Package ‘prada’

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Title Data analysis for cell-based functional assays

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Collate AllGenerics.R AllClasses.R methods-cytoFrame.R
methods-misc.R analysePlate.R as.all.R barploterrbar.R
fitNorm2.R gateMatrix.R getPradaPar.R plotNorm2.R plotPlate.R
readCytoSet.R readFCS.R readSDM.R removeCensored.R thresholds.R
touchFCS.R tcltkProgress.R getAlphanumeric.R

Description Tools for analysing and navigating data from high-throughput phenotyping experiments based on cellular assays and fluorescent detection (flow cytometry (FACS), high-content screening microscopy).

License LGPL

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LazyLoad yes

NeedsCompilation yes

R topics documented:

analysePlate .......................................................... 2
as.all ................................................................. 3
barploterrbar ........................................................ 4
cframe ............................................................... 5
combineFrames ....................................................... 5
csApply ............................................................... 6
cset ................................................................. 7
cytoFrame-class ..................................................... 7
cytoSet-class ....................................................... 9
analysePlate

Apply a statistic to the data from each well in a plate

Description

Apply a statistic to the data from each well in a plate

Usage

```r
analysePlate(x, wellcol="well", wellrange, statfun, platename, plotdir=".", ...)```

Arguments

- `x`: data frame. It must contain a column whose name is the value of `wellcol`, and further columns that are needed by the function named by `stat`.  
- `wellcol`: character of length 1. Name of a column in `x` that contains the well ID.  
- `wellrange`: vector of the same type as `x[, wellcol]`. All values `x[, wellcol]` must be contained in `wellrange`.  
- `statfun`: character of length 1. Name of a function that can calculate a statistic from selected rows of `x`.  
- `platename`: character of length 1. The name or ID of this plate, which will be used for graphics output filenames and as the value of the column `platename` of the return value.  
- `plotdir`: character of length 1. The name of directory where diagnostic plots will be saved.  
- `...`: further arguments that are passed on to `statfun`. 
as.all

Details
The semantics of this function are similar to \texttt{tapply}, but some additional checking and reporting is performed, and the return value is a data frame.

Value
A data frame with number of rows equal to \texttt{length(wellrange)}. Rows (wells) for which there is no data contains NAs. The columns comprise \texttt{platename}, \texttt{well-ID} (from \texttt{x[, wellcol]}), and the results from \texttt{statfun}.

Author(s)
Wolfgang Huber

Examples
 Calling \texttt{as.all} without introduction of NAs

Description
Coercion without introduction of NAs

Usage
\texttt{as.all(x, what)}

Arguments
\begin{itemize}
\item \texttt{x} an object.
\item \texttt{what} character of length 1.
\end{itemize}

Details
The function calls do.call(paste("as.", what, sep=""), list(x)), and checks whether any NAs were introduced.

Value
A vector of type \texttt{what}

Author(s)
Wolfgang Huber

See Also
\texttt{as}

Examples
Calling \texttt{as.all} with introduction of NAs

as.all(runif(5)*10, "integer")
barploterrbar  

**Barplot with error bars.**

**Description**

Barplot with error bars.

**Usage**

```r
barploterrbar(y, yl, yh, barcol="orange", errcol="black", horiz=FALSE, 
w=0.2, ylim=c(0, max(yh)*1.05), ...) 
```

**Arguments**

- `y`: Numeric vector.
- `yl`: Numeric vector of same length as `y`.
- `yh`: Numeric vector of same length as `y`.
- `barcol`: Color of the bars.
- `errcol`: Color of the error bars.
- `horiz`: Logical. As in `barplot`.
- `w`: The plot limits. The default value will cause the error bars to fit nicely on the plotting device.
- `ylim`: Size of the error bar ticks.
- `...`: Further arguments that get passed on to `barplot`.

**Details**

The function calls `barplot` with `y` and decorates it with error bars according to `yl` and `yh`.

**Value**

The function is called for its side effect, producing a plot.

**Author(s)**

Wolfgang Huber [http://www.dkfz.de/abt0840/whuber](http://www.dkfz.de/abt0840/whuber)

**See Also**

`barplot`

**Examples**

```r
y <- matrix(runif(80), ncol=5)
ym <- apply(y, 2, mean)
dy <- apply(y, 2, sd)*2/sqrt(nrow(y))
barploterrbar(ym, ym-dy, ym+dy, barcol="#0000c0", errcol="orange", 
               ylim=c(0, max(ym+dy)))
```
A sample cytoFrame object - German Cancer Research Center Heidelberg -

Description
Archived cytoFrame object from a MAP kinase screen conducted at the German Cancer Research Center Heidelberg. In the fluorescence channel 3 the expression of a YFP tag and in channel 7 the activation state of ERK2 was measured.

Usage
##cytoFrame object, see examples for details

Format
cytoFrame object

Source
German Cancer Research Center Heidelberg, Germany

Examples
data(cytoFrame)

combineFrames
Combine the cytoFrames within a cytoSet according to some grouping factor

Description
Combine the cytoFrames within a cytoSet according to some grouping factor.

Usage
combineFrames(x, by)

Arguments
x cytoSet.
by factor. Length must be same as that of x.

Value
cytoSet.
Author(s)

Wolfgang Huber <huber@ebi.ac.uk>

Examples

