

Package ‘caroline’

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Title A Collection of Database, Data Structure, Visualization, and
Utility Functions for R

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Depends R (>= 1.8.0), methods

Suggests MASS, RSQLite, grid

Enhances sm, DBI, gplots, limma, edgeR

Description The caroline R library contains dozens of functions useful for: database migration (dbWriteTable2), database style joins & aggregation (nerge, groupBy & bestBy), data structure conversion (nv, tab2df), legend table making (sstable & leghead), plot annotation (labsegs & mvlabs), data visualization (violins, pies & raPlot), character string manipulation (m & pad), file I/O (write.delim), batch scripting and more. The package's greatest contributions lie in the database style merge, aggregation and interface functions as well as in its extensive use and propagation of row, column and vector names in most functions.

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addFactLevs	<i>Add new levels to the Factors in a DataFrame.</i>
-------------	--

Description

This function loops through all of the factor columns in a dataframe and adds new.levels to the factor levels list.

Usage

```
addFactLevs(x, new.levels=NA)
```

Arguments

x a dataframe.
new.levels new levels to be added.

See Also

[factor](#), [levels](#)

bestBy *Find the "best" record within subgroups of a dataframe.*

Description

Finding the an extreme record for each group within a dataset is a more challenging routine task in R and SQL. This function provides a easy interface to that functionality either using R (fast for small data frames) or SQL (fastest for large data)

Usage

```
bestBy(df, by, best, c1mns=names(df), inverse=FALSE, sql=FALSE)
```

Arguments

df a data frame.
by the factor (or name of a factor in df) used to determine the grouping.
c1mns the colums to include in the output.
best the column to sort on (both globally and for each sub/group)
inverse the sorting order of the sort column as specified by 'best'
sql whether or not to use SQLite to perform the operation.

Value

A data frame of 'best' records from each factor level

Author(s)

David Schruth

See Also

[groupBy](#)

Examples

```
blast.results <- data.frame(score=c(1,2,34,4,5,3,23),
                           query=c('z','x','y','z','x','y','z'),
                           target=c('a','b','c','d','e','f','g')
                           )
best.hits.R <- bestBy(blast.results, by='query', best='score', inverse=TRUE)
best.hits.R
## or using SQLite
best.hits.sql <- bestBy(blast.results, by='query', best='score', inverse=TRUE, sql=TRUE)
best.hits.sql
```

 dbWriteTable2

Data Import Wrapper for dbWriteTable.

Description

This is wrapper for dbWriteTable written with the the primary improvements focusing on database import into an existing table definition schema. The function matches and rearranges columns of the dataframe to database feilds and additionally performs checks for NA's in required variables, overlength strings, and type mismatches. There also exists support for updating of the PostgreSQL specific sequence for tables with auto incrementing primary keys.

Usage

```
dbWriteTable2(con, table.name, df, fill.null = TRUE, add.id=TRUE,
              row.names=FALSE, pg.update.seq=FALSE, ...)
```

Arguments

con	connection.
table.name	The name of the table to which the data frame is to be loaded.
df	A dataframe to be loaded to the database.
fill.null	Should new db present fields be added to the data.frame before it is loaded?.
add.id	Should a new column should be added for the database id?
row.names	Should the row names be loaded as a seperate column? (unlike the original dbWriteTable, default is FALSE)
pg.update.seq	should the table primary key's sequence be updated to the highest id value +1? (Postgres specific)
...	other parameters passed to dbWriteTable.

Value

If successful, the ids of the newly added database records (invisible)

See Also[dbWriteTable](#)

geomean	<i>Calculate the Geometric Mean</i>
---------	-------------------------------------

Description

A trivial one-line function for `exp(mean(log()))`

Usage

```
geomean(x)
```

Arguments

x a vector of numeric values

Value

the geometric mean (a scalar value)

See Also

`geometric.mean`

Examples

```
geomean(rnorm(20,5))
```

groupBy	<i>Group a dataframe by a factor and perform aggregate functions.</i>
---------	---

Description

The R equivalent of a SQL 'group by' call.

Usage

```
groupBy(df, by, aggregation, cmlns=names(df), collapse=',',  
        distinct=FALSE, sql=FALSE, full.names=FALSE, ...)
```

Arguments

df	a data frame.
by	the factor (or name of a factor in df) used to determine the grouping.
aggregation	the functions to perform on the output (default is to sum). Suggested functions are: 'sum', 'mean', 'var', 'sd', 'max', 'min', 'length', 'paste', NULL.
clmns	the columns to include in the output.
collapse	string delimiter for columns aggregated via 'paste' concatenation.
distinct	used in conjunction with paste and collapse to only return unique elements in a delimited concatenated string
sql	whether or not to use SQLite to perform the grouping (not yet implemented).
full.names	names of the aggregation functions should be appended to the output column names
...	additional parameters (such as na.rm) passed to the underlying aggregate functions.

