most complete lines (see also last example).
**Usage**

```r
orderMatrToRef(
  mat,
  ref,
  addRef = TRUE,
  listReturn = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `mat` (matrix, data.frame) main input of which rows should get re-ordered according to a (character) reference vector `ref`
- `ref` (character) reference imposing new order
- `addRef` (logical) add `ref` to output as new column
- `listReturn` (logical) allows retrieving more information in form of list
- `silent` (logical) suppress messages
- `debug` (logical) display additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

**Value**

This function returns, depending on `listReturn`, either the input-matrix in new order or a list with `$mat` (the input matrix in new order), `$grep` (matched matrix) and `$col` indicating the column of `mat` finally used.

**See Also**

for basic ordering see `match`; `checkGrpOrder` for testing each line for expected order, `checkStrictOrder` to check for strict (ascending or descending) order

**Examples**

```r
mat1 <- matrix(paste0("__",letters[rep(c(1,1,2,2,3),3) +rep(0:2,each=5)], rep(1:5)), ncol=3)
orderMatrToRef(mat1, paste0(letters[rep(c(3,4,5,3,4),3),c(1,3,5,2,4)]))

mat2 <- matrix(paste0("__",letters[rep(c(1,1,2,2,3),3) +rep(0:2,each=5)],
  c(rep(1:5,2),1,1,3,5)), ncol=3)
orderMatrToRef(mat2, paste0(letters[rep(c(3,4,5,3,4),3),c(1,3,5,1,4)]))

mat3 <- matrix(paste0(letters[rep(c(1,1,2,2,3),3) +rep(0:2,each=5)],
  c(rep(1:5,2),1,1,3,3,5)), ncol=3)
orderMatrToRef(mat3, paste0("__",letters[rep(c(3,4,5,3,4),3),c(1,3,5,1,3)]))
```
organizeAsListOfRepl  (re)organize data of (3-dim) array as list of replicates

Description

Organize array of all data (‘arrIn’, long table) into list of (replicate-)arrays (of similar type/layout) based on dimension number ‘byDim’ of ‘arrIn’ (eg 2nd or 3rd dim). Argument inspNChar defines the number of characters to consider, so if the beginning of names is the same they will be separated as list of multiple arrays. Default will search for ‘_’ separator or trim from end if not found in the relevant dimnames.

Usage

organizeAsListOfRepl(
  arrIn,
  inspNChar = 0,
  byDim = 3,
  silent = TRUE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

arrIn (array) main input
inspNChar (interger) if inspNChar=0 the array-names (2nd dim of ‘arrIn’) will be cut before last ‘_’
byDim (integer, length=1) dimension number along which data will be split in separate elements (considering the first inspNChar characters)
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This function returns a list of arrays (typically 1st and 2nd dim for specific genes/objects, 3rd for different measures associated with)

See Also

array

Examples

arr1 <- array(1:24,dim=c(4,3,2),dimnames=list(c(LETTERS[1:4]),
paste("col",1:3,sep=""), c("ch1","ch2")))
organizeAsListOfRepl(arr1)
packageDownloadStat

Simple Package Download Statistics from CRAN

Description

This function allows accessing the most recent counts of package downloads available on http://www.datasciencemeta.com/rpackages, obtaining rank quantiles and to compare (multiple) given packages to the bulk data, optionally a plot can be drawn.

Usage

packageDownloadStat(
  queryPackages = c("wrMisc", "wrProteo", "cif", "bcv", "FinCovRegularization"),
  countUrl = "http://www.datasciencemeta.com/rpackages",
  refQuant = (1:10)/10,
  figure = TRUE,
  log = "",
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

queryPackages (character or integer) package names of interest, if integer, n random packages will be picked by random

countUrl (character) the url where the daily counts are available

refQuant (numeric) add reference quantile values to output matrix

figure (logical) decide of figure should be printed

log (character) set count-axis of figure to linear or log-scale (by setting log="y")

silent (logical) suppress messages

callFrom (character) allow easier tracking of messages produced

debug (logical) additional messages for debugging

Details

Detailed articles on this subject have been published on R-Hub (https://blog.r-hub.io/2020/05/11/packagerank-intro/) and on R-bloggers (https://www.r-bloggers.com/2020/10/a-cran-downloads-experiment/). The task of checking the number of downloads for a given package has also been addressed by several other packages (eg dlstats, cranlogs, adjustedcranlogs).

This function only allows accessing counts as listed on the website of www.datasciencemeta.com which get updated daily. Please note, that reading all lines from the website may take a few seconds!! To get a better understanding of the counts read, reference quantiles for download-counts get added by default (see argument refQuant). The (optional) figure can be drawn in linear scale
pairsAsPropensMatr

Convert Pairs of Node-Names to Non-Oriented Propensity Matrix

Description

Numerous network query tools produce a listing of pairs of nodes (with one pair of nodes per line). Using this function such a matrix (or data.frame) can be combined to this more comprehensive view as propensity-matrix.

Usage

pairsAsPropensMatr(mat, silent = FALSE, debug = FALSE, callFrom = NULL)

Arguments

- mat (matrix) main input, matrix of interaction partners with each line as a separate pair of nodes; the first two columns should contain identifiers of the nodes
- silent (logical) suppress messages
- debug (logical) additional messages for debugging
- callFrom (character) allow easier tracking of messages produced

Value

This function returns a matrix with download counts (or NULL if the web-site can’t be accessed or the query-packages are not found there)

See Also

packages cranlogs and packageRank

Examples

```r
## Let’s try a microscopic test-file (NOT representative for true up to date counts !!)
pack1 <- c("cif", "bcv", "FinCovRegularization", "wrMisc", "wrProteo")
testFi <- file.path(system.file("extdata", package="wrMisc"), "rpackagesMicro.html")
packageDownloadStat(pack1, countUrl=testFi, log="y", figure=FALSE)
## For real online counting simply drop the argument countUrl
```
Details

Note, this has been primarily developed for undirected interaction networks, the resulting propensity-
matrix does not show any orientation any more. In a number of applications (eg in protein-protein
interaction networks, PPI) the resulting matrix may be rather sparse.

Value

This function returns matrix or data.frame

See Also

uses typically input from filterNetw

Examples

```
pairs3L <- matrix(LETTERS[c(1,3,3, 2,2,1)], ncol=2)  # loop of 3
(netw13pr <- pairsAsPropensMatr(pairs3L))          # as prop matr
```

partialDist  Partial distance matrix (focus on closest)

Description

partialDist calculates distance matrix like dist for 1- or 2-dim data, but only partially, ie only
cases of small distances. This function was made for treating very large data-sets where only very
close distances to a given point need to be found, it allows to overcome memory-problems with
larger data (and faster execution with > 50 rows of 'dat').

Usage

```
partialDist(
  dat,
  groups,
  overLap = TRUE,
  method = "euclidean",
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

dat  (matrix of numeric values) main input

groups  (factor) to split using cut or specific custom grouping (length of dat)

overLap  (logical) if TRUE make groups overlapping by 1 value (ie maintain some context-
information)
partUnlist

method 'character' name of method passed to dist
silent (logical) suppress messages
debug (logical) display additional messages for debugging
callFrom (character) allow easier tracking of message(s) produced

Value

This function returns a matrix with partial distances (not of class 'dist' object)

See Also

dist

Examples

set.seed(2016); mat3 <- matrix(runif(300), nr=30)
round(dist(mat3), 1)
round(partialDist(mat3, gr=3), 1)

partUnlist Partial unlist of lists of lists

Description

partUnlist does partial unlist for treating list of lists : New (returned) list has one level less of hierarchy (Highest level list will be appended). In case of conflicting (non-null) listnames a prefix will be added. Behaviour different to unlist when unlisting list of matrixes.

Usage

partUnlist(lst, sep = "_", silent = FALSE, debug = FALSE, callFrom = NULL)

Arguments

lst (list) main input, list to be partially unlisted
sep (character, length=1) separator for names
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of message(s) produced

Value

This function returns a list with partially reduced nested structure

See Also

unlist, asSepList
Examples

```r
partUnlist(list(list(a=11:12,b=21:24), list(c=101:101,d=201:204)))
li4 <- list(c=1:3, M2=matrix(1:4,ncol=2), L3=list(L1=11:12, M3=matrix(21:26,ncol=2)))
partUnlist(li4)
unlist(li4, rec=FALSE)
```

---

### pasteC

**Advanced paste-collapse**

#### Description

This function is a variant of **paste** for convenient use of paste-collapse and separation of last element to paste (via `lastCol`). This function was made for more human like enumerating in output and messages. If multiple arguments are given without names they will all be concatenated, if they contain names lazy evaluation for names will be tried (with preference to longest match to argument names). Note that some special characters (like backslash) may need to be protected when used with `collapse` or `quoteC`. Returns character vector of length 1 (everything pasted together)

#### Usage

```r
pasteC(..., collapse = "", lastCol = " and ", quoteC = "")
```

#### Arguments

- `...`: (character) main input to be collapsed
- `collapse`: (character,length=1) element to use for collapsing
- `lastCol`: (character) text to use before last item enumerated element
- `quoteC`: (character) text to use for citing with quotations (default "")

#### Value

This function returns a character vector of truncated versions of input text

#### See Also

- **paste** for basic paste

#### Examples

```r
pasteC(1:4)
```
Description

This function produces a logical matrix to be used as filter for lines of ‘dat’ for sufficient presence of non-NA values (ie limit number of NAs per line). Filter abundance/expression data for min number and/or ratio of non-NA values in at least 1 of multiple groups. This type of procedure is common in proteomics and transcriptomics, where a NA can many times be associated with quantitation below detection limit.

Usage

```r
presenceFilt(
  dat,
  grp,
  maxGrpMiss = 1,
  ratMaxNA = 0.8,
  minVal = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

dat matrix or data.frame (abundance or expression-values which may contain some NAs).

grp factor of min 2 levels describing which column of ‘dat’ belongs to which group (levels 1 & 2 will be used)

maxGrpMiss numeric at least 1 group has not more than this number of NAs (otherwise mark line as bad)

ratMaxNA numeric at least 1 group reaches this content of non-NA values

minVal default NULL or numeric, any value below will be treated like NA

silent (logical) suppress messages

debug (logical) additional messages for debugging

callFrom (character) allow easier tracking of messages produced

Value

logical matrix (with separate col for each pairwise combination of ‘grp’ levels) indicating if line of ‘dat’ acceptable based on NAs (and values minVal)
presenceGrpFilt

Filter for each group of columns for sufficient data as non-NA

Description

The aim of this function is to filter for each group of columns for sufficient data as non-NA.

Usage

presenceGrpFilt(dat, grp, presThr = 0.75, silent = FALSE, callFrom = NULL)

Arguments

dat  matrix or data.frame (abundance or expression-values which may contain some NAs).
grp  factor of min 2 levels describing which column of 'dat' belongs to which group (levels 1 & 2 will be used)
presThr  (numeric) min ratio of non-NA values (per group) for returning a given line & group as TRUE
silent  (logical) suppress messages
callFrom  (character) allow easier tracking of messages produced
Details

This function allows to identify lines with an NA-content above the threshold presThr per group as defined by the levels of factor grp. With different types of projects/questions different threshold presThrs per group rather low, one could use a value of 0.75 (ie >= 75

Value

logical matrix (with on column for each level of grp)

See Also

presenceFilt, there are also other packages totally dedicated to filtering on CRAN and Bioconductor

Examples

mat <- matrix(NA, nrow=11, ncol=6)
mat[lower.tri(mat)] <- 1
mat <- cbind(mat, mat[,1:4])
colnames(mat) <- c(paste0("re",1:6), paste0("x",1:4))

## accept 1 NA out of 4, 2 NA out of 6 (ie certainly present)
(filt0a <- presenceGrpFilt(mat, rep(1:2, c(6,4)), pres=0.66))

## accept 2 NA out of 4, 2 NA out of 6 (ie min 50% present)
(filt0b <- presenceGrpFilt(mat, rep(1:2, c(6,4)), pres=0.5))

## accept 3 NA out of 4, 4 NA out of 6 (ie possibly present)
(filt0c <- presenceGrpFilt(mat, rep(1:2, c(6,4)), pres=0.19))

protectSpecChar

Protect Special Characters

Description

Some characters do have a special meaning when used with regular expressions. This concerns characters like a point, parenthesis, backslash etc. Thus, when using grep or any related command, such special characters must get protected in order to get considered as they are.

Usage

protectSpecChar(
  x,
  prot = c(".", "\", "|", "("", ")", "[", "]", "{", "^", ",", "+", "?"),
  silent = TRUE,
  callFrom = NULL
)
**Arguments**

- **x** (character) character vector to be prepared for use in regular expressions
- **prot** (character) collection of characters that need to be protected
- **silent** (logical) suppress messages
- **callFrom** (character) allow easier tracking of messages produced

**Value**

This function returns a modified character vector

**Examples**