```r
cset <- readCytoSet(path = system.file("extdata", package = "prada"),
    pattern = "[A-Z][0-9][0-9]"$)
nr1 <- csApply(cset, nrow)
sm1 <- csApply(cset, sum)

fac <- factor(c(1, 1, 2, 2, 2, 2))
cc <- combineFrames(cset, fac)
nr2 <- csApply(cc, nrow)
sm2 <- csApply(cc, sum)

stopifnot(all(nr2 == tapply(nr1, fac, sum)))
stopifnot(all(sm2 == tapply(sm1, fac, sum)))
```

Description

This a wrapper for `sapply` for objects of class `cytoSet`.

Usage

```r
csApply(X, FUN, ..., simplify = TRUE)
```

Arguments

- `X`: cytoSet.
- `FUN`: the function to be applied.
- `...`: optional arguments to `FUN`.
- `simplify`: logical; should the result be simplified to a vector or matrix if possible? Gets passed on the `sapply`.

Details

A wrapper for `sapply`.

Value

Like `sapply`: If `FUN` always returns a scalar, then the value of this function is a named vector. If `FUN` always returns a vector of length `n`, then the value of this function is an `n x length(X)` matrix with dimnames. Else, the value of this function is a named list whose values are the return values of the individual calls to `FUN`.

Author(s)

Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber)
cset

See Also

sapply

Examples

cset=readCytoSet(path=system.file("extdata", package="prada"),
   pattern="[A-Z][0-9][0-9]"
)cApply(cset, nrow)
cApply(cset, colMeans)

cset                                           A sample cytoSet object - German Cancer Research Center Heidelberg

Description

Archived cytoSet object from a MAP kinase screen conducted at the German Cancer Research Center Heidelberg. In the fluorescence channel 3 the expression of a YFP tag and in channel 7 the activation state of ERK2 was measured. The set contains measurements from 5 wells of a 96 well plate

Usage

## cytoSet object, see examples for details

Format

cytoSet object

Source

German Cancer Research Center Heidelberg, Germany

Examples

data(cytoSet)

cytoFrame-class 'cytoFrame': a class for storing observed quantitative properties from a population of cells, most likely from a FACS run or, alternatively, from automated microscopy

Description

This class represents the data contained in a FCS 3.0 file or similar data structures.
Details

Although objects of class cytoFrame can be used to hold arbitrary data of cell populations, the main focus lies on flow-cytometry data.

FCS 3.0 is the Data File Standard for Flow Cytometry, Version FCS3.0. See the vignette of this package for additional information on using the object system for handling of flow-cytometry data.

Creating Objects

Objects can be created using

```
new('cytoFrame',
  exprs = ...., # Object of class matrix
  description = .... # Object of class character
)
```
or the function `readFCS`.

Slots

exprs: Object of class matrix containing the measured intensities. Rows correspond to cells, columns to the different channels. The colnames attribute of the matrix is supposed to hold the names or identifiers for the channels. The rownames attribute would usually not be set.
description: A named character vector containing the experiment description as key-value pairs.
well: A single integer vector giving the position of the well on a microtitre plate. This only applies when using the object within a cytoSet collection and will usually be filled in by the function `readCytoSet`.
gate: An object of class gateSet. This object can be used to select defined subsets of the data, a process referred to as gating in the analysis of flow-cytometry data.

Methods

[  subsetting. Returns an object of class cytoFrame. The subsetting is applied to the exprs slot, while the description slot is unchanged.
  exprs, exprs<-  extract or replace the intensities.
  description, description<-  extract or replace the description.
  show  display summary.
  plot  scatterplot for cytoFrame objects. The additional argument gate can be used to plot subsets of the data defined by either an object of class gate or by a character vector giving the name of one of the gates in the list.
  gate, gate<-  extract or replace the list of gates.
  ncol, nrow  extract the dimensions of the data matrix.
  appendGate  Append a gate or gateSet to the gate slot.
  drawGate  Create an object of class gate or gateSet based on a selection made from the data.
  hist  Draw a histogram of the data

Author(s)

Florian Hahne, Wolfgang Huber

See Also

readFCS, cytoSet, gate, gateSet
Examples

```r
intens <- matrix(runif(100), ncol=4)
colnames(intens) <- c("FL1-H", "FL2-H", "FL3-H", "FL4-H")

a <- new("cytoFrame",
       exprs=intens,
       description=c(name="example data", date=date()))

description(a)
dim(exprs(a))

a[1:3, -4]

plot(a)

## Not run:
g1 <- drawGate(a, name="Gate1")

## End(Not run)
```

Description

This class is a container for a set of `cytoFrame` objects.

Creating Objects

Objects can be created using the function `readCytoSet` or via