Value

an summary/aggregate dataframe

See Also

[aggregate](#), [bestBy](#)

Examples

```
df <- data.frame(a=runif(12),b=c(runif(11),NA),
                 z=rep(letters[13:18],2),w=rep(letters[20:23],3))

groupBy(df=df, by='w', clmns=c(rep(c('a','b'),3),'z','w'),
        aggregation=c('sum','mean','var','sd','min','max','paste','length'),
        full.names=TRUE, na.rm=TRUE)
# or using SQLite
groupBy(df=df, by='w', clmns=c(rep(c('a','b'),2),'z','w'),
        aggregation=c('sum','mean','min','max','paste','length'),
        full.names=TRUE, sql=TRUE)

## passing a custom function
meantop <- function(x,n=2, ...)
  mean(x[order(x, decreasing=TRUE)][1:n], ...)

groupBy(df, by='w', aggregation=rep(c('mean','max','meantop'),2),
        clmns=rep(c('a','b'),3), na.rm=TRUE)
```

heatmatrix *Simple Heatmap Plot*

Description

This is a very simplified heatmap function: basically a convenient wrapper around the 'image' function.

Usage

```
heatmatrix(x, values=TRUE, clp=c('bottom', 'top'), rlp=c('left', 'right'),  
          xadj=.02, yadj=.3, ylab.cntr=FALSE, cex=1, cex.axis=1, ...)
```

Arguments

x	A matrix
values	boolean: should the values be plotted over each cell?
clp	column label position: either 'bottom' or 'top'.
rlp	row label position: either 'right' or 'left'
xadj	x-adjust of the row labels
yadj	y-adjust of the column labels.
ylab.cntr	boolean for justification of row labels.
cex,	character expansion factor for values in cells if values == true
cex.axis	character expansion factor for axis tick mark labels
...	other parameters passed on to image()

Value

a heatmap plot

See Also

[image](#), [heatmap](#), [heatmap.2](#)

Examples

```
data(mtcars)  
x <- as.matrix(mtcars)  
  
heatmatrix(x)
```

Description

This simple function makes R scatter plots interactive by creating an image and wrapping HTML around it: creating a hyperlinked hyperplot. Hover over the points to see what each is. Click to connect to a table below that will tell you more about each point (if `browse == TRUE`).

Usage

```
hyperplot(x, y = NULL, annout = 1:length(x),
          name = "hyperplot.imagemap", w = 72 * 8, h = 72 * 6,
          link = "internal", browse = TRUE, cex = 1, ...)
```

Arguments

x	a plot-able object, a numeric vector or the name of a numeric vector column in annout.
y	a numeric vector or the name of a numeric vector column in annout. Must be the same length as x.
annout	a named data.frame or table of outliers to annotate the points in the plot. 'x' and 'y' params can indicate column names or numbers of annout.
name	base name of the image & html (map) page that get generated.
w	width of the png image in inches.
h	height of the png image in inches.
link	create a linked lookup table from image to the annout table.
browse	load the html page automatically via R.
cex	character expansion for points
...	other paramters passed on to plot()

Value

HTML page with annotation mapped image

See Also

[browseURL](#)

Examples

```

if(capabilities()["png"] && interactive()) {

  main.hov <- 'Hover over a point to see the name'
  main.subsets <- '(annotated subset in red only)'
  main.click.in <- 'click on points to visit table'
  main.click.out <- 'click on points to visit external site'
  cols <- c('black','red')
  ext.url <- 'http://cran.r-project.org'

  #####
  ## x and y as numeric vectors ##
  #####
  x.out <- nv(rnorm(13,2,sd=5),toupper(letters)[1:13])
  x.in <- nv(rnorm(13,1,sd=.5),toupper(letters)[14:26])
  y.out <- nv(rnorm(13,2,sd=5),toupper(letters)[1:13])
  y.in <- nv(rnorm(13,1,sd=.5),toupper(letters)[14:26])
  x <- c(x.out, x.in)
  y <- c(y.out, y.in)

  ## simplest version
  hyperplot(x,y, main=main.hov)

  ## same but with annotations being supplied as a paramter (instead of names on x)
  names(x) <- NULL
  hyperplot(x,y, annout=toupper(letters), main=main.hov)

  ## annotate only a subset
  hyperplot(x,y, annout=1:13, col=cols[rep(c(T,F), each=13)+1],
            main=paste(main.hov,main.subsets,sep='\n'))

  #####
  ## annout as dataframe #
  #####
  ## x and y as vectors
  x <- nv(x,toupper(letters)) # reinstate the names of x
  df <- data.frame(ab=rep(c('a','b'),13),row.names=toupper(letters))
  hyperplot(x,y, annout=df,
            main=paste(main.hov, main.click.in, sep='\n'))

  ## x and y as names of columns in df
  df <- cbind.data.frame(data.frame(x=x, y=y), df)
  hyperplot(x='x',y='y', annout=df,
            main=paste(main.hov, main.click.in, sep='\n'))

  ## using 'link' column name parameter to specify external links
  df <- cbind.data.frame(df,
                        data.frame(url=ext.url, stringsAsFactors=FALSE))
  hyperplot(x='x',y='y', annout=df, link='url',
            main=paste(main.hov, main.click.out,sep='\n'))
}

```

```
## using reserved column name 'out' as a way to annotate a subset
df <- cbind.data.frame(df, data.frame(out=rep(c(TRUE,FALSE), each=13)))
hyperplot(x='x',y='y', annout=df, col=cols[df$out+1],
          main=paste(main.hov, main.click.in, main.subsets,sep='\n'))

}
```

install.prev.pkg *Install the next oldest package*

Description

This function will recursively search the package archives backwards in time until it finds a version that installs successfully. This function is useful for installing or troubleshooting package dependency trees where one or more packages "require" the very most recent version of R. Rather than upgrading your base R installation, you can use this function to search back in time until you find a version of the package that works with your existing version of R.