```r
aa <- c("abc","abcde","ab.c","ab.c.e","ab*c","ab\d")
grepl("b.", aa) # all TRUE
grepl("b\\.", aa) # manual protection
grepl(protectSpecChar("b."), aa)
```

---

**Description**

This function takes a numeric vector of p-values and returns a vector of lfdr-values (local false discovery) using the package `fdrtool`. Multiple testing correction should be performed with caution, short series of p-values typically pose problems for transforming to lfdr. The transformation to lfdr values may give warning messages, in this case the resultant lfdr values may be invalid!

**Usage**

```r
pVal2lfdr(x, silent = TRUE, callFrom = NULL)
```

**Arguments**

- **x** (numeric) vector of p.values
- **silent** (logical) suppress messages
- **callFrom** (character) allow easier tracking of messages produced

**Value**

This function returns a (numeric) vector of lfdr values (or NULL if data insufficient to run the function 'fdrtool')

**See Also**

lfdr from `fdrtool`, other p-adjustments (multiple test correction, eg FDR) in `p.adjust`
Examples

```r
## Note that this example is too small for estimating really meaningful fdr values
## In consequence, a warning will be issued.
set.seed(2017); t8 <- matrix(round(rnorm(160,10,0.4),2), ncol=8,
dimnames=list(letters[1:20], c("AA1","BB1","CC1","DD1","AA2","BB2","CC2","DD2")))
t8[3:6,1:2] <- t8[3:6,1:2]+3  # augment lines 3:6 (c-f) for AA1&BB1
t8[5:8,5:6] <- t8[5:8,5:6]+3  # augment lines 5:8 (e-h) for AA2&BB2 (c,d,g,h should be found)
head(pVal2lfdr(apply(t8, 1, function(x) t.test(x[1:4], x[5:8])$p.value)))
```

### randIndFx

**Distance of categorical data (Jaccard, Rand and adjusted Rand index)**

**Description**

randIndFx calculates distance of categorical data (as Rand Index, Adjusted Rand Index or Jaccard Index). Note: uses/requires package flexclust Methods so far available (via flexclust): "ARI" .. adjusted Rand Index, "RI" .. Rand index, "J" .. Jaccard, "FM" .. Fowlkes-Mallows.

**Usage**

```r
randIndFx(ma, method = "ARI", adjSense = TRUE, silent = FALSE, callFrom = NULL)
```

**Arguments**

- `ma`: (matrix) main input for distance calculation
- `method`: (character) name of distance method (eg "ARI","RI","J","FM")
- `adjSense`: (logical) allows introducing correlation/anticorrelation (interprete neg distance results as anti)
- `silent`: (logical) suppres messages
- `callFrom`: (character) allow easier tracking of messages produced

**Value**

This function returns a distance matrix

**See Also**

comPart in randIndex

**Examples**

```r
set.seed(2016); tab2 <- matrix(sample(1:2, size=42, replace=TRUE), ncol=7)
if(requireNamespace("flexclust")) { flexclust::comPart(tab2[1,], tab2[2,])
  flexclust::comPart(tab2[1,], tab2[3,])
  flexclust::comPart(tab2[1,], tab2[4,]) }
## via randIndFx():
randIndFx(tab2, adjSense=FALSE)
cor(t(tab2))
randIndFx(tab2, adjSense=TRUE)
```
rankToContigTab  Contingency tables for fit of ranking

Description

Count the number of instances where the corresponding columns of 'dat' have a value matching the group number as specified by 'grp'. Counting will be performed/repeated independently for each line of 'dat'. Returns array (1st dim is rows of dat, 2nd is unique(grp), 3rd dim is ok/bad), these results may be tested using eg fisher.test. This function was made for prearing to test the ranking of multiple features (lines in 'mat') including replicates (levels of 'grp').

Usage

rankToContigTab(dat, grp)

Arguments

dat  (matrix or data.frame of integer values) ranking of multiple features (lines), equal ranks may occur

grp  (integer) expected ranking

Value

array (1st dim is rows of dat, 2nd is unique(grp), 3rd dim is ok/bad)

See Also

lm

Examples

# Let's create a matrix with ranks (equal ranks do occur)
ma0 <- matrix(rep(1:3,each=6), ncol=6, dimnames=list(
  c("li1","li2","ref"), letters[1:6]))
ma0[1,6] <- 1  # create item not matching correctly
ma0[2,] <- c(3:1,2,1,3)  # create items not matching correctly
gr0 <- gl(3,2)  # the expected ranking (as duplicates)
(count0 <- rankToContigTab(ma0,gr0))
(cTab <- t(apply(count0, c(1,3) ,sum))
# Now we can compare the ranking of line1 to ref ...  
fisher.test(cTab[,c(3,1)])  # test li1 against ref
fisher.test(cTab[,c(3,2)])  # test li2 against ref
ratioAllComb

Calculate all ratios between x and y

Description

This function calculates all possible pairwise ratios between all individual values of x and y, or samples up to a maximum number of combinations.

Usage

ratioAllComb(
  x,
  y,
  maxLim = 10000,
  isLog = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

x (numeric) vector, numerator for constructing ratios
y (numeric) vector, denominator for constructing ratios
maxLim (integer) allows reducing complexity by drawing for very long x or y
isLog (logical) adjust ratio calculation to log-data
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Examples

set.seed(2014); ra1 <- c(rnorm(9,2,1),runif(8,1,2))
ratioAllComb(ra1[1:9],ra1[10:17])
boxplot(list(norm=ra1[1:9], unif=ra1[10:17], rat=ratioAllComb(ra1[1:9],ra1[10:17])))
Description

This function transforms ratio 'x' to ppm (parts per million). If 'y' not given (or different length as 'x'), then 'x' is assumed as ratio otherise rations are constructed as x/y is used lateron. Does additional checking : negative values not expected - will be made absolute!

Usage

```r
ratioToPpm(
  x,
  y = NULL,
  nSign = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `x` (numeric) main input
- `y` (numeric) optional value to construct ratios (x/y). If NULL (or different length as 'x'), then 'x' will be considered as ratio.
- `nSign` (numeric) number of significan digits
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced

Value

This function returns a numeric vector of ppm values

See Also

`XYToDiffPpm` for ppm of difference as used in mass spectrometrie

Examples

```r
set.seed(2017); aa <- c(1.000001,0.999999,1+rnorm(10,0,0.001))
cbind(x=aa, ppm=ratioToPpm(aa, nSign=4))
```
**readCsvBatch**

**Read batch of csv-files**

**Description**

This function was designed to read screening data split in parts (with common structure) and saved to multiple files, to extract the numeric columns and to compile all (numeric) data to a single array (or list). Some screening platforms save results while progressing through a pile of microtiter-plates separately. The organization of the resultant files is structured through file-names and all files have exactly the same organization of lines and columns/ European or US-formatted csv files can be read, if argument fileFormat is NULL both types will be tested, otherwise it allows to specify a given format. The presence of headers (to be used as column-names) may be tested using checkFormat.

**Usage**

```r
readCsvBatch(
  fileNames = NULL,
  path = ".",
  fileFormat = "Eur",
  checkFormat = TRUE,
  returnArray = TRUE,
  columns = c("Plate", "Well", "StainA"),
  excludeFiles = "All infected plates",
  simpleNames = TRUE,
  minNamesLe = 4,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

**Arguments**

- **fileNames** (character) names of files to be read, if NULL all files fitting 'fileFormat'
- **path** (character) where files should be read (folders should be written in R-style)
- **fileFormat** (character) may be NULL (both US and European formats will be tried), 'Eur' or 'US'
- **checkFormat** (logical) if TRUE: check header, remove empty columns, 1st line if all empty, set output format for each file to matrix, if rownames are increasing integeres try to use 2nd of 'columns' as rownames
- **returnArray** (logical) allows switching from array to list-output
- **columns** (NULL or character) column-headers to be extracted (if specified), 2nd value may be columnw with rownames (if rownames are encountered as increasing rownames)
- **excludeFiles** (character) names of files to exclude (only used when reading all files of given directory)
simpleNames  (logical) allows truncating names (from beginning) to get to variable part (using .trimFromStart()), but keeping 'minNamesLe'
minNamesLe (integer) min length of column-names if simpleNames=TRUE
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allows easier tracking of messages produced

Value

This function returns an array (or list if returnArray=FALSE) of all numeric data read (numerical columns only) from individual files

See Also

read.table, writeCsv, readXlsxBatch

Examples

path1 <- system.file("extdata", package="wrMisc")
fiNa <- c("pl01_1.csv", "pl01_2.csv", "pl02_1.csv", "pl02_2.csv")
datAll <- readCsvBatch(fiNa, path1)
str(datAll)
## batch reading of all csv files in specified path :
datAll2 <- readCsvBatch(fileNames=NULL, path=path1, silent=TRUE)

Description

This function allows batch reading of multiple tabulated text files in batch. The files can be designed specifically, or, alternatively all files from a given directory can be read. If package data.table is available, faster reading of files will be performed using the function fread.

Usage

readTabulatedBatch(
  query,
  path = NULL,
  dec = ".", 
  header = "auto", 
  strip.white = FALSE, 
  blank.lines.skip = TRUE, 
  fill = FALSE, 
  filtCol = 2, 
  filterAsInf = TRUE, 
  filtVal = 5000,
silent = FALSE,  
callFrom = NULL,  
debug = FALSE)
}

Arguments

query (character) vector of file-names to be read, if "." all files will be read (no matter what their extension might be)

path (character) path for reading files, if NULL or NA the current directory will be used

dec (character, length=1) decimals to use, will be passed to fread or read.delim

header (character, length=1) path for reading files, if NULL or NA the current directory will be used, will be passed to fread or read.delim

strip.white (logical, length=1) Strips leading and trailing whitespaces of unquoted fields, will be passed to fread or read.delim

blank.lines.skip (logical, length=1) If TRUE blank lines in the input are ignored. will be passed to fread or read.delim

fill (logical, length=1) If TRUE then in case the rows have unequal length, blank fields are implicitly filled, will be passed to fread or read.delim

filtCol (integer, length=1) which columns should be used for filtering, if NULL or NA all data will be returned

filterAsInf (logical, length=1) filter as inferior or equal (TRUE) or superior or equal threshold

filtVal (numeric, length=1) which numeric threshold should be used for filtering, if NULL or NA all data will be returned

silent (logical) suppress messages

callFrom (character) allow easier tracking of messages produced

debug (logical) display additional messages for debugging

Details

If you want to provide a flexible pattern of file-names, this has to be done before calling this function, eg using grep to provide an explicit collection of files. However, it is possible to read different files from different locations/directories, the length of path must match the length of query

Value

This function returns a list of data.frames

See Also

fread, read.delim, for reading batch of csv files: readCsvBatch
Examples

```r
path1 <- system.file("extdata", package="wrMisc")
fiNa <- c("a1.txt","a2.txt")
allTxt <- readTabulatedBatch(fiNa, path1)
str(allTxt)
```

Description

Reading the content of files where the number of separators (eg tabulation) is variable poses problems with traditional methods for reding files, like `read.table`. This function reads each line independently and then parses all separators therein. The first line is assumed to be column-headers. Finally, all data will be returned in a matrix adopted to the line with most separators and if the number of column-headers is insufficient, new (unique) column-headers will be generated. Thus, the lines may contain different number of elements, empty elements (ie tabular fields) will always get added to right of data read and their content will be as defined by argument `emptyFields` (default NA).

Usage