```r
new('cytoSet',
    frames = ...., # environment with cytoFrames
    phenoData = .... # object of class phenoData
    colnames = .... # object of class character
)
```

Slots

- `frames`: An environment containing one or more `cytoFrame` objects.
- `phenoData`: A `phenoData`. Each row corresponds to one of the cytoFrames in the `frames` slot. It is mandatory that the pData has column named `name`.
- `colnames`: A character object with the (common) column names of all the data matrices in the cytoFrames.

Methods

- `[`, `[[` subsetting. If `x` is `cytoSet`, then `x[i]` returns a `cytoSet` object, and `x[[i]]` a `cytoFrame` object. The semantics is similar to the behavior of the subsetting operators for lists.
- `colnames`, `colnames<-` extract or replace the `colnames` slot.
- `phenoData`, `phenoData<-` extract or replace the `phenoData` slot.
show display summary.

plot Scatterplot of one or all (consecutively) cytoFrame objects. The additional argument gate can be used to plot subsets of the data defined by an object of class gate or gateSet.

hist Draw histogram of the data. The additional argument variable can be used to subset to one variable prior to plotting.

Important note on storage and performance

The bulk of the data in a cytoSet object is stored in an environment, and is therefore not automatically copied when the cytoSet object is copied. If x is an object of class cytoSet, then the code

\[ y \leftarrow x \]

will create a an object y that contains copies of the phenoData and administrative data in x, but refers to the same environment with the actual fluorescence data. See below for how to create proper copies.

The reason for this is performance. The pass-by-value semantics of function calls in R can result in numerous copies of the same data object being made in the course of a series of nested function calls. If the data object is large, this can result in a considerable cost of memory and performance. cytoSet objects are intended to contain experimental data in the order of hundreds of Megabytes, which can effectively be treated as read-only: typical tasks are the extraction of subsets and the calculation of summary statistics. This is afforded by the design of the cytoSet class: an object of that class contains a phenoData slot, some administrative information, and a reference to an environment with the fluorescence data; when it is copied, only the reference is copied, but not the potentially large set of fluorescence data themselves.

However, note that subsetting operations, such as

\[ y \leftarrow x[i] \]

do create proper copies, including a copy of the appropriate part of the fluorescence data, as it should be expected. Thus, to make a proper copy of a cytoSet x, use

\[ y \leftarrow x[\text{seq(along}=x)] \]

Author(s)

Florian Hahne, Wolfgang Huber http://www.ebi.ac.uk/huber

See Also

readCytoSet, cytoFrame, gate, gateSet

Examples

cset<-readCytoSet(path=system.file("extdata", package="prada"),
                  pattern="[A-Z]\[0-9]\[0-9]\$")
cset
pData(cset)
cset[[1]]
cset["fas-Bcl2-plate323-04-04.A02"]
cset["fas-Bcl2-plate323-04-04.A02"]
cset[1:3]
devDims

Device Dimensions for plate plots

Description

Calculate device dimensions for plate plots

Usage

devDims(width, height, ncol=12, nrow=8, res=72)

Arguments

width

Device width in inches.

height

Device width in inches.

ncol

Number of columns for plate plot.

nrow

Number of rows for plate plot.

res

The resolution of the graphic device used for plotting.

Details

The function computes the device dimensions needed to create plate plots that fit perfectly in the
device. This is necessary to retain the aspect ratio of the plots.

One of width or height need to be specified, the missing value will be computed.

Value

A list with items width, height, pwidth and pheight. These are the width and height values in
inches and pixels respectively.

Author(s)

Florian Hahne

See Also

plotPlate

Examples

devDims(width=10)
Description

Calculates what R thinks to be the resolution of the current graphic device.

Usage

devRes()

Details

This function may be used to get the resolution of the current graphics device. This can be important when calculating pixel coordinates for the output graphic.

Value

A vector with items x res and y res, the resolution in x and y direction respectively.

Author(s)

Florian Hahne

See Also

plotPlate

Examples

devRes()

fitNorm2

Fit bivariate normal distribution.

Description

Fits a bivariate normal distribution into a data set of paired values and selects data points according to their standard deviation from the fitted distribution.

Usage

fitNorm2(x, y=NA, scalefac=1, method="covMcd", noise, gateName = "fitNorm")
Arguments

x | Numeric vector containing x-value or n by 2 matrix containing x and y values or object of class cytoFrame.
y | Numeric vector containing y-value (optional). The length of x must be the same as that of y.
scalefac | Numeric vector giving factor of standard deviations used for data selection (all points within scalefac standard deviations are selected).
method | One of covMcd or cov.rob defining method used for computation of covariance matrix.
noise | Numeric or logical index vector defining value pairs in x that are not used for fitting of distributions. Can be used to deal with noisy data.
gateName | Character giving the name of the gate object.

Details
Computes the densities of a bivariate normal distribution from the covariance matrix of the paired data. Covariance matrices are acquired either by function covMcd (considerably faster) or by function cov.rob.

Value
A list containing items mu (midpoint of distribution), S (covariance matrix), p (density values for each data pair), sel (selection of data points), scalefac (factor of standard deviations used for data selection), data (x and y values of data points) and gate, an object of class gate containing the selection.

Author(s)
Florian Hahne

See Also
cov.rob, covMcd, plotNorm2

Examples
sampdat <- readFCS(system.file("extdata", "fas-Bcl2-plate323-04-04.A01", package="prada"))
nfit <- fitNorm2(exprs(sampdat[,1:2]), scalefac=2)
plotNorm2(nfit, selection=TRUE, ellipse=TRUE)

description
In flow-cytometry analysis, regions in two-dimensional projections of the data space often have to be selected. Objects of this class can store the properties of these selections.
Creating Objects

Objects can be created using methods of the generic function `drawGate` or via
new("gate",
gateFun = ...., # function returning logical vector
colnames = .... # object of class character and length 2
logic = .... # object of class character )

Slots

name: A character vector for the name of the gate object. You can reference the object by its name
for subsequent operations (e.g. plotting).
gateFun: A function call together with necessary arguments to produce a logical vector when
applied on the data.
colnames: The colnames of the data matrix to which the gating function is to be applied.
logic: A character object, either & or |. This specifies the logical operation that will be applied
when combining the selection from the gate with other object of that class. See link(gateSet)
for additional information on combining gates.
type: A character giving the type of the object. This is currently not used but might become
important in the future.
boundaries: A matrix with two columns giving the boundaries of the gate in two dimensional
space. Can be used to superimpose the gate boundaries on a plot using lines().

Methods

applyGate: applyGate(x, data) applies the gating of object x on data objects of class cytoFrame
or matrix. In the former case x may be of class gate, gateSet, character, numeric or
logical. See vignette for details.
show display summary.
names, names<- access and replace slot name.
as.gateSet Convert gate object to gateSet object
combineGates Combine multiple gate objects to one gateSet object
lines Draw the boundaries of the gate.

Author(s)

Florian Hahne

See Also
cytoFrame, gateSet

Examples

sampdat <- readFCS(system.file("extdata", "fas-Bcl2-plate323-04-04.A01",
package="prada"))
g1 <- new("gate", name="test1", gateFun=function(x)x[,"FSC-H"]<500, logic="&",
colnames="FSC-H", type="misc")
g1
gateSet-class

In flow-cytometry analysis, regions in two-dimensional projections of the data space often have to be selected. Objects of this class can store the properties for several of these selections.

Creating Objects

Objects can be created using methods of the generic function `drawGate` or via