Usage

```
install.prev.pkg(pkg.nm, version=NULL,
                 repo.url='https://cran.r-project.org/src/contrib/Archive/')
```

Arguments

pkg.nm	The package name.
version	The version number as #.#-# or #.#.# .
repo.url	The base url for archives of old versions of packages on CRAN.

Value

NULL

Examples

```
#install.prev.pkg('mnormt')
#install.prev.pkg('mvtnorm')
```

labsegs *Buffered Segments for Point Labels*

Description

This function is a wrapper for segments which trigonometrically shortens the lines that are near the "1" end so as not to clutter or overplot the text label it is attached to.

Usage

```
labsegs(x0, y0, x1, y1, buf=.3, ...)
```

Arguments

x0	initial x point coordinate
y0	initial y point coordinate
x1	initial x point coordinate
y1	initial y point coordinate
buf	the buffer between the label at point "1" and the actual segment
...	other parameters passed to segments.

See Also

[segments](#)

Examples

```
x <- rnorm(1000,0,.5)
y <- rnorm(1000,-.3,.15)

labdb <- data.frame(x=seq(-.5,.5,by=.5), y=rep(.85,3))
xlims <- c(-1,1)
ylims <- c(-.5,1)

x0.lbd <- x[rev(order(y))][1:3]
y0.lbd <- y[rev(order(y))][1:3]

par(mfrow=c(1,2))
plot(x,y, xlim=xlims, ylim=ylims, main='segments')
segments(x0=x0.lbd, y0=y0.lbd, x1=labdb$x, y1=labdb$y, col=rainbow(3), lwd=3)
text(x=labdb$x, y=labdb$y, labels=letters[1:3], cex=3, col=rainbow(3))

plot(x,y, xlim=xlims, ylim=ylims, main='labsegs')
labsegs(x0=x0.lbd, y0=y0.lbd, x1=labdb$x, y1=labdb$y, col=rainbow(3), lwd=3, buf=.07)
text(x=labdb$x, y=labdb$y, labels=letters[1:3], cex=3, col=rainbow(3))
```

 leghead

Generate a Color Coded Legend dataframe via head and sum.

Description

'leghead' is part 'head' and part 'summary'. It works best on a sorted dataframe where all you are interested in only the most (or least) abundant rows. An ideal place to use it is in a legend for ~lognormally distributed data. Additionally, an optional row-wise color coding column is added (the color 'gray' is used for missing row names).

Usage

```
leghead(x, n=7, tabulate=FALSE, colors=TRUE, na.name='NA',
        na.col = "white", other.col = "gray", na.last = TRUE)
```

Arguments

x	dataframe or table you wish to summarize
n	the number of rows you wish to display as is
colors	list of vectors or a dataframe
tabulate	the column name to tabulate on if x is an untabulated dataframe and FALSE otherwise
na.name	the new rowname for a row with a missing name
na.col	color for rows labeled as 'NA'
other.col	color for the rows labeled as 'unknown'
na.last	boolean specifying if the na category should be listed last in the table.

Value

A truncated dataframe with a new bottom row summarizing all the truncated ones.

See Also

[summary](#), [head](#), [sstable](#)

Examples

```
e <- data.frame(a=runif(12),b=runif(12), z=rep(letters[13:18],2),w=rep(letters[20:23],3))
tab <- sstable(e, idx.clmns=c('z'), ct.clmns=c('a','b'))
lh <- leghead(tab)
plot(x=lh$a, y=lh$b, cex=lh$sum*3, col=lh$color, pch=20)
legend('topleft', legend=rownames(lh), col=lh$color, pch=20)
```

 m *Regex Match Operator*

Description

A grep/sub-like function that returns one or more back-referenced pattern matches in the form of a vector or as columns in a dataframe (respectively). Unlike sub, this function is more geared towards data extraction rather than data cleaning. The name is derived from the popular PERL regular expression 'match' operator function 'm' (eg. 'extraction =~ m/sought_text/').

Usage

```
m(pattern, vect, names="V", types="character", mismatch=NA, ...)
```

Arguments

pattern	A regular expression pattern with at least one back reference.
vect	A string or vector of strings one which to apply the pattern match.
names	The vector of names of the new variables to be created out of vect. Must be the same length as vect.
types	The vector of types of the new variables to be created out of vect. Must be the same length as vect.
mismatch	What do to when no pattern is found. NA returns NA, TRUE returns original value (currently only implimented for single match, vector returns)
...	other parameters passed on to grep

Value

Either a vector or a dataframe depending on the number of backreferences in the pattern.

See Also

[sub](#), [gsub](#), [regexpr](#), [grep](#), [gregexpr](#).

Examples

```
## single vector output examples
m(pattern="asdf.([A-Z]{4}).",
  vect=c('asdf.AS.fds', 'asdf.ABCD.asdf', '12.ASDF.asdf', 'asdf.REWQ.123'))

Rurls <- c('http://www.r-project.org', 'http://cran.r-project.org',
  'http://journal.r-project.org', 'http://developer.r-project.org')
m(pattern="http://([a-z]+).r-project.org", vect=Rurls)
```

```
# dataframe output examples

data(mtcars)
m(pattern="^[A-Za-z]+) ?(.*)$",
  vect=rownames(mtcars), names=c('make', 'model'), types=rep('character',2))
```

makeElipseCoords	<i>Make Elipse Coordinates</i>
------------------	--------------------------------

Description

Create x & y coordinates for an ellipse from parameters. save.