```r
readVarColumns(
  fiName,
  path = NULL,
  sep = "\t",
  header = TRUE,
  emptyFields = NA,
  refCo = NULL,
  supNa = NULL,
  silent = FALSE,
  callFrom = NULL
)
```

Arguments

- `fiName` (character) file-name
- `path` (character) optional path
- `sep` (character) separator (between columns)
- `header` (logical) indicating whether the file contains the names of the variables as its first line.
- `emptyFields` (NA or character) missing headers will be replaced by the content of 'emptyFields', if NA the last column-name will be re-used and a counter added
- `refCo` (integer) for custom choice of column to be used as row-names (default will use 1st text-column)
supNa (character) base for constructing name for columns wo names (+counter starting at 2), default column-name to left of 1st col wo colname
silent (logical) suppress messages
callFrom (character) allow easier tracking of messages produced

Details
Note, this functions assumes one line of header and at least one line of data! Note, for numeric data the comma is assumed to be US-Style (as '.'). Note, that it is assumed, that any missing fields for the complete tabular view are missing on the right (ie at the end of line)!

Value
This function returns a matrix (character or numeric)

See Also
for regular 'complete' data read.table

Examples
path1 <- system.file("extdata",package="wrMisc")
fiNa <- "Names1.tsv"
datAll <- readVarColumns(fileName=file.path(path1,fiNa))
str(datAll)

readXlsxBatch  Read Batch of Excel xlsx-Files

Description
readXlsxBatch reads data out of multiple xlsx files, the sheet indicated by 'sheetInd' will be considered. All files must have the same organization of data, as this is typically the case when high-throughput measurements are automatically saved while experiments progress. In particular, the first file read is used to structure the output.

Usage
readXlsxBatch(
  fileNames = NULL,
  path = ".",
  fileExtension = "xlsx",
  excludeFiles = NULL,
  sheetInd = 1,
  checkFormat = TRUE,
  returnArray = TRUE,
  columns = c("Plate", "Well", "StainA"),
)
Arguments

fileNames  (character) provide either explicit list of file-names to be read or leave NULL for reading all files ending with ‘xlsx’ in path specified with argument path

path  (character) there may be a different path for each file

fileExtension  (character) extension of files (default='xlsx')

excludeFiles  (character) names of files to exclude (only used when reading all files of given directory)

sheetInd  (character or integer) specify which sheet to extract (as exact name of sheet or sheet-number, eg sheetInd=2 will extract always the 2nd sheet (no matter the name); if given as sheet-name but nor present in file an empty list-elements will be returned

checkFormat  (logical) if TRUE: check header, remove empty columns, if rownames are increasing integers it will search for first column with different entries to use as rownames

returnArray  (logical) allows switching from array to list-output

columns  (NULL or character) column-headers to be extracted (if specified, otherwise all columns will be extracted)

simpleNames  (integer), if NULL all characters of fileNames will be maintained, otherwise allows truncating names (from beginning) to get to variable part (using .trimFromStart()), but keeping at least the number of characters indicated by this argument

silent  (logical) suppress messages

callFrom  (character) allows easier tracking of messages produced

Details

By default all columns with text-content may be eliminated to keep the numeric part only, which may then get organized to a 3-dim numeric array (where the additional files will be used as 2nd dimension and multiple columns per file shown as 3rd dimension).

NOTE: (starting from version wrMisc-1.5.5) requires packages readxl and Rcpp being installed! (This allows much faster and memory efficient processing than previous use of package ‘xlsx’)

Value

This function returns a list of data.frames

See Also

read_excel; for simple reading of (older) xls-files under 32-bit R one may also see the package RODBC
Examples

```r
path1 <- system.file("extdata", package="wrMisc")
fiNa <- c("pl01_1.xlsx","pl01_2.xlsx","pl02_1.xlsx","pl02_2.xlsx")
datAll <- readXlsxBatch(fiNa, path1)
str(datAll)
## Now let's read all xlsx files of directory
datAll2 <- readXlsxBatch(path=datAll, silent=TRUE)
identical(datAll, datAll2)
```

---

**reduceTable**

Reduce table by aggregating smaller groups

Description

reduceTable treats/reduces results from `table` to 'nGrp' groups, optional indiv resolution of 'sep-
marFirst' (numeric or NULL). Mainly made for reducing the number of classes for better plots with `pie`

Usage

```r
reduceTable(tab, separFirst = 4, nGrp = 15)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>tab</code></td>
<td>output of <code>table</code></td>
</tr>
<tr>
<td><code>separFirst</code></td>
<td>(integer or NULL) optinal separation of n <code>separFirst</code> groups (value &lt;2 or NULL will privilege more uniform size of groups, higher values will cause small initial and larger tailing groups)</td>
</tr>
<tr>
<td><code>nGrp</code></td>
<td>(integer) number of groups expected</td>
</tr>
</tbody>
</table>

Value

This function returns a numeric vector with number of counts and class-borders as names (like `table`).

See Also

`table`

Examples

```r
set.seed(2018); dat <- sample(11:60,200,repl=TRUE)
pie(table(dat))
pie(reduceTable(table(dat), sep=NULL))
pie(reduceTable(table(dat), sep=NULL), init.angle=90,
clockwise=TRUE, col=rainbow(20)[1:15], cex=0.8)
```
regrBy1or2point

Rescaling according to reference data using linear regression.

Description

regrBy1or2point does rescaling: linear transform simple vector 'inDat' that (mean of) elements of names cited in 'refLst' will end up as values 'regrTo'. Regress single vector according to 'refLst' (describing names of inDat). If 'refLst' contains 2 groups, the 1st group will be set to the 1st value of 'regrTo' (and the 2nd group of 'refLst' to the 2nd 'regrTo')

Usage

regrBy1or2point(
  inDat,
  refLst,
  regrTo = c(1, 0.5),
  silent = FALSE,
  callFrom = NULL
)

Arguments

inDat matrix or data.frame
refLst list of names existing in inDat (one group of names for each value in 'regrTo'), to be transformed in values precised in 'regrTo'; if no matches to names of 'inDat' found, the 2 lowest and/or highest highest values will be chosen
regrTo (numeric,length=2) range (at scale 0-1) of target-values for mean of elements cited in 'refLst'
silent (logical) suppress messages
callFrom (character) allows easier tracking of message(s) produced

Value

normalized matrix

See Also

adjBy2ptReg, regrMultBy1or2point

Examples

set.seed(2016); dat1 <- 1:50 +(1:50)*round(runif(50),1)
names(dat1) <- 1:length(dat1)
reg1 <- regrBy1or2point(dat1,refLst=c("2","49"))
plot(reg1,dat1)
regrMultBy1or2point

Rescaling of multiple data-sets according to reference data using regression

Description

regrMultBy1or2point regresses each col of matrix according to 'refLst' (describing rownames of inDat). If 'refLst' contains 2 groups, the 1st group will be set to the 1st value of 'regrTo' (and the 2nd group of 'refLst' to the 2nd 'regrTo')

Usage

regrMultBy1or2point(
  inDat,
  refLst,
  regrTo = c(1, 0.5),
  silent = FALSE,
  callFrom = NULL
)

Arguments

inDat  matrix or data.frame
refLst  list of names existing in inDat (one group of names for each value in 'regrTo'), to be transformed in values precised in 'regrTo'; if no matches to names of 'inDat' found, the 2 lowest and/or highest highest values will be chosen
regrTo  (numeric, length=2) range (at scale 0-1) of target-values for mean of elements cited in 'refLst'
silent  (logical) suppress messages
callFrom  (character) allow easier tracking of message(s) produced

Value

normalized matrix

See Also

adjBy2ptReg, regrBy1or2point

Examples

set.seed(2016); dat2 <- round(cbind(1:50 + (1:50) * runif(50), 2.2 * (1:50) + rnorm(50, 0, 3)), 1)
rownames(dat2) <- 1:nrow(dat2)
reg1 <- regrBy1or2point(dat2[,1], refLst=list(as.character(5:7), as.character(44:45)))
reg2 <- regrMultBy1or2point(dat2, refLst=list(as.character(5:7), as.character(44:45)))
plot(dat2[,1], reg2[,1])
identical(reg1, reg2[,1])
identical(dat2[,1], reg2[,1])
renameColumns  

**Description**

This function renames columns of 'refMatr' using 2-column matrix (or data.frame) indicating old and new names (for replacement).

**Usage**

```
renameColumns(refMatr, newName, silent = FALSE, debug = FALSE, callFrom = NULL)
```

**Arguments**

- `refMatr`: matrix (or data.frame) where column-names should be changed
- `newName`: (matrix of character) giving correspondence of old to new names (number of lines must match number of columns of 'refMatr')
- `silent`: (logical) suppres messages
- `debug`: (logical) additional messages for debugging
- `callFrom`: (character) allows easier tracking of messages produced

**Value**

This function returns a matrix (or data.frame) with renamed columns

**Examples**

```
ma <- matrix(1:8,ncol=4,dimnames=list(1:2,LETTERS[1:4]))
replBy1 <- cbind(new=c("dd","bb","z_"),old=c("D","B","zz"))
replBy2 <- matrix(c("D","B","zz","dd","bb","z_"),ncol=2)
replBy3 <- matrix(c("X","Y","zz","xx","yy","z_"),ncol=2)
renameColumns(ma,replBy1)
renameColumns(ma,replBy2)
renameColumns(ma,replBy3)
```

reorgByCluNo  

**Description**

Reorganize input matrix as sorted by cluster numbers (and geometric mean) according to vector with cluster names, and index for sorting per cluster and per geometric mean. In case mat is an array, the 3rd dimension will be considered as 'column' with arguments useColumn (and cluNo, if it designs a 'column' of mat).
Usage

reorgByCluNo(
  mat,
  cluNo,
  useColumn = NULL,
  meanCol = NULL,
  addInfo = TRUE,
  retList = FALSE,
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

mat (matrix or data.frame) main input
cluNo (positive integer, length to match nrow(mat) initial cluster numbers for each line of ‘mat’ (obtained by separate clustering or other segmentation) or may desinn column of mat to use as cluster-numbers
useColumn (character or integer) the columns to use from mat as main data (default will use all, exept cluCol and/or meanCol if they design columns))
meanCol (character or integer) alternative summarizing data for intra-cluster sorting (instead of geometric mean)
addInfo (logical) allows adding of columns ‘index’, ‘geoMean’ and ‘cluNo’ (or array if FALSE)
retList (logical) return as list of matrixes (or array if FALSE)
silent (logical) suppress messages
callFrom (character) allow easier tracking of messages produced
debug (logical) additional messages for debugging

Value

This function returns a list or array (as 2- or 3 dim) with possible number of occurrences for each of the 3 elements in nMax. Read results vertical : out[[1]] or out[,1] .. (multiplicative) table for 1st element of nMax; out[,2] .. for 2nd

See Also

pairwise combinations combn, clustering kmeans

Examples

dat1 <- matrix(round(runif(24),2), ncol=3, dimnames=list(NULL,letters[1:3]))
clu <- stats::kmeans(dat1, 5)$cluster
reorgByCluNo(dat1, clu)

dat2 <- cbind(dat1, clu=clu)
reorgByCluNo(dat2, "clu")
replicateStructure

Search and Select Groups of Replicates

Description

This function was designed for mining annotation information organized in multiple columns to identify the (potential) grouping of multiple samples, i.e., to determine factor levels. The argument method allows further fine-tuning if high or low number of groups should be preferred, if multiple columns may be combined, or to choose a particular custom column for designating factor levels.