```r
new("gateSet",
glist = ....,  # object of class list
)
```

Slots

- **name**: Object of class character giving the name of the object. You can reference the object by its name for subsequent operations (e.g. plotting).
- **glist**: Object of class "list" with items of class `gate`. The individual `gate` objects will be combined according to the value of their slot `logic`.

Methods

- **applyGate**: `applyGate(x, data)` applies the gating of object `x` on data objects of class `cytoFrame` or matrix
- **length**: length of slot `glist`
- **show**: display summary
- **names, names<-**: extract or replace the names of the individual `gate` objects.
- `[]` subset to `gateSet` objects.
- `[[` subset to individual `gate` objects.
- **appendGates**: append a `gate` or `gateSet` to a `cytoFrame`

Author(s)

Florian Hahne
getAlphaNumeric

Convert from plate coordinates to alphanumeric notation.

description

Given an array of (x,y) well coordinates, this function returns the corresponding alphanumeric notation.

Usage

getAlphaNumeric(horizontal, vertical)

Arguments

  horizontal  Integer coordinate of horizontal well location.
  vertical    Integer coordinate of vertical well location.

Value

getAlphaNumeric returns a list containing id, the full alphanumeric id of the well(s), id.alpha, the alpha part of the id, and id.num, the numeric part of the id.

Author(s)

Joseph Barry <joseph.barry@embl.de>
getPradaPar

See Also
convertWellCoordinates

Examples

## To obtain the alpha, numeric and alphanumeric information for a single well
getAlphaNumeric(horizontal=1, vertical=1)

## To obtain only the alphanumeric ids of a tetrad in the corner of a 1536 well plate
getAlphaNumeric(horizontal=c(31,31,32,32), vertical=c(47,48,47,48))$id

getPradaPar Set and query global parameters for functions in the prada package

Description
Set and query global parameters for functions in the prada package

Usage

setPradaPars(pars)
getPradaPar(parname)

Arguments
pars Named list
parname Character string of length 1

Details
TBA

Value
For getPradaPar, the value of the list element with name parname of the global parameters list. The function setPradaPars is invoked for its side effect, which is assigning a value to the global parameters list. It returns the value invisible(NULL).

Author(s)
Wolfgang Huber http://www.ebi.ac.uk/huber

Examples

setPradaPars(list(tomato=1, apple="two", pear=as.integer(3)))
getPradaPar("pear")
plotNorm2

Description

Plots objects derived from function `fitNorm2` in false color representation.

Usage