Usage

```
makeElipseCoords(x0 = 0, y0 = 0, b = 1, a = 1, alpha = 0, pct.range = c(0,1), len = 50)
```

Arguments

x0	x coordinate of center of ellipse.
y0	y coordinate of center of ellipse.
b	y axis stretch factor.
a	x axis stretch factor.
alpha	rotation factor.
pct.range	percentage of the way around the ellipse.
len	number of points used to draw ellipse.

Value

a 2 column (x and y) dataframe with coordinates for drawing an ellipse

Examples

```
makeElipseCoords(x0 = 0, y0 = 0, b = 1, a = 2, alpha = 0)
```

Description

There is no easy way to move point labels around interactively on an a plot in R. This function allows a point and click way to select (using identify) and move (using locator) points by modifying the underlying dataframe.

Usage

```
mvlabs(df, n=nrow(df), x='x', y='y', l='lab', cols=colors()[grep("dark",colors())], ...)
```

Arguments

df	A dataframe with x and y coordinates and text labels
n	the number of points you wish to move
x	the column name of the x axis coordinates
y	the column name of the y axis coordinates
l	the column name of the point labels
cols	the color vector to iterate through while assigning new positions.
...	other paramters passed on to text

Value

a series of violin plots

See Also

[locator](#), [identify](#), [labsegs](#)

Examples

```
x <- rnorm(20); y <- rnorm(20)
df <- data.frame(x,y, lab=as.character(letters[1:20]))
plot(df$x, df$y, pch=''); text(df$x, df$y, df$lab)
## df <- mvlabs(df, 'x','y','lab', n=3)
plot(df$x, df$y, pch=''); text(df$x, df$y, df$lab)
```

 nerge

Named Merge

Description

This function is a wrapper for merge that supports merging multiple vectors and or dataframes

Usage

```
nerge(l, ...)
```

Arguments

l A named list of named vectors (and/or dataframes)
 ... Other parameters passed on to each sub-merge

See Also

[merge](#)

Examples

```
df <- data.frame(a=c(6,7,8), b=c(9,8,7))
rownames(df) <- c('a','d','c')
```

```
l <- list(x=nv(c(1,2),c('a','b')),y=nv(c(2,3),c('b','d')),z=nv(c(4,1),c('c','d')), w=df)
nerge(l, all=TRUE)
```

```
l2 <- list(a=nv(c(1.23, 1.423, 2.343), c('z','y','x')),b=nv(c(6.34,7.34, 12.545),c('z','w','y')))
nerge(l2, all=TRUE)
```

 nv

Create a named vector from a dataframe, table or vector

Description

The '\$' or "[" operators for dataframes and tables do not carry along with them the row names. This function provides a solution for this problem. Additionally this function will accept a vector of values and a corresponding vector of value names—an ideal, in-line way for setting named-vectors as default parameters in new functions.)

Usage

```
nv(x, name)
```

Arguments

x	The source dataframe, table, vector, or factor
name	The column name you would like to pull out as a named vector. OR the names of the vector (if x is a vector)

Value

a named vector or factor

Author(s)

David Schruth

See Also

[vector,name](#)

Examples

```
## example 1: pulling a row.named vector out of a dataframe
df <- data.frame(a=c(1,2,34,4,5,3,23), b=c('z','x','y','z','x','n','p'))
rownames(df) <- letters[1:nrow(df)]
nv(df, 'a')
nv(df, 'b')
```

```
## example 2: a naming vectors from scratch
nv(c(1,2,3), c('a','b','c'))
nv(df$a, df$b)
```

pad

Pad a vector of numerical string with zeros.

Description

This function helps to pad numbers on the left side with zeros so that they may be used to create strings used in filesystem names (for example).

Usage

```
pad(vect, np)
```

Arguments

`vect` a vector of strings representing numbers.
`np` number of zeros to pad to the left of the string.

Value

a (vector of) string(s) with `np` number of zeros padded on as a prefix

Author(s)

Jeremy Tantrum

Examples

```
pad(c(1,10,1000,10000), 4)
```

`parseArgString`

Process Command Line Arguments

Description

generic function for parsing delimited lists from BATCH mode argument strings.

Usage

```
parseArgString(string, delimiter=',', min.param.ct=2, max.param.ct=2, param.range=NULL)
```

Arguments

`string` string to parse.
`delimiter` how the string is delimited into a vector.
`min.param.ct` minimum number of parameters in the vector.
`max.param.ct` maximum number of parameters in the vector.
`param.range` the range of the parameter values.

Value

a vector or value that has been check for validity

Examples

```
## passes
parseArgString('apple,banana,pear', param.range=c("apple","banana","pear","pineapple"))
parseArgString('1,2,3', param.range=c(1,4))

## fails
## Not run:
parseArgString('apple,banana,pear', param.range=c("apple","banana"))
parseArgString('1,2,3', param.range=c(1,2))

## End(Not run)
```

pct

Add Percentage Columns to a Dataframe

Description

This function will add extra columns to an existing dataframe. The second argument 'clmn' should specify which column(s) of the dataframe the percentage should be calculated by dividing each column's row-element by it's sum.

Usage

```
pct(df, clmns)
```

Arguments

df	A dataframe with numeric columns.
clmns	the names of the columns for which the percentage column should be calculated from.

Value

The original dataframe plus extra percentage columns corresponding to original columns in the dataframe.