Usage

replicateStructure(
  x,
  method = "median",
  sep = "__",
  exclNoRepl = TRUE,
  trimNames = FALSE,
  includeOther = FALSE,
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

x (matrix or data.frame) the annotation to inspect; each column is supposed to describe another set of annotation/metadata for the rows of x (min 1 row and 1 column),

method (character, length=1) the procedure to choose column(s) with properties of information, may be highest or max (max number of levels) lowest or min (min number of levels), median (median of all options for number of levels), combAll (combine all columns of x) or combNonOrth (combine only non-orthogonal columns of x, to avoid n lines with n levels); lazy evaluation of the argument is possible

sep (character) separator used when a method combining multiple columns (eg combAll, combNonOrth) is chosen (should not appear anywhere in x)

exclNoRepl (logical) decide whether columns with all values different (ie no replicates or max divergency) should be excluded

trimNames (logical) optional trimming of names in x by removing redundant heading and tailing text

includeOther (logical) include $allCols with pattern of (all) other columns

silent (logical) suppress messages

callFrom (character) allow easier tracking of messages produced

debug (logical) additional messages for debugging
Details

Statistical tests require specifying which samples should be considered as replicates of whom. In some cases, like the Sdrf-format, automatic mining of such annotation to indentify an experiment’s underlying structure of replicates may be challanging, since the key information may not always be found in the same column. For this reason this function allows inspecting all columns of a matrix of data.frame to identify which columns may serve describing groups of replicates.

The argument exclNoRepl=TRUE allows excluding all columns with different content for each line (like line-numbers), ie information without any replicates. It is set by default to TRUE to exclude such columns, since statistical tests usually do require some replicates.

When using as method="combAll", there is risk all lines (samples) will be be considered different and no replicates remain. To avoid this situation the argument can be set to method="combNonOrth". Using this mode it will be checked if adding more columns will lead to complete loss of replicates, and -if so- concerned columns omitted.

Value

This function returns a list with $col (column index relativ to x), $lev (abstract labels of level), $meth (note of method finally used) and $allCols with general replicate structure of all columns of x.

See Also
duplicated, uses trimRedundText

Examples

```r
## a is all different, b is groups of 2,  
## c & d are groups of 2 nut NOT 'same general' pattern as b 
strX <- data.frame(a=letters[18:11], b=letters[rep(c(3:1,4), each=2)],  
c=letters[rep(c(5,8:6), each=2)], d=letters[c(1:2,1:3,3:4,4)],  
e=letters[rep(c(4,8,4,7),each=2)], f=rep("z",8) )
strX
replicateStructure(strX[,1:2])
replicateStructure(strX[,1:4], method="combAll")
replicateStructure(strX[,1:4], method="combAll", exclNoRepl=FALSE)
replicateStructure(strX[,1:4], method="combNonOrth", exclNoRepl=TRUE)
replicateStructure(strX, method="lowest")
replicateStructure(strX, method=3, includeOther=TRUE)  # custom choice of 3rd column
```

replNAbyLow Replace NAs by low values
Description

With several screening techniques used in high-throughput biology values at/below detection limit are returned as NA. However, the resultant NA-values may be difficult to analyse properly, simply ignoring NA-values may not be a good choice. When (technical) replicate measurements are available, one can look for cases where one gave an NA while the other did not with the aim of investigating such 'NA-neighbours'. replNAbyLow locates and replaces NA values by (random) values from same line & same group 'grp'. The origin of NAs should be predominantly absence of measure (quantitation) due to signal below limit of detection and not saturation at upper detection limit or other technical problems. Note, this approach may be not optimal if the number of NA-neighbours is very low. Replacamet is done -depending on argument 'unif'- by Gaussian random model based on neighbour values (within same group), using their means and sd, or a uniform random model (min and max of neighbour values). Then numeric matrix (same dim as 'x') with NA replaced is returned.

Usage

replNAbyLow(
  x, 
  grp, 
  quant = 0.8, 
  signific = 3, 
  unif = TRUE, 
  absOnly = FALSE, 
  seed = NULL, 
  silent = FALSE, 
  callFrom = NULL 
)

Arguments

  x (numeric matrix or data.frame) main input
  grp (factor) to organize replicate columns of (x)
  quant (numeric) quantile form 'neighbour' values to use as upper limit for random values
  signific number of signif digits for random values
  unif (logical) toggle between uniform and Gaussian random values
  absOnly (logical) if TRUE, make negative NA-replacement values positive as absolute values
  seed (integer) for use with set.seed for reproducible output
  silent (logical) suppress messages
  callFrom (character) allow easier tracking of message(s) produced

Value

numeric matrix (same dim as 'x') with NA replaced
See Also

naOmit, na.fail

Examples

dat <- matrix(round(rnorm(30),2),ncol=6); grD <- gl(2,3)
dat[sort(sample(1:30,9,repl=FALSE))] <- NA
dat; replNAbyLow(dat,gr=grD)

replPlateCV

CV of replicate plates (list of matrixes)

Description

replPlateCV gets CVs of replicates from list of 2 or 3-dim arrays (where 2nd dim is replicates, 3rd dim may be channel). Note: all list-elements of must MUST have SAME dimensions! When treating data from microtiter plates (eg 8x12) data are typically spread over multiple plates, ie initial matrices that are the organized into arrays. Returns matrix or array (1st dim is intraplate-position, 2nd .. plate-group/type, 3rd .. channels)

Usage

replPlateCV(lst, callFrom = NULL)

Arguments

lst
list of matrixes: suppose lines are independent elements, columns are replicates of the 1st column. All matrixes must have same dimensions

callFrom
(character) allows easier tracking of messages produced

Value

matrix or array (1st dim is intraplate-position, 2nd .. plate-group/type, 3rd .. channels)

See Also

rowCVs, @seealso arrayCV

Examples

set.seed(2016); ral <- matrix(rnorm(3*96),nrow=8)
pla1 <- list(ral[,1:12],ral[,13:24],ral[,25:36])
replPlateCV(pla1)
arrL1 <- list(a=array(as.numeric(ral)[1:192],dim=c(8,12,2)),
             b=array(as.numeric(ral)[97:288],dim=c(8,12,2)))
replPlateCV(arrL1)
Description

`rmDupl2colMatr` removes lines of matrix that are redundant /duplicated for 1st and 2nd column (irrespective of content of their columns). The first occurrence of redundant /duplicated elements is kept.

Usage

```r
dlDupl2colMatr(mat, useCol = c(1, 2))
```

Arguments

- `mat` (matrix or data.frame) main input
- `useCol` (integer, length=2) columns to consider/use when looking for duplicated entries

Value

matrix with duplicated lines removed

See Also

`unlist`

Examples

```r
mat <- matrix(1:12, ncol=3)
mat[3,1:2] <- mat[1,1:2]
rmDupl2colMatr(mat)
```

Description

This function allows identifying, removing or renaming enumerator tag/name (or remove entire enumerator) from tailing enumerators (eg 'abc_No1' to 'abc_1'). A panel of potential candidates as combination of separator-symbols and separtor text/words will be tested to find if one matches all data. In case the main input is a matrix, all columns will be tested independently to find the first column where one specific combination of separator-symbols and separtor text/words is found. Several options exist for the output, the combination of separator-symbols and separtor text/words may be included, too.
Usage

```r
rmEnumeratorName(
  dat,
  nameEnum = c("Number", "No", ",#", "Replicate", "Sample"),
  sepEnum = c(" ", ",", ",_"),
  newSep = ",",
  incl = c("anyCase", "trim2"),
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- **dat** (character vecor or matrix) main input
- **nameEnum** (character) potential enumerator-names
- **sepEnum** (character) potential separators for enumerator-names
- **newSep** (character) potential enumerator-names
- **incl** (character) options to include further variants of the enumerator-names, use "rmEnum" for completely removing enumerator tag/name and digits for different options of trimming names/tags from nameEnum one may use anyCase, trim3 (trimming down to max 3 letters), trim2 (trimming to max 2 letters) or trim1 (trimming down to single letter); trim@ works like trim1 but also includes ",", ie no enumerator tag/name in front of the digit(s)
- **silent** (logical) suppress messages
- **debug** (logical) display additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced

Details

Please note, that checking a variety of different separator text-word and separator-symbols may give an important number of combinations to check. In particular, when automatic trimming of separator text-words is added (eg incl="trim2"), the complexity of associated searches increases quickly. Thus, with large data-sets restricting the content of the arguments nameEnum, sepEnum and (in particular) newSep to the most probable terms/options is suggested to help reducing demands on memory and CPU.

In case the input dat is a matrix and multiple different numerator-types are found, only the first colum (from the left) will be treated. If you which to remove/subsitute mutiple types of enumerators the function `rmEnumeratorName` must be run independently, see last example below.

Value

This function returns a corrected vector (or matrix), or a list if incl="rmEnumL" containing $dat (corrected data), $pattern (the combination of separator-symbols and separaror text/words found), and if input is matrix $column (which column of the input was identified and treated)
See Also

when the exact pattern is known grep and sub may allow direct manipulations much faster

Examples

```r
xx <- c("hg_Re1","hjRe2_Re2","hk-Re3_Re33")
rnEnumeratorName(xx)
rnEnumeratorName(xx, newSep="--")
rnEnumeratorName(xx, incl="anyCase")

xy <- cbind(a=11:13, b=c("11#11","2_No2","333_samp333"), c=xx)
rnEnumeratorName(xy)
rnEnumeratorName(xy, incl=c("anyCase","trim2","rmEnumL"))

xz <- cbind(a=11:13, b=c("23#11","4#2","567#333"), c=xx)
apply(xz, 2, rnEnumeratorName, sepEnum=c("","-_"), newSep="_", silent=TRUE)
```

---

**rnormW**

Normal random number generation with close fit to expected mean and sd

---

**Description**

This function allows creating a vector of random values similar to rnorm, but resulting value get recorrected to fit to expected mean and sd. When the number of random values to generate is low, the mean and sd of the resultant values may deviate from the expected mean and sd when using the standard rnorm function. In such cases the function rnormW helps getting much closer to the expected mean and sd.

**Usage**

```r
rnormW(
  n,
  mean = 0,
  sd = 1,
  seed = NULL,
  digits = 8,
  silent = FALSE,
  callFrom = NULL
)
```

**Arguments**

- `n` (integer, length=1) number of observations. If length(n) > 1, the length is taken to be the number required.
- `mean` (numeric, length=1) expected mean
sd (numeric, length=1) expected sd
seed (integer, length=1) seed for generating random numbers
digits (integer, length=1 or NULL) number of significant digits for output, set to NULL to get all digits
silent (logical) suppress messages
callFrom (character) allow easier tracking of messages produced

Details

For making result reproducible, a seed for generating random numbers can be set via the argument seed. However, with n=2 the resulting values are ‘fixed’ since no random component is possible at n <3.

Value

This function returns a numeric vector of random values

See Also

Normal

Examples

x1 <- (11:16)[-5]
mean(x1); sd(x1)
## the standard way
ra1 <- rnorm(n=length(x1), mean=mean(x1), sd=sd(x1))
## typically the random values deviate (slightly) from expected mean and sd
mean(ra1) - mean(x1)
sd(ra1) - sd(x1)
## random numbers with close fit to expected mean and sd :
ra2 <- rnormW(length(x1), mean(x1), sd(x1))
mean(ra2) - mean(x1)
sd(ra2) - sd(x1) # much closer to expected value

Description

This function returns CV for values in each row (using speed optimized standard deviation). Note : NaN values get replaced by NA.

Usage

rowCVs(dat, autoconvert = NULL)
Arguments

dat (numeric) matrix
autoconvert (NULL or character) allows converting simple vectors in matrix of 1 row (autoconvert="row")

Value
This function returns a (numeric) vector with CVs for each row of 'dat'

See Also

colSums, rowGrpCV, rowSds

Examples

```r
set.seed(2016); dat1 <- matrix(c(runif(200) + rep(1:10, 20)), ncol=10)
head(rowCVs(dat1))
```

---

### rowGrpCV

**Row group CV**

Description
This function calculates CVs for matrix with multiple groups of data, ie one CV for each group of data. Groups are specified as columns of 'x' in 'grp' (so length of grp should match number of columns of 'x', NAs are allowed)

Usage

`rowGrpCV(x, grp, means = NULL, listOutp = FALSE)`

Arguments

- `x` numeric matrix where replicates are organized into separate columns
- `grp` (factor) defining which columns should be grouped (considered as replicates)
- `means` (numeric) alternative values instead of means by .rowGrpMeans()
- `listOutp` (logical) if TRUE, provide output as list with $CV$, $mean$ and $n$

Value
This function returns a matrix of CV values

See Also

`rowCVs, arrayCV, replPlateCV`
Examples