```r
plotNorm2(fn, colrange=c("gray82", "blue"), center=TRUE, selection=FALSE,
    ellipse=FALSE, pch=20, cex=1, col="dens", ...)
```

Arguments

- **fn**: List. Object derived from function `fitNorm2`
- **colrange**: Character vector with valid color identifiers (e.g., name or RGB values) from which a smooth color palette is derived.
- **center**: Logical. Assign center of distribution.
- **selection**: Logical. Mark all points beyond selection.
- **ellipse**: Logical. Plot area and borders of selection as ellipse.
- **pch**: see `par`
- **cex**: see `par`
- **col**: see `par` or special cases `dens` for coloring according to density and `prob` for coloring according to probability.
- **...**: further arguments that are passed on to `plot`.

Details

Produces a scatterplot of paired data showing the densities of the fitted bivariate distribution from function `fitNorm2` in false color representation. Additionally, a selection of data points can be highlighted either by marking outliers or by showing its area.

Value

A list containing items `p`, `cov`, `mu`, `S` (density values for each data pair, resulting object from call to `cov.rob`, midpoint of distribution, covariance matrix).

Author(s)

Florian Hahne

See Also

`fitNorm2`

Examples

```r
sampdat <- readFCS(system.file("extdata",
    "fas-Bcl2-plate323-04-04.A01", package="prada"))
nfit <- fitNorm2(exprs(sampdat[,1:2]), scalefac=2)
plotNorm2(nfit, selection=TRUE, ellipse=TRUE)
```
plotPlate

Plot a well statistic for microtiter plates.

Description

Plot a well statistic in false color representation or using a self-defined grid plotting function. The plot is supposed to resemble the physical geometry of a microtitre plate.

Usage

plotPlate(x,nrow = 8, ncol = 12, col=c("red", "blue"),
ind = 1:(ncol*nrow), xrange=function(y) range(y, na.rm=TRUE), na.action = "zero",
main, char, desc = character(2), add=FALSE, gridFun="default",
funArgs=NULL,...)

Arguments

x Numeric vector of length ncol*nrow or matrix with ncol*nrow rows (except if argument ind is specified). If of class matrix, the use of argument gridFun is expected.
nrow Numeric of length 1. The number of rows of the plate.
col Numeric of length 1. The number of columns of the plate.
char Character vector. Usually the names of two or three colors between which the color map is interpolated, using the function colorRampPalette.
ind Optional integer vector of equal length as x. It indicates the position of the respective value of x on the plate. Can be used to adress the problem of missing values. Each well that is not allocated a value of x by ind will not be plotted.
xrange Numeric vector of length two giving thwe range of x that is mapped into the color scale. Alternatively, this can be a function which takes the values of x as input and creates such a vector.
na.action Character. One of "zero" "omit" or "xout". How should the wells for which x is NA be treated? For "zero", they are plotted as if the value were 0. For "omit", they are omitted. For "xout", they are crossed out. When x is a matrix, na.action is only applied to rows containing nothing but NAs. Further special treatment of NA values in matrices need to be implemented in gridFun.
main Character of length 1. Plot title.
char An optional character vector of equal length as x (except if argument ind is specified) to be used for well annotation. Each element of the vector may contain a string to be superimposed on the respective well or NA for no plotting.
desc Character of length 2. Legend for the two extremes of the colorbar, e.g. 'act' and 'inh'.
add Logical. If TRUE add plate plot to current plot. May be used when plotting in grid layout panels.
gridFun Character. The name of the plotting function to create individual graphs for each well. See functions .drawCircle and .drawPie for examples.
funArgs Dataframe with argument values to be passed to gridCall. For each argument specified in gridCall there must be one column with the argument name as col-name and the argument values for every well.
Further graphical parameters that can be used to control the output of plotPlate:

- **cex.main**: expansion factor for title.
- **cex.lab**: expansion factor for label.
- **cex.char**: expansion factor for well annotation.
- **cex.legend**: expansion factor for well legend labels.
- **cex.desc**: expansion factor for well legend description.

**Details**

You may use this function either to create plots showing a single-value per well statistic for microtiter plates, or you can use a self-made plotting function using a combination of any valid grid commands to produce arbitrary plots in a plate array format. These plots may also show multifactorial data. Self-defined plotting functions need to have `data` as first argument. `plotPlate` passes all data values for the respective well to the plotting function. Any further arguments may be passed on using argument `funArgs`. See `.drawCircle` and `.drawPie` for examples of valid plotting functions and the vignette for detailed information. Note that using `funCall` overrides some of the default functionalities, e.g. plotting of legends and alters the treatment of NA values.

Argument `ind` allows the user to indicate the position (well number) for each element of vector `x` on the plate. This can be used either to change the order in which elements of `x` are to be plotted or to deal with the problem of missing data for some of the wells on a plate.

To further increase the amount of information of the platePlot one may decorate wells with short annotations using argument `char`. Each element of `char` != NA will be superimposed on the respective well (see examples).

**Value**

The function produces a plot in the active graphics device. It returns a list with four elements. The element `which` is a vector with the indices of those elements in `x` that were plotted (see argument `na.action`). The element `coord` is a `length(which)` by 4 matrix in which each row specifies the corners of a rectangle that contains a well. It is intended to be used as an argument to a subsequent call to `imageMap`. Elements `width` and `height` may be used to open a graphic devices that can hold the plate plot with the correct aspect ratio.

**Author(s)**

Florian Hahne, Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber)

**See Also**

- `imageMap`

**Examples**