Examples

```
df <- data.frame(a=c(1,2,3), b=c('x','y','z'), c=c(5,3,2))
pct(df, c('a','c'))
```

 pies

Pie chart scatterplot

Description

Plot pie charts in an XY scatterplot. An overhauled wrapper of the original pie plot function. It is currently very slow: a recommended work around is to plot to something other than the default device (aka png, pdf, etc).

Usage

```
pies(x, show.labels = FALSE, show.slice.labels = FALSE, color.table = NULL,
     radii = rep(2,length(x)), x0=NULL, y0=NULL,
     edges = 200, clockwise = FALSE,
           init.angle = if (clockwise) 90 else 0, density = NULL, angle = 45,
           border = NULL, lty = NULL,
           other.color='gray', na.color='white', ...)
```

Arguments

x	a list of named vectors.
show.labels	boolean specifying if the pie point labels should be plotted.
show.slice.labels	boolean specifying if the pie slice labels should be plotted.
color.table	a named vector of colors. names should correspond to all possible levels of x
radii	a vector of radii used to size the pie points.
x0,y0	a vector of x and y positions for the pie points.
edges	the circular outline of the pie is approximated by a polygon with this many edges.
clockwise	logical indicating if slices are drawn clockwise or counter clockwise (i.e., mathematically positive direction), the latter is default.
init.angle	number specifying the starting angle (in degrees) for the slices. Defaults to 0 (i.e., 3 o'clock) unless clockwise is true where init.angle defaults to 90 (degrees), (i.e., 12 o'clock).
density	the density of shading lines, in lines per inch. The default value of NULL means that no shading lines are drawn. Non-positive values of density also inhibit the drawing of shading lines.
angle	the slope of shading lines, given as an angle in degrees (counter-clockwise).
border	(possibly vectors) arguments passed to polygon which draws each slice.
lty	(possibly vectors) arguments passed to polygon which draws each slice.
other.color	color used for x vector elements for names without corresponding names in the color table
na.color	color used for x vector elements with missing names
...	other arguments passed to polygon

Value

Pie charts as points on a plot

See Also

[pie](#)

Examples

```
## these examples are to the default plot window, which can be slow
## try instead to plot to png or pdf for example

## example 1
pies(
  list(
    a=nv(c(1,2,3),c('one','two','thre')),
    b=nv(c(2,2,3),c('one','two','thre')),
    c=nv(c(1,2,3),c('one','two','thre'))
  ),
  x0=c(0,.5,1),
  y0=c(0,.5,1), radii=6, border=c('gray','black','red')
)

## example 2
n <- 200
n.groups <- 10
n.subgroups <- 6

grps <- paste('gene',seq(1,n.groups), sep='')[round(runif(n,1,n.groups))]
subgrps <- paste('species',seq(1,n.subgroups), sep='')[round(runif(n,1,n.subgroups))]
group.df <- cbind.data.frame(grps,subgrps)
subgroup.list <- by(group.df, group.df$grps, function(x) x$subgrps)

pie.list <- lapply(subgroup.list, table)
col.tab <- nv(rainbow(6), unique(subgrps))

pies(x=pie.list, x0=rnorm(n.groups), y0=rnorm(n.groups),
     radii=10, show.labels=TRUE, show.slice.labels=TRUE, color.table=col.tab)

## example 3 reading from external flat file
## salt.df <- read.delim('/path/to/my/file.tab')
## create a dummy dataset that might live inside the above file
salt.df <- data.frame(salinity=rnorm(25,5), temperature=rnorm(25,25),spec_a=rpois(25,4),
  spec_b=rpois(25,4),
  spec_c=rpois(25,4),
  spec_d=rpois(25,4),
  spec_e=rpois(25,4)
)
## pull out the column names that are specific to pie wedge numbers
```

```

salt.spec.nms <- names(salt.df)[grep('spec',names(salt.df))]
## turn them into a list
pie.list <- lapply(1:nrow(salt.df),
  function(i) as.table(nv(as.vector(as.matrix(salt.df[i,salt.spec.nms])),salt.spec.nms)))
names(pie.list)<- letters[1:25]
with(salt.df, pies(x=pie.list, x0=salinity, y0=temperature, radii=2))

```

plotClock

Plot a simple clock.

Description

Used to create a clock on a plot as a way to keep track of the additional parameter of time for use in animated movies of multiple plots.

Usage

```
plotClock(hour, minute, x0 = 0, y0 = 0, r = 1)
```

Arguments

hour	integer specifying the position of the hour hand.
minute	integer specifying the position of the minute hand.
x0	number specifying the x position of the clock.
y0	number specifying the y position of the clock.
r	number specifying the radius of the clock.

Value

a plot of a clock

raAddArms

Add Arms to a RA plot.

Description

.

Usage

```
raAddArms(epsilon=.55, start=1, end=6, A.shift=0, R.shift=0, ...)
```

Arguments

epsilon .
start .
end .
A.shift .
R.shift .
... other parameters passed to lines.

See Also

[raPlot](#)

raAddAxLabs

Add axis labels to an RA plot.

Description

.

Usage

```
raAddAxLabs(conditions=nv(c('a','b'),c('ref','obs')), normalize=T, add=TRUE, line=2)
```

Arguments

conditions .
normalize .
add .
line .

See Also

[raPlot](#)

raAddSigLines	<i>Add Significance Lines to an RA plot.</i>
---------------	--

Description

.