```r
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
head(rowGrpCV(dat1, gr=gl(4,3,labels=LETTERS[1:4])[2:11]))
```

**Description**

`rowGrpMeans` calculates column-means for matrix with multiple groups of data, i.e. similar to `rowMeans` but one mean for each group of data. Groups are specified as columns of 'x' in 'grp' (so length of 'grp' should match number of columns of 'x', NAs are allowed).

**Usage**

```r
rowGrpMeans(x, grp, na.rm = TRUE)
```

**Arguments**

- `x`: matrix or data.frame
- `grp`: (character or factor) defining which columns should be grouped (considered as replicates)
- `na.rm`: (logical) a logical value indicating whether NA-values should be stripped before the computation proceeds.

**Value**

matrix with mean values

**See Also**

`rowSds`, `colSums`

**Examples**

```r
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)), ncol=10)
head(rowGrpMeans(dat1, gr=gl(4, 3, labels=LETTERS[1:4])[2:11]))
```
Description

This function allows easy counting the number of NAs per row in data organized in multiple subgroups as columns.

Usage

rowGrpNA(mat, grp)

Arguments

- **mat**: (matrix of data.frame) data to count the number of NAs
- **grp**: (character or factor) defining which columns should be grouped (considered as replicates)

Value

matrix with number of NAs per group

See Also

- rowGrpMeans, rowSds, colSums

Examples

```r
mat2 <- c(22.2, 22.5, 22.2, 22.2, 21.5, 22.1, 21.7, 21.5, 22, 22.2, 22.7,
         NA, NA, NA, NA, NA, NA, NA, NA, NA, NA, NA, NA,
         NA, 22.6, 23.2, 23.2, 22.4, 22.8, 22.8, NA, 23.3, 23.2, NA, 23.7,
         NA, 23.0, 23.1, 23.0, 23.2, 23.2, NA, 23.3, NA, NA, 23.3, 23.8)
mat2 <- matrix(mat2, ncol=12, byrow=TRUE)
gr4 <- gl(3, 4, labels=LETTERS[1:3])
# overall number of NAs per row
rowSums(is.na(mat2))
# number of NAs per row and group
rowGrpNA(mat2, gr4)
```
rowGrpSds

**Description**

rowGrpSds calculate Sd (standard-deviation) for matrix with multiple groups of data, ie one sd for each group of data. Groups are specified as columns of 'x' in 'grp' (so length of grp should match number of columns of 'x', NAs are allowed).

**Usage**

rowGrpSds(x, grp)

**Arguments**

- **x**: matrix where replicates are organized into separate columns
- **grp**: (character or factor) defining which columns should be grouped (considered as replicates)

**Value**

This function returns a matrix of sd values

**See Also**

rowGrpMeans, rowCVs, rowSEMs, sd

**Examples**

```r
set.seed(2016); dat1 <- matrix(c(runif(200) +rep(1:10,20)), ncol=10)
head(rowGrpSds(dat1, gr=gl(4,3,labels=LETTERS[1:4])[2:11]))
```

---

rowGrpSums

**Description**

This function calculates row-sums for matrix with multiple groups of data, ie similar to rowSums but one summed value for each line and group of data. Groups are specified as columns of 'x' in 'grp' (so length of grp should match number of columns of 'x', NAs are allowed).

**Usage**

rowGrpSums(x, grp, na.rm = TRUE)
rowMedSds

Arguments

- **x**: matrix or data.frame
- **grp**: (character or factor) defining which columns should be grouped (considered as replicates)
- **na.rm**: (logical) a logical value indicating whether NA-values should be stripped before the computation proceeds.

Value

This function returns a matrix with row/group sum values

See Also

- rowGrpMeans, rowGrpSds, rowSds, colSums

Examples

```r
set.seed(2016); dat1 <- matrix(c(runif(200) + rep(1:10, 20)), ncol=10)
head(rowGrpMeans(dat1, gr=gl(4, 3, labels=LETTERS[1:4])[, 1:11]))
```

```
rowMedSds       sd of median for each row by bootstrap
```

Description

rowMedSds determines the standard error (sd) of the median for each row by bootstrapping each row of 'dat'. Note: requires package boot

Usage

```r
rowMedSds(dat, nBoot = 99)
```

Arguments

- **dat**: (numeric) matrix, main input
- **nBoot**: (integer) number of iterations for bootstrap

Value

This function returns a (numeric) vector with estimated standard errors

See Also

- boot

Examples

```r
set.seed(2016); dat1 <- matrix(c(runif(200) + rep(1:10, 20)), ncol=10)
rowMedSds(dat1); plot(rowSds(dat1), rowMedSds(dat1))
```
Description

This function was designed for normalizing data that is supposed to be particularly similar, like a collection of technical replicates. Thus, initially for each row an independent normalization factor is calculated and the median or mean across all factors will be finally applied to the data. This function has a special mode of operation with higher content of NA values (which may pose problems with other normalization approaches). If the NA-content is higher than the threshold set in `sparseLim`, a special procedure for sparse data will be applied (iteratively treating subsets of `nCombin` columns that will be combined in a later step).

Usage

```r
rowNormalize(
  dat,
  method = "median",
  refLines = NULL,
  refGrp = NULL,
  proportMode = TRUE,
  minQuant = NULL,
  sparseLim = 0.4,
  nCombin = 3,
  omitNonAlignable = FALSE,
  maxFact = 10,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- `dat`: matrix or data.frame of data to get normalized
- `method` (character) may be "mean","median" (plus "NULL","none"); When NULL or 'none' is chosen the input will be returned as is
- `refLines` (NULL or numeric) allows to consider only specific lines of 'dat' when determining normalization factors (all data will be normalized)
- `refGrp` (integer) Only the columns indicated will be used as reference, default all columns (integer or colnames)
- `proportMode` (logical) decide if normalization should be done by multiplicative or additive factor
- `minQuant` (numeric) optional filter to set all values below given value as NA
- `sparseLim` (integer) decide at which min content of NA values the function should go in sparse-mode
Arguments were kept similar with function `normalizeThis` as much as possible. In most cases data get normalized by proportional factors. In case of log2-data (very common in omics-data) normalizing by an additive factor is equivalent to a proportional factor.

This function has a special mode of operation for sparse data (ie containing a high content of NA values). 0-values by themselves will be primarily considered as true measurement outcomes and not as missing. However, by using the argument `minQuant` all values below a given threshold will be set as NA and this may possibly trigger the sparse mode of normalizing.

Note: Using a small value of `nCombin` will give the highest chances of finding sufficient complete combination of columns with sparse data. However, this will also increase (very much) the computational efforts and time required to produce an output.

When using default proportional mode a potential division by 0 could occur, when the initial normalization factor turns out as 0. In this case a small value (default the maximum value of `dat` / 10 will be added to all data before normalizing. If this also creates 0-vales in the data this factor will be multiplied by 0.03.

Value

This function returns a matrix of normalized data

See Also

`exponNormalize`, `adjBy2ptReg`, `justvsn`

Examples

```r
# sparse matrix normalization
set.seed(2); AA <- matrix(rbinom(110,10,0.05), nrow=10)
AA[2,c(2,6,7)] <- 1; AA[3,8] <- 1;

(AA1 <- rowNormalize(AA))
(AA2 <- rowNormalize(AA, minQuant=1))  # set all 0 as NAs
(AA3 <- rowNormalize(AA, refLines=1:6, omitNonAlignable=FALSE, minQuant=1))
```
Description
This function is speed optimized sd per line (takes matrix or data.frame and treats each line as set of data for sd, )equiv to using apply. NAs are ignored from data unless entire line NA). Speed improvements may be seen at more than 100 lines. Note: NaN instances will be transformed to NA

Usage
rowSds(dat)

Arguments
dat matrix (or data.frame) with numeric values (may contain NAs)

Value
numeric vector of sd values

See Also
sd

Examples
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
rowSds(dat1)

Description
This function speed optimized SEM (standard error of the mean) for each row. The function takes a matrix or data.frame and treats each row as set of data for SEM; NAs are ignored from data. Note: NaN instances will be transformed to NA

Usage
rowSEMs(dat)

Arguments
dat matrix or data.frame
Value

This function returns a numeric vector with SEM values

See Also

rowSds, colSds, colSums

Examples

```r
set.seed(2016); dat1 <- matrix(c(runif(200)+rep(1:10,20)),ncol=10)
head(rowSEMs(dat1))
```

Description

When multiple series of data are tested simultaneously (eg using moderTestXgrp), multiple pairwise comparisons get performed. This function helps locating the samples, i.e. mean-columns, corresponding to a specific pairwise comparison.

Usage

```r
sampNoDeMArrayLM(MArrayObj, useComp, groupSep = "-", lstMeans = "means", lstP = c("BH", "FDR", "p.value"), silent = FALSE, debug = FALSE, callFrom = NULL)
```

Arguments

- `MArrayObj` (list or MArray-object) main input
- `useComp` (character or integer) index or name of pairwise-comparison to be addressed
- `groupSep` (character, length=1) separator for pair of names
- `lstMeans` (character, length=1) the list element containing the individual sample names, typically the matrix containing the replicate-mean values for each type of sample, the column-names get used
- `lstP` (character, length=1) the list element containing all pairwise comparisons performed, the column-names get used
- `silent` (logical) suppress messages
- `debug` (logical) additional messages for debugging
- `callFrom` (character) allow easier tracking of messages produced
Details
As main input one gives a list or MArrayLM-object containing testing results contain the pairwise comparisons and a specific comparison indicated by useComp to get located in the element of mean-columns (lstMeans) among all pairwise comparisons.

Value
This function returns a numeric vector (length=2) with index indicating the columns of (replicate) mean-values corresponding to the comparison specified in useComp

See Also
moderTestXgrp, this function gets used eg in MAplotW or VolcanoPlotW

Examples
```r
grp <- factor(rep(LETTERS[c(3,1,4)],c(2,3,3)))
set.seed(2017); t8 <- matrix(round(rnorm(208*8,10,0.4),2), ncol=8,
dimnames=list(paste(letters[],rep(1:8,each=26),sepal=""), paste(grp,c(1:2,1:3,1:3),sepal="")))
test8 <- moderTestXgrp(t8, grp)
head(test8$p.value) # all pairwise comparisons available
if(requireNamespace("limma", quietly=TRUE)) { # need limma installed...
sampNoDeMArrayLM(test8,1)
head(test8$means[,sampNoDeMArrayLM(test8,1)])
head(test8$means[,sampNoDeMArrayLM(test8,"C-D")] )}
```

scaleXY

Scale data to given minimum and maximum

Description
This is a convenient way to scale data to given minimum and maximum without full standarization, i.e., without dividing by the sd.

Usage
```r
scaleXY(x, min = 0, max = 1)
```

Arguments
- `x`: (numeric) vector to rescale
- `min`: (numeric) minimum value in output
- `max`: (numeric) maximum value in output

Value
vector of rescaled data (in dimensions as input)
searchDataPairs

See Also

scale

Examples

dat <- matrix(2*round(runif(100),2), ncol=4)
range(dat)
dat1 <- scaleXY(dat, 1,100)
range(dat1)
summary(dat1)

## scale for each column individually
dat2 <- apply(dat, 2, scaleXY, 1, 100)
range(dat2)
summary(dat2)

searchDataPairs

Search duplicated data over multiple columns, ie pairs of data

Description

searchDataPairs searches matrix for columns of similar data, ie 'duplicate' values in separate columns or very similar columns if realDupsOnly=FALSE. Initial distance measures will be normalized either to diagonale (normRange=TRUE) or to the real max distance observed (equal or less than diagonale). Return data.frame with names for sample-pair, percent of identical values (100 for complete identical pair) and relative (Euclidean) distance (ie max dist observed =1.0). Note, that low distance values do not necessarily imply correlating data.