```r
plotPlate(runif(96), main="example 1", col=c("#0000e0", "#e00000"), desc=c("act", "inh"))
plotPlate(runif(384), nrow=16, ncol=24, main="example 2", col=c("#0000e0", "white", "#e00000"))
plotPlate(runif(48), main="example 3", col=c("#0000e0", "#e00000"), ind=c(1:24, 73:96))
x <- runif(96)
x[sample(96, 10)] <- NA
plotPlate(x, main="example 4", col=c("#0000e0", "#e00000"),
char=c(rep(NA, 72), LETTERS[1:24]), na.action="xout")
plotPlate(runif(96, min=0.1, max=0.5), gridFun=".drawCircle")
plotPlate(matrix(runif(288), ncol=3), gridFun=".drawPie",
```
funArgs=as.data.frame(matrix(2:4, ncol=3, nrow=96, byrow=TRUE))

---

**Description**

Show progress of a task in a tcltk window as percentage

**Usage**

```r
progress(title="processing task...", message="", sub="")
updateProgress(percentage, autoKill=FALSE, sub="")
killProgress()
```

**Arguments**

- `title` The title of the tcltk window
- `message` A short test message to add to the window
- `sub` An additional text field that can be updated via updateProgress
- `percentage` An integer giving the percentage of completion
- `autoKill` Logical indicating whether to kill the display after 100 is reached

**Details**

Function `progress` creates the progress window and sets up the necessary environment. `updateProgress` takes as argument an integer value indicating the percentage of completion and updates the display. The integer value that gets passed to `updateProgress` will usually be generated by an iterator (e.g. in a for loop). `killProgress` may be called explicitly to kill the progress window. Alternatively, one can set the argument `autoKill` of `updateProgress` to `TRUE` to automatically kill the window once a value of 100 is reached.

**Value**

The functions are called for their side effects.

**Author(s)**

Florian Hahne

**Examples**

```r
if(interactive() && capabilities()["tcltk"]) {
  progress(message="This is a progress display...", sub="(step 1 of 50)")
  for(i in 1:50) {
    zz = rnorm(1e5)
    updateProgress(i*2, autoKill=TRUE, sub=paste("(step", i, " of 50)"))
  }
}
```
readCytoSet  

Create a cytoSet object from one or more FCS 3.0 files

Description
Create a cytoSet object from one or more FCS 3.0 files

Usage
readCytoSet(files=NULL, path=".", pattern=NULL, phenoData, sep="\t", ...)

Arguments
- files: Optional character vector with filenames
- path: Directory where to look for the files
- pattern: This argument is passed on to `dir` (see details).
- phenoData: Either an object of class `phenoData` or character.
- sep: Separator character that gets passed on to `read.AnnotatedDataFrame`.
- ...: Further arguments that get passed on to `read.AnnotatedDataFrame`, see details.

Details
There are three different ways to specify the file names:
First, if the argument `phenoData` is present and is of class `AnnotatedDataFrame`, then it is obtained from its column name. The column is mandatory, and an error will be generated if it is not there. Alternatively, the argument `phenoData` can be of class character, in which case this function tries to read a `AnnotatedDataFrame` object from the file with that name by calling `read.AnnotatedDataFrame` with arguments `file.path(path, phenoData), ...`.

Second, if the argument `phenoData` is not present and the argument `files` is not `NULL`, then `files` is expected to be a character vector with the file names.

Third, if neither the argument `phenoData` is present nor `files` is not `NULL`, then the file names are obtained by calling `dir(path, pattern)`.

Value
An object of class `cytoSet`.

Author(s)
Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber)

See Also
- readFCSdata

Examples
## Please see man page for cytoSet-class
Description

Read one or several FCS files: Data File Standard for Flow Cytometry

Usage

```r
read.fcs(filename=NULL, objectModel="prada", ...)
readFCS(filename)
```

Arguments

- `filename`: Character of length 1: filename
- `objectModel`: Character of length 1: the object model to use for the output. Currently only ’prada’ for cytoFrame objects is supported.
- `...`: Arguments that get passed on to higher-level import functions.

Details

The function `readFCS` works with the output of the FACS machine software from a number of vendors. However, the FCS 3.0 standard includes some options that are not yet implemented in this function. If you need extensions, please let me know. The output of the function is an object of class `cytoFrame`.

`read.fcs` is a wrapper function that allows the user to specify the class of the output. The purpose of the function is to standardize the way flow cytometry data is imported into R using the prada package. If the `filename` argument to `read.fcs` is a character vector of length > 1, multiple FCS files can be imported. Please see the documentation for `readCytoSet` for alternatives ways to import multiple FCS files and for more details on the higher-level import function.

For specifications of FCS 3.0 see [http://www.isac-net.org](http://www.isac-net.org) and the file `../doc/fcs3.html` in the doc directory of the package.

Value

An object of class `cytoFrame`.

Author(s)

Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber), Florian Hahne

See Also

`readCytoSet`
Examples