Usage

```
raAddSigLines(n, end=20, alpha=1e-3, nr=0, A.shift=0, plot=FALSE, ...)
```

Arguments

n	.
end	.
alpha	.
nr	a numeric value indicating the asymptotic normalization ratio line.
A.shift	.
plot	.
...	other parameters passed to lines.

See Also

[raPlot](#)

raPlot	<i>Generate a Ratio Average [RAy] Plot.</i>
--------	---

Description

A plot which turns two vectors of count data into log scaled fold change ratio and average abundance. The plot derives from a Bland-Altman plot and is also very similar to an MA plot. The RA plot is unique, however, in its creative inclusion of the vector-unique 'arms' which are artificially introduced into the plot by adding a <1 epsilon factor before the log function is applied. The name RAy comes from the fact that the aforementioned 'uniques' arms addition makes it strongly resemble a geometric ray. Many of the parameters to the function play off of this convenient anatomical analogy.

Usage

```
raPlot(a, b=NULL, uniques=5, normalize=FALSE,
       nr=0, alpha = 0.01, jitter=FALSE, jit.wgts=NULL,
       rex=1, flat=TRUE, tail=.5, arms=.5, spine=1, border=NULL, plot=TRUE, ...)
```


Arguments

a	a vector of counts for a. can also be a matrix with two columns 1 for a and 2 for b.
b	a vector of counts for b.
uniques	a boolean specifying whether or not to plot the library-unique genes (those with zero counts in one or the other library).
normalize	A boolean specifying whether or not to normalize the counts into proportions.
nr	a numeric value indicating the asymptotic normalization ratio line.
alpha	a statistical significance value.
jitter	whether or not or how much to jitter the a and b counts into surrounding, non-overlapping space.
jit.wgts	a weight vector used to spread the counts of a and b into surrounding, non-overlapping space.
rex	a numeric value specifying the radial expansion of the plot points.
flat	a boolean for the radial expansion of points as a function of both R and A axes.
tail	a numeric or boolean value indicating the line thickness of the two trailing curved significance lines of the RAY.
arms	a numeric or boolean value indicating the line thickness of the two leading straight separator lines of the RAY.
spine	a numeric or boolean value indicating the line thickness of the normalization line (whose y position is specified by mm).
border	a vector of strings used to color the borders of the points.
plot	whether or not to do the actual plot.
...	other parameters passed to plot.

Value

a RAY plot

See Also

[plotMA](#), [link\[edgeR\]maPlot](#)

Examples

```
a <- rnbinom(n=10000, mu=5, size=2)
b <- rnbinom(n=10000, mu=5, size=2)

## the alternative
plot(a,b)
## the raPlot version
raPlot(a, b)

## highlight the condition unique points in the same way as edgeR's "maPlot"
```

```
RA <- raPlot(a, b, pch='')
cond.unique <- apply(cbind(a,b), 1, function(d) any(d==0))
points(RA$A,RA$R, col=c('black','orange')[cond.unique+1])

## try playing with jittering over plotted points
raPlot(a, b, jitter=.3)
```

read.tab

Read in a Tab Delimited File.

Description

This function is a slight (genome annotation friendly) variant of the built-in `read.delim` function in R. Two non-standard defaults have been set: `stringsAsFactors=TRUE`, `quote=""`. An additional parameter `"check.row.ct"`, triggering a `count.fields` call, has been added to further ensure the integrity of large data files.

Usage

```
read.tab(file, check.row.ct = TRUE, stringsAsFactors = FALSE,
         quote = "", header=TRUE, ...)
```

Arguments

<code>file</code>	the name of the file which the data are to be read from.
<code>check.row.ct</code>	logical: use 'count.fields' to independently verify the number of rows read.table reads into memory?
<code>stringsAsFactors</code>	logical: should character vectors be converted to factors?.
<code>quote</code>	the set of quoting characters.
<code>header</code>	boolean specifying if the first row serves as labels for the columns
<code>...</code>	other paramters passed to <code>read.delim</code> .

Value

a dataframe.

regroup	<i>Regroup a dataframe.</i>
---------	-----------------------------

Description

Used to group a dataframe of numbers by a factor that need not be the same length. Find the a factor in the old df and use it to group by the new trumping factor (NA's allowed)

Usage

```
regroup(df, old, new, clmns, funcs=rep('sum',length(clmns)), combine=TRUE)
```

Arguments

df	a dataframe.
old	the ids to match the rows in df to the 'new' grouping ids.
new	the new ids (must be a vector of the same length as 'old'.
clmns	the colums to include in the output.
funcs	the functions to perform on the output (default is to sum) .
combine	Determines wether to combine with existing groupings or to start fresh.

Value

a dataframe with number of rows equal to the number of factor levels in 'new'

Examples

```
df <- data.frame(a=rnorm(20),b=rpois(20,1))  
  
mapping <- data.frame(old=rownames(df), new=rep(c('a','b'),10))  
regroup(df, old=mapping$old, new=mapping$new)
```

rerowname	<i>Rename select rows of a dataframe</i>
-----------	--

Description

Used to easily rename the rows of a dataframe.

Usage

```
rerowname(df, old='NA', new='unknown')
```

Arguments

df	A dataframe with rownames.
old	The row name to be replaced.
new	The replacement row name.