Usage

searchDataPairs(
  dat,
  disThr = 0.01,
  byColumn = TRUE,
  normRange = TRUE,
  altNa = NULL,
  realDupsOnly = TRUE,
  silent = FALSE,
  callFrom = NULL
)

Arguments

dat matrix or data.frame (main input)
disThr (numeric) threshold to decide when to report similar data (applied on normalized distances, low val fewer reported), applied on normalized distances (norm to diagonale of all data for best relative 'unbiased' view)
searchLinesAtGivenSlope

byColumn  (logical) rotates main input by 90 degrees (using t), thus allows to read by rows instead of by columns
normRange  (logical) normalize each columns separately if TRUE
altNa  (character, default NULL) vector with alternative names (for display)
realDupsOnly  (logical) if TRUE will consider equal values only, otherwise will also consider very close values (based on argument disThr)
silent  (logical) suppres messages
callFrom  (character) allows easier tracking of messages produced

Value

This function returns a data.frame with names of sample-pairs, percent of identical values (100 for complete identical pair) and rel (Euclidean) distance (ie max dist observed =1.0)

See Also

duplicated

Examples

mat <- round(matrix(c(11:40,runif(20)+12,11:19,17,runif(20)+18,11:20), nrow=10), 1)
colnames(mat) <- 1:9
searchDataPairs(mat,disThr=0.05)

searchLinesAtGivenSlope

Search points forming lines at given slope

Description

searchLinesAtGivenSlope searches among set of points (2-dim) those forming line(s) with user-defined slope (‘coeff’), ie search optimal (slope-) offset parameter(s) for (regression) line(s) with given slope (‘coeff’). Note: larger data-sets : segment residuals to ‘coeff’ & select most homogenous

Usage

searchLinesAtGivenSlope(
  dat,
  coeff = 1.5,
  filtExtr = c(0, 1),
  minMaxDistThr = NULL,
  lmCompare = TRUE,
  indexPoints = TRUE,
  displHist = FALSE,
  displScat = FALSE,
  bestCluByDistRat = TRUE,
searchLinesAtGivenSlope

neighbDiLim = NULL,
silent = FALSE,
driv = FALSE,
callFrom = NULL
)

Arguments

dat matrix or data.frame, main input
coeff (numeric) slope to consider
filtExtr (integer) lower & upper quantile values, remove points with extreme deviation
to offset=0, (if single value: everything up to or after will be used)
minMaxDistThr (logical) optional minimum and maximum distance threshold
lmCompare (logical) add’l fitting of linear regression to best results, return offset AND slope
based on lm fit
indexPoints (logical) return results as list with element ‘index’ specifying retained points
displHist (logical) display histogram of residues
displScat (logical) display (simple) scatter plot
bestCluByDistRat (logical) initial selection of decent clusters based on ratio overallDist/averNeighbDist
(or by CV & cor)
neighbDiLim (numeric) additional threshold for (trimmed mean) neighbour-distance
silent (logical) suppress messages
driv (logical) for bug-tracking: more/enhanced messages
callFrom (character) allow easier tracking of messages produced

Value

This functions returns a matrix of line-characteristics (or if indexPoints is TRUE then list (linecharacteristics & index & lm-results)

See Also

lm

Examples

set.seed(2016); ral <- runif(300)
dat1 <- cbind(x=round(c(1:100+r*ral[1:100]/5,4*r*ral[1:50]),1),
y=round(c(1:100+r*ral[101:200]/5,4*r*ral[101:150]), 1))
(l11 <- searchLinesAtGivenSlope(dat1, coeff=1))
**simpleFragFig**

*Simple figure showing line from start- to end-sites of edges (or fragments) defined by their start- and end-sites. simpleFragFig draws figure showing start- and end-sites of edges (or fragments)*

**Description**

Simple figure showing line from start- to end-sites of edges (or fragments) defined by their start- and end-sites.

`simpleFragFig` draws figure showing start- and end-sites of edges (or fragments).

**Usage**

```r
simpleFragFig(
  frag,
  fullSize = NULL,
  sortByHead = TRUE,
  useTit = NULL,
  useCol = NULL,
  displNa = TRUE,
  useCex = 0.7
)
```

**Arguments**

- `frag` (matrix) 2 columns defining begin- and end-sites (as integer values)
- `fullSize` (integer) optional max size used for figure (x-axis)
- `sortByHead` (logical) sort by begin-sites (if `TRUE`) or sort by end-sites
- `useTit` (character) custom title
- `useCol` (character) specify colors, if numeric vector will be considered as score values
- `displNa` (character) display names of edges (figure may get crowded)
- `useCex` (numeric) expansion factor, see also `par`

**Value**

matrix with mean values

**See Also**

`buildTree, countSameStartEnd, contribToContigPerFrag`

**Examples**

```r
frag2 <- cbind(beg=c(2,3,7,13,15,7,9,7,3,7,5,7,3),end=c(6,12,8,18,20,20,19,12,12,4,12,7,12,4))
rownames(frag2) <- c("A","E","B","C","D","F","H","G","I","J","K","L","M","N")
simpleFragFig(frag2,fullSize=21,sortByHead=TRUE)
buildTree(frag2)
```
singleLineAnova

2-factorial Anova on single line of data

Description

This function runs 2-factorial Anova on a single line of data (using aov from package stats) using a model with two factors (without factor-interaction) and extracts the corresponding p-value.

Usage

singleLineAnova(
  dat,
  fac1,
  fac2,
  inclInteraction = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dat numeric vector

fac1 (character or factor) vector describing grouping elements of dat for first factor, must be of same length as fac2

fac2 (character or factor) vector describing grouping elements of dat for second factor, must be of same length as fac1

inclInteraction (logical) decide if factor-interactions (eg synergy) should be included to model

silent (logical) suppress messages

debug (logical) additional messages for debugging

callFrom (character) allow easier tracking of messages produced

Value

This function returns the (uncorrected) p for factor 'Pr(>F)' (see aov)

See Also

aov, anova; for repeated tests using the package limma including lmFit and eBayes see test2factLimma

Examples

set.seed(2012); dat <- round(runif(8),1)
singleLineAnova(dat, gl(2,4),rep(1:2,4))
sortBy2CategorAnd1IntCol

Sort matrix by two categorical and one integer columns

Description

This function sorts matrix 'mat' subsequently by categorical and numerical columns of 'mat', i.e., lines with identical values for categor are sorted by numeric value.

Usage

sortBy2CategorAnd1IntCol(
  mat,  
categCol, 
  numCol,  
  findNeighb = TRUE, 
  decreasing = FALSE, 
  silent = FALSE, 
  debug = FALSE, 
  callFrom = NULL
)

Arguments

mat matrix (or data.frame) from which by 2 columns will be selected for sorting
categCol (integer or character) which columns of 'mat' to be used as categorical columns
numCol (integer or character) which column of 'mat' to be used as integer columns
findNeighb (logical) if 'findNeighb' neighbour cols according to 'numCol' will be identified as groups & marked in new col 'neiGr', orphans marked as NA
decreasing (logical) order of sort
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This function returns a sorted matrix (same dimensions as 'mat')

Examples

mat <- cbind(aa=letters[c(3,rep(7:8,3:4),4,4:6,7)],bb=LETTERS[rep(1:5,c(1,3,4,4,1))],
  nu=c(23:21,23,21,22,18:12))
mat[cbind(c(3:5,1:2,6:9,13:10),c(3:5,1:2,6:9,13:10))]
sortBy2CategorAnd1IntCol(mat,cate=c("bb","aa"),num="nu",findN=FALSE,decr=TRUE)
sortBy2CategorAnd1IntCol(mat,cate=c("bb","aa"),num="nu",findN=TRUE,decr=FALSE)
sortByNRepeated

Make a list of common occurrences sorted by number of repeats

Description

The aim of this function is to count the number of occurrences of words when comparing separate vectors \((x, y \text{ and } z)\) or from a list (given as \(x\)) and to give an output sorted by their frequency. The output lists the various values/words by their frequency, the names of the resulting list-elements indicate number of times the values/words were found repeated.

Usage

```
sortByNRepeated(
  x,
  y = NULL,  
  z = NULL,
  filterIntraRep = TRUE,
  silent = TRUE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- **x**: (list, character or integer) main input, if list, arguments \(y\) and \(z\) will not be used
- **y**: (character or integer) supplemental vector to compare with \(x\)
- **z**: (character or integer) supplemental vector to compare with \(x\)
- **filterIntraRep**: (logical) allow making vectors \(x\), \(y\) and \(z\) unique before comparing (defaults to TRUE)
- **silent**: (logical) suppress messages
- **debug**: (logical) additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

Details

In order to compare the frequency of values/words between separate vectors or vectors within a list, it is necessary that these have been made unique before calling this function or using `filterIntraRep=TRUE`. In case the input is given as list (in \(x\)), there is no restriction to the number of vectors to be compared. With very long lists, however, the computational effort increases (like it does when using `table`).

Value

This function returns a list sorted by number of occurrences. The names of the list indicate the number of repeats.
stableMode

See Also

table, replicateStructure

Examples

sortByNRepeated(x=LETTERS[1:11], y=LETTERS[3:13], z=LETTERS[6:12])
sortByNRepeated(x=LETTERS[1:11], y=LETTERS[c(3:13,5:4)], z=LETTERS[6:12])

stableMode Estimate mode (most frequent value)

Description

Estimate mode, ie most frequent value. In case of continuous numeric data, the most frequent values may not be the most frequently repeated exact term. This function offers various approaches to estimate the mode of a numeric vector. Besides, it can also be used to identify the most frequent exact term (in this case also from character vectors).

Usage

stableMode(
  x,
  method = "density",
  finiteOnly = TRUE,
  bandw = NULL,
  rangeSign = 1:6,
  silent = FALSE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

x (numeric, or character if 'method='mode') data to find/estimate most frequent value

method (character) There are 3 options: BBmisc, binning and density (default). If "binning" the function will search context dependent, ie like most frequent class of histogram. Using "binning" mode the search will be refined if either 80 percent of values in single class or >50 percent in single class.

finiteOnly (logical) suppress non-finite values; allows avoiding NULL as result in presence of some Inf values; NA will be ignored in any case

bandw (integer) only used when method="binning" or method="density" : defines the number of points to look for density or number of classes used; very "critical" parameter, may change results in strong way. Note: with method="binning": At higher values for "bandw" you will finally loose advantage of histLike-type search of mode !
rangeSign: (integer) only used when method="binning": range of numbers used as number of significant values
silent: (logical) suppress messages
callFrom: (character) allows easier tracking of messages produced
debug: (logical) additional messages for debugging

Details
The argument method allows to choose among (so far) 4 different methods available. If "density" is chosen, the most dense region of sqrt(n) values will be chosen; if "binning", the data will be binned (like in histograms) via rounding to a user-defined number of significant values ("rangeSign"). If method is set to "BBmisc", the function computeMode() from package BBmisc will be used. If "mode" is chosen, the first most frequently occurring (exact) value will be returned, if "allModes", all ties will be returned. This last mode also works with character input.

Value
This function returns a numeric vector with value of mode, the name of the value indicates it’s position

See Also
computeMode() in package BBmisc

Examples
```r
set.seed(2012); dat <- round(c(rnorm(50), runif(100)),3)
stableMode(dat)
```

Description
This function works similar to scale, however, it evaluates the entire input and not column-wise (and independently as scale does). With Standarizing we speak of transforming the data to end up with mean=0 and sd=1. Furthermore, in case of 3-dim arrays, this function returns also an object with the same dimensions as the input.

Usage
```r
standardW(
  mat,
  byColumn = FALSE,
  na.rm = TRUE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```
stdErrMedBoot

Arguments

mat (matrix, data.frame or array) data that need to get standardized.
byColumn (logical) if TRUE the function will be run independently over all columns such as apply(mat,2,standardW)
na.rm (logical) if NAs in the data don’t get ignored via this argument, the output will be all NA
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced

Value

This functions trunrs a vector of rescaled data (in dimensions as input)

See Also

scale

Examples

dat <- matrix(2*round(runif(100),2), ncol=4)
mean(dat); sd(dat)

dat2 <- standardW(dat)
apply(dat2, 2, sd)
summary(dat2)

dat3 <- standardW(dat, byColumn=TRUE)
apply(dat2, 2, sd)
summary(dat2)
mean(dat2); sd(dat2)

stdErrMedBoot  Standard error of median by boot-strap

Description

stdErrMedBoot estimate standard eror of median by boot-strap approach. Note: requires package boot

Usage

stdErrMedBoot(x, nBoot = 99)
summarizeCols

Arguments

- **x** (numeric) vector to estimate median and its standard error
- **nBoot** (integer) number for iterations

Value

This function returns a (numeric) vector with estimated standard error

See Also

- boot

Examples

```r
set.seed(2014); ra1 <- c(rnorm(9,2,1),runif(8,1,2))
rat1 <- ratioAllComb(ra1[1:9],ra1[10:17])
median(rat1); stdErrMedBoot(rat1)
```

summarizeCols

Summarize columns (as median, mean, min, last or other methods)

Description

summarizeCols summarizes all columns of matrix (or data.frame). In case of text-columns the sorted middle (~median) will be given, unless `maxLast`, `minLast`, `maxLast`, `maxAbsLast` or `minLast`... consider only last column of `matr` : choose from all columns the line where (max of) last col is at min; `medianComplete` or `meanComplete` considers only lines/rows where no NA occur (NA have influence other columns !)