```r
sampdat <- readFCS(system.file("extdata", "fas-Bcl2-plate323-04-04.A01", package="prada"))
files <- dir(system.file("extdata", package="prada"), pattern="[A-H][0-9][0-9]")
sampdat2 <- read.fcs(system.file("extdata", "fas-Bcl2-plate323-04-04.A01", package="prada"))
sampdat3 <- read.fcs(files, path=system.file("extdata", package="prada"))
sampdat
description(sampdat[[1:3,]])
class(sampdat)
```

---

**readFCSaux**

Auxiliary functions for readFCS

**Description**

Auxiliary functions for readFCS - not normally called by the user

**Usage**

```r
readFCSgetPar(x, pnam)
readFCSheader(con)
readFCStext(con, offsets)
readFCSdata(con, offsets, x)
```

**Arguments**

- `x` Named character vector.
- `pnam` Character vector, its elements must be contained in `names(x)`.
- `con` Connection.
- `offsets` Integer vector of length 6 with byte offsets of the header, text, and data blocks.

**Details**

These functions are not normally called by the user. See `readFCS` instead.

**Value**

Various.

**Author(s)**

Wolfgang Huber [http://www.ebi.ac.uk/huber](http://www.ebi.ac.uk/huber)

**See Also**

`readFCS`
removeCensored

### Description

Remove rows that contain censored data in the columns of x specified by columns.

### Usage

```r
## S4 method for signature 'matrix'
removeCensored(x, values, columns, na.rm=TRUE)
## S4 method for signature 'data.frame'
removeCensored(x, values, columns, na.rm=TRUE)
## S4 method for signature 'cytoFrame'
removeCensored(x, values, columns, na.rm=TRUE)
```

### Arguments

- **x**: Object of class matrix, data.frame, or cytoFrame.
- **values**: Values that correspond to censored data. If missing, `range(x)` is used.
- **columns**: Numeric or character vector specifying the columns of x that are compared against values. If missing, `1:ncol(x)` is used.
- **na.rm**: Logical. If TRUE, rows that contain NA values are also removed.

### Details

The function removes all rows that contain, in the columns specified by the columns argument, values that are contained in the values argument. If na.rm is TRUE, then rows that contain NA values are also removed.

An application is with FACS data, where measurements outside of the detector’s dynamic range produce minimal or maximal values. For example, if a 16-bit A/D converter was used, top-censored data would have a value of 65535.

### Value

Object of the same class as x, with some rows removed.

### Author(s)

Florian Hahne, Wolfgang Huber

### Examples

```r
set.seed(8215)
mat <- matrix(floor(runif(20000)*1024), ncol=4)
range(mat[,1])
mat <- removeCensored(mat, columns=1:2)
range(mat[,1])
range(mat[,3])
```
threePanelPlot

Visualize cytometry data

Description

Function to visualize multivariate (cytometry) data in three two-dimensional plots.

Usage

threePanelPlot(data, x.panels = c(1, 4, 5), y.panels = c(2, 3, 6),
     tot.width = 15, tot.height = 5.4, maxcells = 20000,
     limits = c(0, 1023), remove.extremes = TRUE,
     plotTitle = "Three-Panel Plot", use.smoothScatter = TRUE,
     palette = colorRampPalette(brewer.pal(9, "Blues")),
     new.device = TRUE, verbose = TRUE,
     addPoints = NULL, addCol = "red", ...)

Arguments

data data matrix to visualize
x.panels which variables (columns) are to be plotted at the x-axis of the three variables
y.panels which variables (columns) are to be plotted at the y-axis of the three variables
tot.width width of a new device to open, see argument new.device
tot.height height of a new device to open, see argument new.device
maxcells maximum number of observations (cells) for plotting; higher numbers reduce performance
limits minimum and maximum value (theoretically) observed in the data; e.g., with 10-channel digitized data it is c(0,1023)
remove.extremes logical; are extreme values (equal to theoretical limits) to be removed before plotting
plotTitle title for the plot
use.smoothScatter logical, should the function smoothScatter be employed for plotting the data (plots data densities rather than individual points)
palette if smoothScatter is used, which colour palette is it to use
new.device logical; should a new device be opened for the three plots; if FALSE the three plots will be plotted to the currently active device
verbose logical; do you want extended output to STDOUT
addPoints should special points be marked after plotting the data; is expected to be a subser of argument data with the same number of columns (=variables); if NULL no points are marked
addCol in which colour are the points in addPoints to be marked
...
further arguments passed on to plot.default

Value

no value is returned; the function is called to produce three plots
thresholds

**Author(s)**
Joern Toedling <toedling@ebi.ac.uk>

**See Also**
`plot.default`

**Examples**
```r
# generate some data:
toyData <- cbind(matrix(pmax(0,pmin(runif(3000)+rnorm(3000),4)),ncol=3),
                 matrix(pmax(0,pmin(rnorm(3000,2,1),4)),ncol=3))
colnames(toyData) <- paste("Var",1:6,sep="")
toyQuantiles <- apply(toyData,2,quantile,probs=c(0.25,0.5,0.75))