Value

A dataframe with one new rowname

Examples

```
df <- data.frame(a=c(1,2,3), b=c('x','y','z'), c=c(5,3,2))
rownames(df) <- c('p','q','NA')
rerowname(df)
```

spie *Spie charts*

Description

Spie Chart

Usage

```
spie(p1, p2, init.angle=pi, multi, col = rainbow(length(x$radii)), bg=col, lwd=2,
     pie.labs=TRUE, grid=TRUE, grid.labs=TRUE, scale=TRUE, p1.circle=TRUE)
```

Arguments

p1	a positive numeric vector.
p2	a positive numeric vector. Angles are the same than those used for the first pie but radii change according to the values in .
init.angle	initial angle
multi	radius scale multiplier
col	colors of the p2 (foreground) slices
bg	colors of the p1 (background) slices
lwd	line width of the pie wedge boundaries
pie.labs	boolean labels for the pies
grid	boolean
grid.labs	boolean, scale indicators
scale	boolean
p1.circle	boolean

Author(s)

Romain Francois <francoisromain@free.fr> & David Schruth <dschruth@uw.edu>

References

D. G. Feitelson (2003), "Comparing Partitions with Spie Charts". School of Computer Science and Engineering, The Hebrew University of Jerusalem. <https://www.cs.huji.ac.il/w~feit/papers/Spie03TR.pdf>

Michael Friendly (2022), Spie chart – a comparison of two pie charts. <https://www.datavis.ca/gallery/bright-ideas.php>

See Also

[pie](#)

Examples

```
p1 <- c(0.12, 0.3, 0.26, 0.16, 0.04, 0.12)
p2<- c(0.06, 0.15, 0.52, 0.14, 0.08, 0.05)
plot(p1, p2, multi=c(.5, 1, 1.5, 2))
```

sstable

Sum Sorted Tabulation

Description

A wrapper for the "table()" function that also calculates the row-wise sum and sorts by the new column.

Usage

```
sstable(x, idx.clmns, ct.clmns = NULL, na.label = "NA")
```

Arguments

x	list of vectors or a dataframe
idx.clmns	index columns
ct.clmns	count columns
na.label	row label used for na columns

Value

A dataframe sorted by the count columns.

Author(s)

David Schruth

See Also

ledghead, table, order, sort

Examples

```
e <- data.frame(a=runif(12),b=runif(12), z=rep(letters[13:18],2),w=rep(letters[20:23],3))
e <- data.frame(a=runif(10),b=runif(10), z=rep(letters[12:16],2),w=rep(letters[20:24],2))
sstable(e, idx.clmns=c('z','w'), ct.clmns='a')
sstable(e, idx.clmns=c('z'), ct.clmns=c('a','b'))
sstable(e, idx.clmns=c('z','w'))
e <- data.frame(a=10,b=0, z=as.factor(NA))
sstable(e, 'z', c('a','b'))
e <- data.frame(a=10,b=0, z=NA, w=NA)
sstable(e, 'z', c('a','b'))
e <- data.frame(a=runif(10),b=runif(10),m=rep(c('one','two'),5),
               z=factor(rep('z',10), levels=c('z','x')))
sstable(e, idx.clmns=c('m','z'))
```

 stats

Descriptive Statistics

Descriptionadapted from <http://legacy.ncsu.edu/ST370/distance/r1ab/>**Usage**

```
stats(x, by, quantiles=c(.25,.75))
```

Arguments

x	list of distributions to characterize
by	grouping variable presumes that x is univariate
quantiles	any pair of values >0 : <1

Value

statistics on each distribution

Author(s)

Dave McArthur <dmca@ucla.edu>

See Also

violins, summary

 tab2df

Table to Data Frame

Description

Convert a table to a dataframe while perserving the same number of columns and rows and names of each.

Usage

```
tab2df(x, ...)
```

Arguments

`x` a table or matrix class object (output from the table command).
`...` other arguments passed to `data.frame(...)`.

Value

a dataframe

See Also

table

Examples

```
x <- data.frame(a=runif(10),b=runif(10), z=rep(letters[1:5],2))
as.data.frame(x)
tab2df(x)
x <- nv(rnorm(10), letters[1:10])
as.data.frame(x)
tab2df(x)
x <- nv(rnorm(2), c('x.b','y.b'))
as.data.frame(x)
tab2df(x)
x <- nv(rnorm(2), c('b.x','b.y'))
as.data.frame(x)
tab2df(x)
e <- data.frame(a=runif(10),b=runif(10), z=rep(letters[13:17],2))
x <- as.table(sapply(c('a','b'),function(cc) by(e[, 'a'],list(e$z), sum)))
as.data.frame(x)
tab2df(x)
x <- as.table(by(1:10, list(a=rep(1:5,2),b=rep(1:2,5)), sum))
as.data.frame(x)
```

```

tab2df(x)
x <- as.table(nv(c(54,34), c('a','b')))
  as.data.frame(x)
tab2df(x)

x <- table(a='x',b='y')
tab2df(x)

```

textplot

A Text-Only Plot

Description

Generate a new plot window with just text centered in the middle. This is ideally used in conjunction with the 'layout' command to label columns and rows of the grid.

Usage

```
textplot(..., x=1, y=1)
```

Arguments

...	parameters passed to the 'text' function
x	the x position of the text.
y	the y position of the text.