Usage

```r
summarizeCols(
  matr,  # data.frame matrix of data to be summarized by column (may do different method for text and numeric columns)
  meth = "median",  # (character) summarization method (eg 'maxLast','minLast','maxLast','maxAbsLast','minLast', 'medianComplete' or 'meanComplete')
  silent = FALSE,  # (logical) suppress messages
  debug = FALSE,  # (logical) additional messages for debugging
  callFrom = NULL  # (character) allow easier tracking of messages produced
)
```

Arguments

- **matr**
- **meth**
- **silent**
- **debug**
- **callFrom**
Value

vector with summary for each column

See Also

rowMeans in colSums

Examples

```r
t1 <- matrix(round(runif(30,1,9)), nc=3); rownames(t1) <- letters[c(1:5,3:4,6:4)]
summarizeCols(t1, me="median")
t(sapply(by(t1,rownames(t1), function(x) x), summarizeCols,me="maxLast"))
t3 <- data.frame(ref=rep(11:15,3), tx=letters[1:15],
  matrix(round(runif(30,-3,2),1), ncol=2), stringsAsFactors=FALSE)
by(t3,t3[,1], function(x) x)
t(sapply(by(t3,t3[,1], function(x) x), summarizeCols,me="maxAbsLast"))
```

---

sumNaperGroup  

`Count number of NAs per sub-set of columns`

Description

This function will count the number of NAs per group (defined by argument grp) while summing over all lines of a matrix or data.frame. The row-position has no influence on the counting. Using the argument asRelative=TRUE the result will be given as (average) number of NAs per row and group.

Usage

```r
sumNaperGroup(
  x,
  grp,
  asRelative = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
```

Arguments

- **x**: matrix or data.frame which may contain NAs
- **grp**: factor describing which column of `dat` belongs to which group
- **asRelative**: (logical) return as count of NAs per row and group
- **silent**: (logical) suppress messages
- **debug**: (logical) additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced
Value

This function returns an integer vector with count of NAs per group

See Also

NA, filter NAs by line presenceFilt

Examples

```r
mat <- matrix(1:25, ncol=5)
mat[lower.tri(mat)] <- NA
sumNAperGroup(mat, rep(1:2,c(3,2)))
sumNAperGroup(mat, rep(1:2,c(3,2)), asRelative=TRUE)
```

---

**sysDate**

*System-date (compressed format)*

Description

This function returns current date (based on Sys.Date) in different format options.

Usage

```
sysDate(style = "univ1")
```

Arguments

- **style** (character) choose style (default 'univ1' for very compact style)

Details

Multiple options for formatting exist: 'univ1' or 'wr' ... (default) compact style using day, first 3 letters of English name of month (lowercaps) and last 2 letters of year as ddmmmyy, eg 14jun21
- 'univ2' ... as ddMmmyy, eg 14Jun21
- 'univ3' ... as ddMonthyyyy, eg 14June2021
- 'univ4' ... as ddmmyy, eg 14June2021
- 'univ5' ... as yyyy-mm-dd (output of Sys.Date()), eg 2021-06-14
- 'univ6' ... as yyyy-number of day (in year), eg 2021-165
- 'local1' ... compact sytle using day, first 3 letters of current locale name of month (not necessarily unique !) and last 2 letters of year as ddmmyy, eg 14jui21
- 'local2' ... as ddmmyy, month based on current locale (not necessarily unique !), eg 14Jui21
- 'local3' ... as ddMonthyyyy, month based on current locale , eg 14Juin2021
- 'local4' ... as ddmmyy, month based on current locale , eg 14juin2021
- 'local5' ... as dd-month-yyyy, month based on current locale , eg 14-juin-2021
- 'local6' ... as yyyymonthddd, month based on current locale , eg 2021juin14
tableToPlot

Print matrix-content as plot

Description

This function prints all columns of matrix in plotting region for easier inclusion to reports (default values are set to work for output as A4-sized pdf). It was made for integrating listings of text to graphical output to devices like png, jpeg or pdf.

Usage

```r
tableToPlot(
  matr,
  colPos = c(0.05, 0.35, 0.41, 0.56),
  useCex = 0.7,
  useAdj = c(0, 1, 1, 0),
  titOffS = 0,
  useCol = 1,
  silent = FALSE,
  callFrom = NULL
)
```

Arguments

- **matr**: (matrix) main (character) matrix to display
- **colPos**: (numeric) postion of columns on x-scale (from 0 to 1)
- **useCex**: (numeric) cex expansion factor for size of text (may be different for each column)
- **useAdj**: (numeric) left/cneter/right alignment for text (may be different for each column)
- **titOffS**: (numeric) offset for title line (relative to 'colPos')
- **useCol**: color specification for text (may be different for each column)
- **silent**: (logical) suppress messages
- **callFrom**: (character) allow easier tracking of message(s) produced

Value

character vector with formatted date

See Also

date, Sys.Date and Sys.time,
Details
This function was initially designed for listings with small/medium 1st col (eg counour or index),
2nd & 3rd col small and long 3rd col (like file paths). Obviously, the final number of lines one
can pack and still read correctly into the graphical output depends on the size of the device (on
a pdf of size A4 one can pack up to apr. 110 lines). Of course, Sweave, combined with LaTeX,
provides a powerful alternative for wrapping text to pdf-output (and further combining text and
graphics). Note: The final result on pdf devices may vary depending on screen-size (ie with of
current device), the parameters ’colPos’ and ’titOffS’ may need some refinements. Note: In view
of typical page/figure layouts like A4, the plotting region will be split to avoid too wide spacing
between rows with less than 30 rows.

Value
This function returns NULL (no R-object returned), print ’plot’ in current device only

See Also
Sweave for more flexible framework

Examples
## as example let's make a listing of file-names and associated parameters in current directory
mat <- dir()
mat <- cbind(no=1:length(mat),fileName=mat,mode=file.mode(mat),
           si=round(file.size(mat)/1024),path=getwd())
## Now, we wrap all text into a figure (which could be saved as jpg, pdf etc)
tableToPlot(mat[,1:5],colPos=c(0.01,0.4,0.46,0.6),titOffS=c(0.05,-0.03,-0.01,0.06))
tableToPlot(mat,colPos=c(0.016,0.36,0.42,0.75),useAdj=0.5,titOffS=c(-0.01,0,-0.01,0,-0.1))

Description
The aim of this function is to provide convenient acces to two-factorial (linear) testing withing
the framework of makeMAList including the empirical Bayes shrinkage. The input data 'datMatr'
which should already be organized as limma-type MAList, eg using using makeMAList. Note: This
function uses the Bioconductor package limma.

Usage
test2factLimma(
  datMatr,
  fac1,
  fac2,
  testSynerg = TRUE,
  testOrientation = "=",
  addResults = c("lfdr", "FDR", "Mval", "means"),
)
addGenes = NULL,
silent = FALSE,
callFrom = NULL,
debug = FALSE
)

Arguments

datMatr       matrix or data.frame with lines as independent series of measures (eg different
genes)
fac1          (character or factor) vector describing grouping elements of each line of 'dat-
Matr' for first factor, must be of same length as fac2
fac2          (character or factor) vector describing grouping elements of each line of 'dat-
Matr' for second factor, must be of same length as fac1
testSynerg   (logical) decide if factor-interactions (eg synergy) should be included to model
testOrientation       (character) default (or any non-recognized input) ‘=’, otherwise either ‘>’, ‘ge-
nerter’, ‘sup’, ‘upper’ or ‘<’, ‘inf’, ‘lower’
addResults       (character) vector defining which types of information should be included to
output, may be ‘lfdr’, ‘FDR’ (for BY correction), ‘Mval’ (M values), ‘means’
(add matrix with mean values for each group of replicates)
addGenes       (matrix or data.frame) additional information to add to output
silent         (logical) suppress messages
callFrom       (character) allow easier tracking of messages produced
debug          (logical) additional messages for debugging

Value

This function returns an object of class "MArrayLM" (from limma) containing/enriched by the
testing results

See Also

makeMAList, single line testing lmFit and the eBayes-family of functions in package limma

Examples

set.seed(2014)
dat0 <- rnorm(30) + rep(c(10,15,19,20),c(9,8,7,6))
fa <- factor(rep(letters[1:4],c(9,8,7,6)))
dat2 <- data.frame(facA=rep(c("-","A","-","A"), c(9,8,7,6)),
facB= rep(c("-","-","B","B"), c(9,8,7,6)),
dat1=dat0, dat2=runif(30))
grpNa <- sub("-", "", sub("\", "", apply(dat2[,1:2], 1, paste, collapse="")))
test2f <- test2factLimma(t(dat2[,3:4]), dat2$facA, dat2$facB, testS=FALSE)
test2f
# Now you can easily summarize results using topTable from limma
if(requireNamespace("limma", quietly=TRUE)) {
  library(limma)
transpGraySca

Make single vector gray-gradient

Description

This function helps making gray-gradients. Note: The resulting color gradient does not seem linear
to the human eye, you may try gray.colors instead

Usage

transpGraySca(startGray = 0.2, endGrey = 0.8, nSteps = 5, transp = 0.3)

Arguments

  startGray      (numeric) gray shade at start
  endGrey        (numeric) gray shade at end
  nSteps         (integer) number of levels
  transp         (numeric) transparency alpha

Value

  character vector (of same length as x) with color encoding

See Also

  gray.colors

Examples

layout(1:2)
col1 <- transpGraySca(0.8,0.3,7,0.9)
pie(rep(1,length(col1)), col=col1, main="from transpGraySca")
col2 <- gray.colors(7,0.9,0.3,alph=0.9)
pie(rep(1,length(col2)), col=col2, main="from gray.colors")
treatTxtDuplicates  
*Locate duplicates in text and make non-redundant*

**Description**

treatTxtDuplicates locates duplicates in character-vector 'x' and return list (length=3): with $init$ (initial), $nRed$ .. non-redundant text by adding number at end or beginning, and $nrLst$ .. list-version with indexes per unique entry. Note: NAs (if multiple) will be renamed to NA_1, NA_2

**Usage**

treatTxtDuplicates(
  x,
  atEnd = TRUE,
  sep = "_",
  onlyCorrectToUnique = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

**Arguments**

- **x**  
  (character) vector with character-entries to identify (and remove) duplicates

- **atEnd**  
  (logical) decide location of placing the counter (at end or at beginning of ID)  
  (see `correctToUnique`)

- **sep**  
  (character) separator to add before counter when making non-redundant version

- **onlyCorrectToUnique**  
  (logical) if TRUE, return only vector of non-redundant

- **silent**  
  (logical) suppress messages

- **debug**  
  (logical) additional messages for debugging

- **callFrom**  
  (character) allow easier tracking of messages produced

**Value**

list with $init$, $nRed$, $nrLst$

**See Also**

For simple correction use `correctToUnique`

**Examples**

treatTxtDuplicates(c("li0",NA,rep(c("li2","li3"),2)))
correctToUnique(c("li0",NA,rep(c("li2","li3"),2)))
triCoord

Pairwise x,y combinations

Description

triCoord gets pairwise combinations for 'n' elements; returns matrix with x & y coordinates to form all pairwise groups for 1:n elements

Usage

triCoord(n, side = "upper")

Arguments

n (integer) number of elements for making all pair-wise combinations
side (character) "upper" or "lower"

Value

2-column matrix with indexes for all pairwise combinations of 1:n

See Also

lower.tri or upper.tri, simpler version upperMaCoord

Examples

triCoord(4)

trimRedundText

Trim redundant text

Description

This function allows trimming/removing redundant text-fragments (redundant from head or tail) out of character vector 'txt'.