# plot it and mark the quantiles:
threePanelPlot(toyData,addPoints=toyQuantiles,
               addCol=c("orange","red","purple"),limits=c(0,4),pch=20)
```

**thresholds**

Discretize a two-dimensional data space into quadrants by applying thresholds.

**Usage**

```r
thresholds(x, y, xthr, ythr)
```

**Arguments**

- `x`: Vector containing x or matrix containing x and y values of bivariate data.
- `y`: Optional vector containing y values of bivariate data.
- `xthr`: x value separating 'left' and 'right'.
- `ythr`: y value separating 'up' and 'down'.

**Details**

The function returns a 2x2 matrix giving the counts for each quadrant. Events with values equal to the thresholds are counted to the left or down respectively.

**Value**

2x2 matrix.

**Author(s)**
Florian Hahne

**Examples**

```r
thresholds(cbind(c(1, 1, 2, 2, 2, 4), c(1, 4, 2, 4, 5, 4)), xthr=3, ythr=3)
```
touchFCS  
*Check for FCS files*

**Description**

The function reads the header of a file or of a range of files and checks whether they are valid FCS 2.0 or FCS 3.0 files.

**Usage**

```r
touchFCS(path = ".", file)
```

**Arguments**

- `path` character, the path to a folder containing files
- `file` character, the path to a single file

**Details**

The user may either specify the path to a directory in which to search for FCS files or the path to a single file.

**Value**

A character vector with names of the valid FCS files found.

**Author(s)**

fhahne

---

vpLocation  
*Absolute location of current viewport*

**Description**

Calculates the absolute location and size of the current grid viewport in inches and pixels.

**Usage**

```r
vpLocation()
```

**Details**

This function may be used to get the absolute location of the current viewport on the current graphics device. It uses function `devRes` to get the device resolution for calculating pixel values. Locations are given by the two extreme coordinates in x and y direction.

**Value**

A list with items `location`, `size`, `ilocation` and `isize`, the location and size of the viewport in pixels and inches respectively.
Author(s)
    Florian Hahne

See Also
    plotPlate, devRes

Examples
    vpLocation()
Index

*Topic IO
  readCytoSet, 22
  readFCS, 23
  readFCSaux, 24
  touchFCS, 28

*Topic classes
  cytoFrame-class, 7
  cytoSet-class, 9
  gate-class, 13
  gateSet-class, 15

*Topic datasets
  cframe, 5
  cset, 7

*Topic hplot
  barploterrbar, 4
  plotPlate, 19
  threePanelPlot, 26

*Topic manip
  analysePlate, 2
  as.all, 3
  csApply, 6
  getPradaPar, 17

*Topic misc
  progress, 21
  [, cytoFrame, ANY, ANY, ANY-method (cytoFrame-class), 7
  [, cytoSet, ANY, missing, missing-method (cytoSet-class), 9
  [, gateSet, ANY, missing, missing-method (gateSet-class), 15
  [[, cytoSet, ANY, missing-method (cytoSet-class), 9
  [[, gateSet, ANY, missing-method (gateSet-class), 15
  [[<-, cytoSet-method (cytoSet-class), 9
  $ . cytoFrame (cytoFrame-class), 7
  analysePlate, 2
  AnnotatedDataFrame, 22
  appendGates (gateSet-class), 15
  appendGates, gateSet-method (gateSet-class), 15
  appendGates, cytoFrame-method (cytoFrame-class), 7
  appendGates, gateSet-method (gate-class), 13
  applyGate (gateSet-class), 15
  applyGate, cytoFrame, character-method (cytoFrame-class), 7
  applyGate, cytoFrame, gate-method (cytoFrame-class), 7
  applyGate, cytoFrame, gateSet-method (cytoFrame-class), 7
  applyGate, cytoFrame, logical-method (cytoFrame-class), 7
  applyGate, cytoFrame, numeric-method (cytoFrame-class), 7
  applyGate, matrix, gate-method (gate-class), 13
  applyGate, matrix, gateSet-method (gateSet-class), 15
  as, 3
  as.all, 3
  as.gateSet (gate-class), 13
  as.gateSet, gate-method (gate-class), 13
  barplot, 4
  barploterrbar, 4
  cframe, 5
  colnames, cytoFrame-method (cytoFrame-class), 7
  colnames, cytoSet-method (cytoSet-class), 9
  colnames<-, cytoFrame-method (cytoFrame-class), 7
  colnames<-, cytoSet-method (cytoSet-class), 9
  colorRampPalette, 19
  combineFrames, 5
  combineGates (gate-class), 13
  convertWellCoordinates, 17
  cov.rob, 13
  csApply, 6
  cset, 7
  cytoFrame, 9, 10, 14–16, 23
  cytoFrame (cytoFrame-class), 7
  cytoFrame-class, 7
split, cytoSet, ANY-method (cytoSet-class), 9
split, cytoSet-method (cytoSet-class), 9
tapply, 3
threePanelPlot, 26
thresholds, 27
touchFCS, 28
updateProgress (progress), 21
vpLocation, 28