Value

A new plot window with just text

See Also

[layout](#), [text](#)

Examples

```

layout(rbind(c(1,1,1),c(2,3,4), c(5, 6,7)),
        widths=c(5, 10,10) , heights=c(5, 10,10))
textplot('title', cex=2)
textplot('row 1', srt=90, cex=2)
plot(1,2)
hist(c(1,2,34,4,3,2,2))
textplot('row 2', srt=90, cex=2)
pie(c(1,23,3,1,1,2,3,4,54,5))
plot(c(1,2,4,4,23,2), c(1,2,4,3,2,2))

```

`usr2lims`*Grab and adjust the current plot dimensions*

Description

This is a simple function which grabs the current plot dimensions and adjusts them by shrinking them by 4

Usage

```
usr2lims(adj=.04)
```

Arguments

`adj` The automatic adjustment factor 'plot' adds to buffer the specified plot dimensions.

Value

A 2 item (x and y) list of 2 item (min and max) vectors for x and y limits of the current plot area

See Also

`par`

Examples

```
plot(c(0,1), c(0,1))
usr2lims()
```

`vennMatrix`*Create a Venn Ready Matrix out of a List of Factors*

Description

The limma package has great functions for making venn diagrams from a matrix. This function is provides upstream functionality to turn a list of factors into this required input format.

Usage

```
vennMatrix(l)
```

Arguments

`l` a named list of factors

Value

a matrix with columns for list elements and rows with globally unique factor levels

See Also

venCounts

Examples

```
l <- list(a=factor(c('x','y','z')), b=factor(c('w','x','v')))
vennMatrix(l)
```

volins

Plot a Series of Vioplot Violins

Description

This is an improvement on the original vioplot function (combination of a box plot and a kernel density plot). You can now pass a list of violins and a list of colors. Optionally it also returns statistics on the distributions as well.

Usage

```
violins(x, by, range = 1.5, h = NULL, ylim = NULL, names =
  NULL, horizontal = FALSE, col = "transparent", border
  = "black", lty = 1, lwd = 1, rectCol = "grey50",
  colMed = "grey80", pchMed = 19, at, add = FALSE, wex =
  1, drawRect = TRUE, main = "", xlab = "", ylab = "",
  connect = c("median", "mean", "hubermu", "deciles"),
  SD.or.SE = c("SD"), connectcol = c("lightblue",
  "cyan", "darkred", "grey"), las = 2, stats = FALSE,
  quantiles = c(0.1, 0.9), CImed = TRUE, deciles = TRUE)
```

Arguments

x	list of vectors or a dataframe
by	accepts 'by' object for grouping
range	a factor to calculate the upper/lower adjacent values.
h	the height for the density estimator, if omit as explained in sm.density, h will be set to an optimum.
ylim	y limits.

names	one label, or a vector of labels for the datas must match the number of datas given.
horizontal	description of var 1.
col,border,lty,lwd	Graphical parameters for the violin passed to lines and polygon.
rectCol,colMed,pchMed	Graphical parameters to control the look of the box.
at	position of each violin. Default to 1:n
add	logical. if FALSE (default) a new plot is created
wex	relative expansion of the violin.
drawRect	logical. the box is drawn if TRUE.
main	main title for the plot.
connect	connects a violin plot series at the medians with line segments
connectcol	line color of the median connecting line segments
xlab	x axis label
ylab	y axis label
stats	print out statistics
SD.or.SE	Boolean for standard deviation or standard error
las	axis tick mark labels orientation
quantiles	map any pair of quantiles (as dotted box) in addition to Q1 & Q3, but are not shown when "c(0,0)" and arg is passed to descriptive stats when 'stats'=TRUE
CImed	portrays 95 percent confidence intervals for the median (as solid box)
deciles	maps deciles 0.1:0.9 (as thin lines) independently of 'quantiles' and can be connected when 2 or more variables are plotted.

Value

a series of violin plots

Author(s)

Daniel Adler <dadler@uni-goettingen.de>, David Schruth, Dave McArthur <dmca@ucla.edu>

See Also

vioplot(vioplot), [sm.density](#)

Examples

```
n <- rnorm(130, 10, 3)
p <- rpois(110, 4)
u <- runif(300, 0, 20)
l <- rlnorm(130, log(2))
```

```

g <- rgamma(140, 3)
e <- rexp(160)

violins(list(e=e, p=p,u=u,n=n,l=l,g=g), ylim=c(0,20),
        col=c('purple', 'lightblue', 'lightgreen', 'red', 'orange', 'yellow'),
        stats=TRUE)

```

wjitter	<i>Weighted Jitter</i>
---------	------------------------

Description

Use weights to jitter values away from their current value.

Usage

```
wjitter(x, w, amount=.43)
```

Arguments

x	a vector of values
w	a vector of weights of the same length as x
amount	the amount to jitter (passed to the parameter by the same name in the jitter function)

Value

A weighted jittered vector of the same length as x

Examples

```

x <- seq(1,20)
w <- runif(20, 0,1)
plot(x,wjitter(w,x))

```

write.delim	<i>Write a (tab) delimited text file.</i>
-------------	---

Description

A simple wrapper for write.table with the same options as read.delim

Usage

```
write.delim(df, file, quote = FALSE, row.names = FALSE, sep = "\t", ...)
```

Arguments

<code>df</code>	a dataframe.
<code>file</code>	outputfile path.
<code>quote</code>	should elements of the dataframe be quoted for output.
<code>row.names</code>	should the output include rownames.
<code>sep</code>	the delimiter between fields.
<code>...</code>	other parameters passed to <code>write.table</code> .

Value

A tab delimited text file

See Also

[read.delim](#)

Examples

```
## Not run:  
x <- data.frame(a = I("a \" quote"), b = pi)  
write.delim(x, file = "foo.tab")  
  
## End(Not run)
```

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