Usage

trimRedundText(
  txt,
  minNchar = 1,
  side = "both",
  spaceElim = FALSE,
  silent = TRUE,
  callFrom = NULL,
  debug = FALSE
)

Arguments

txt  character vector to be treated
minNchar  (integer) minimum number of characters that must remain
side  (character) may be be either 'both', 'left' or 'right'
spaceElim  (logical) optional removal of any heading or tailing white space
silent  (logical) suppress messages
callFrom  (character) allow easier tracking of messages produced
debug  (logical) display additional messages for debugging

Value

This function returns a modified character vector

See Also

Inverse : Find/keep common text keepCommonText; you may also look for related functions in package stringr

Examples

txt1 <- c("abcd_ccc","bcd_ccc","cde_ccc")
trimRedundText(txt1, side="right")  # trim from right

txt2 <- c("ddd_ab","ddd_bcd","ddd_cde")
trimRedundText(txt2, side="left")  # trim from left

tTestAllVal

t.test on all individual values against all other values

Description

Run t.test on each indiv value of x against all its neighbours (=remaining values of same vector) in order to test if tis value is likely to belong to vector x. This represents a repeated leave-one-out testing. Multiple choices for multiple testing correction are available.

Usage

tTestAllVal(
  x,
  alph = 0.05,
  alternative = "two.sided",
  p.adj = NULL,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)
Arguments

- **x**: matrix or data.frame
- **alpha**: (numeric) threshold alpha (passed to `t.test`)
- **alternative**: (character) will be passed to `t.test` as argument 'alternative', may be "two.sided"...
- **p.adj**: (character) multiple test correction: may be NULL (no correction), "BH", "BY", "holm", "hochberg" or "bonferroni" (but not 'fdr' since this may be confounded with local false discovery rate), see `p.adjust`
- **silent**: (logical) suppress messages
- **debug**: (logical) additional messages for debugging
- **callFrom**: (character) allow easier tracking of messages produced

Value

This function returns a numeric vector with p-values or FDR (depending on argument p.adj)

See Also

- `t.test`, `p.adjust`

Examples

```r
set.seed(2016); x1 <- rnorm(100)
allTests1 <- tTestAllVal(x1)
hist(allTests1,breaks="FD")
```

Description

The aim of this function is to provide help in automatically harmonizing enumerators at the end of sample-names. When data have same grouped setup/design, many times this is reflected in their names, eg 'A_sample1', 'A_sample2' and 'B_sample1'. However, human operators may use multiple similar (but not identical) ways of expressing the same meaning, eg writing 'A_Samp_1'. This function allows testing a panel of different extensions of enumerators and (if recognized) to replace them by a user-defined standard text/enumerator. Please note that the more recent function `rmEnumeratorName` offers better/more flexible options.

Usage

```r
unifyEnumerator(
  x, 
  refSep = "_", 
  baseSep = c("\-", "\ ", "\."), 
  supplEnu = c("Repl", "Rep", "R", "Number", "No", "Sample", "Samp"),
)```
unifyEnumerator

```r
stringentMatch = TRUE,
silent = FALSE,
debug = FALSE,
callFrom = NULL
```

Arguments

- `x` (character): main input
- `refSep` (character): separator for output
- `baseSep` (character): basic separators to test (you have to protect special characters)
- `suplEnu` (character): additional text
- `stringentMatch` (logical): decide if enumerator text has to be found in all instances or only once
- `silent` (logical): suppress messages
- `debug` (logical): display additional messages for debugging
- `callFrom` (character): allow easier tracking of messages produced

Details

This function has been developed for matching series of the same samples passing in parallel through different evaluation software (see R package wrProteo). The way human operators may name things may easily leave room for surprises and this function allows testing only a limited number of common ways of writing. Thus, in any case, the user is advised to inspect the results by eye and - if needed- to adjust the parameters.

Basically enumerator separators can be constructed by combing a base-separator `baseSep` (like '-', '_', etc) and an enumerator-abbreviation `suplEnu`. Then, all possible combinations will be tested if they occur in the text `x`. Furthermore, the text searched has to be followed by one or multiple digits at the end of text entry (decimal comma-separators etc are not allowed). Thus, if there is other 'free text' following to the right after the enumerator-text this function will not find any enumerators to replace.

The argument `stringentMatch` allows defining if this text has to be found in all text-entries of `x` or just one of them. When using `stringentMatch=FALSE` there is risk that other text not meant to design enumerators may be picked up and modified.

Please note, that with large data-sets (ie many columns) testing/checking a larger panel of enumerator-abbreviations may result in slower performance. In cases of larger data-sets it may be more effective to first study the data and then run simple substitutions using sub targeted for this very case.

Value

This function returns a character vector of same length as input `x`, with it's content as adjusted enumerators

See Also

- `rmEnumeratorName` for better/more flexible options; `grep` or `sub()`, etc if exact and consistent patterns are known
uniqCountReport

Examples

unifyEnumerator(c("ab-1","ab-2","c-3"))
unifyEnumerator(c("ab-R1","ab-R2","c-R3"))
unifyEnumerator(c("ab-1","c3-2","dR3"), strin=FALSE);

uniqCountReport Report number of unique and redundant elements (optional figure)

Description

Make report about number of unique and redundant elements of vector 'dat'. Note: fairly slow for long vectors !!

Usage

uniqCountReport(
  dat,
  frL = NULL,
  plotDispl = FALSE,
  tit = NULL,
  col = NULL,
  radius = 0.9,
  sizeTo = NULL,
  clockwise = FALSE,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

dat (character or numeric vector) main input where number of unique (and redundant) should be determined
frL (logical) optional (re-)introducing results from duplicated to shorten time of execution
plotDispl (logical) decide if pie-type plot should be produced
tit (character) optional title in plot
col (character) custom colors in pie
radius (numeric) radius passed to pie
sizeTo (numeric or character) optional reference group for size-population relative adjusting overall surface of pie
clockwise (logical) argument passed to pie
silent (logical) suppress messages
debug (logical) additional messages for debugging
callFrom (character) allow easier tracking of messages produced
upperMaCoord

Value

vector with counts of n (total), nUnique (wo any repeated), nHasRepeated (first of repeated), nRedundant (first of repeated), optional figure

See Also

correctToUnique, unique

Examples

layout(1:2)
uniqCountReport(rep(1:7,1:7),plot=TRUE)
uniqCountReport(rep(1:3,1:3),plot=TRUE,sizeTo=rep(1:7,1:7))

upperMaCoord(n)

Arguments

n (integer) number of elements for making all pair-wise combinations

Value

2-column matrix with indexes for all pairwise combinations of 1:n

See Also

lower.tri, more evolved version triCoord

Examples

upperMaCoord(4)
withinRefRange  

Check for values within range of reference

Description
withinRefRange checks which values of numeric vector 'x' are within range +/- 'fa' x 'ref' (ie within range of reference).

Usage
withinRefRange(x, fa, ref = NULL, absRef = TRUE, asInd = FALSE)

Arguments
x matrix or data.frame
fa (numeric) absolute or relative tolerance value (numeric, length=1), interpreted according to 'absRef' as absolute or relative to 'x'(ie fa*ref)
ref (numeric) (center) reference value for comparison (numeric, length=1), if not given mean of 'x' (excluding NA or non-finite values) will be used
absRef (logical) return result as absolute or relative to 'x'(ie fa*ref)
asInd (logical) if TRUE return index of which values of 'x' are within range, otherwise return values if 'x' within range

Value
numeric vector (containing only the values within range of reference)

Examples
## within 2.5 +/- 0.7
withinRefRange(-5:6,fa=0.7,ref=2.5)
## within 2.5 +/- (0.7*2.5)
withinRefRange(-5:6,fa=0.7,ref=2.5,absRef=FALSE)

writeCsv  

Write (and convert) csv files

Description
This functions is absed on write.csv allows for more options when writing data into csv-files. The main input may be given as R-object or read from file 'input'. Then, one can (re-)write using specified conversions. An optional filter to select columns (column-name specified via 'filterCol') is available. The output may be simultaneously written to multiple formats, as specified in 'expTy', tabulation characters may be converted to avoid accidentally split/shift text to multiple columns. Note: Mixing '.' and ',' as comma separators via text-columns or fused text&data may cause problems lateron, though.
Usage

writeCsv(
  input,
  inPutFi = NULL,
  expTy = c("Eur","US"),
  imporTy = "Eur",
  filename = NULL,
  quote = FALSE,
  filterCol = NULL,
  replMatr = NULL,
  returnOut = FALSE,
  SYLKprevent = TRUE,
  digits = 22,
  silent = FALSE,
  debug = FALSE,
  callFrom = NULL
)

Arguments

input either matrix or data.frame
inPutFi (character or NULL) file-name to be read (format as US or Euro-type may specified via argument importTy)
expTy (character) 'US' and/or 'Eur' for sparator and decimal type in output
imporTy (character) default 'Eur' (otherwise set to 'US')
filename (character) optional new file name(s)
quote (logical) will be passed to function write.csv
filterCol (integer or character) optionally, to export only the columns specified here
replMatr optional, matrix (1st line:search, 2nd li:use for replacing) indicating which characters need to be replaced
returnOut (logical) return output as object
SYLKprevent (logical) prevent difficulty when opening file via Excel. In some cases Excel presumes (by error) the SYLK format and produces an error when trying to open files : To prevent this, if necessary, the 1st column-name will be changed from 'ID' to 'Id'.
digits (integer) limit number of signif digits in output (ie file)
silent (logical) suppress messages
debug (logical) for bug-tracking: more/enhanced messages
callFrom (character) allow easier tracking of messages produced

Value

This function writes a file to disk and returns NULL unless returnOut=TRUE
XYToDiffPpm

Express difference as ppm

Description
This function transforms offset (pairwise-difference) between 'x' & 'y' to ppm (as normalized difference ppm, parts per million, i.e. (x-y)/y). This type of expression for differences is used e.g. in mass-spectrometry.

Usage
XYToDiffPpm(x, y, nSign = NULL, silent = FALSE, debug = FALSE, callFrom = NULL)

Arguments
- **x** (numeric) typically for measured variable
- **y** (numeric) typically for theoretical/expected value (vector must be of same length as 'x')
- **nSign** (integer) number of significant digits in output
- **silent** (logical) suppress messages
- **debug** (logical) additional messages for debugging
- **callFrom** (character) allow easier tracking of messages produced

Value
This function returns a numeric vector of (ratio-) ppm values

See Also
ratToPpm for classical ppm

Examples
dat1 <- data.frame(ini=letters[1:5], x1=1:5, x2=11:15, t1=c("10,10","20.20","11,11","21,21","33.33"),
t2=c("10,11","20.21","kl;kl","az;az","ze;ze"))
fiNa <- file.path(tempdir(), paste("test",1:2,".csv",sep=""))
writeCsv(dat1, filename=fiNa[1])
dir(path=tempdir(), pattern="cs")

(writeCsv(dat1, replM=rbind(bad=c(";",""), replBy="__"), expTy=c("Eur"),
          returnOut=TRUE, filename=fiNa[2])))

See Also
write.csv in write.table, batch reading using this package readCsvBatch
Examples

```r
set.seed(2017); aa <- runif(10,50,900)
cbind(x=aa,y=aa+1e-3,ppm=XYToDiffPpm(aa,aa+1e-3,nSign=4))
```
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