Package 'plotROC'

October 6, 2023

Type Package
Title Generate Useful ROC Curve Charts for Print and Interactive Use
Version 2.3.1
Date 2023-10-05
Maintainer Michael C. Sachs <sachsmc@gmail.com>
Description Most ROC curve plots obscure the cutoff values and inhibit interpretation and comparison of multiple curves. This attempts to address those shortcomings by providing plotting and interactive tools. Functions are provided to generate an interactive ROC curve plot for web use, and print versions. A Shiny application implementing the functions is also included.
License MIT + file LICENSE

LICENSE MIT + IIIE LICENSE

URL https://sachsmc.github.io/plotROC/

BugReports https://github.com/sachsmc/plotROC/issues/

Depends R (>= 3.0.0), ggplot2

Imports methods, grid, gridSVG, shiny, plyr, rlang

Suggests knitr, testthat, stringr, survivalROC, rmarkdown

VignetteBuilder knitr

RoxygenNote 7.2.3

NeedsCompilation no

Author Michael C. Sachs [aut, cre], Robert W. Corty [ctb], Luis Crouch [ctb] (Modification to calc_auc)

Repository CRAN

Date/Publication 2023-10-06 12:40:02 UTC

R topics documented:

calculate_multi_roc	2
calculate_roc	3
calc_auc	3
direct_label	4
export_interactive_roc	5
GeomRoc	6
geom_rocci	9
getD3	11
ggroc	11
melt_roc	12
multi_ggroc	13
plotROC	14
plot_interactive_roc	15
plot_journal_roc	16
roc_key	17
shiny_plotROC	17
StatRoc	
	19
style_roc	
verify_d	23
	24

calculate_multi_roc Calculate the Empirical ROC curves for multiple biomarkers

Description

Deprecated, use geom_roc instead

Usage

Index

```
calculate_multi_roc(data, M_string, D_string)
```

Arguments

data	data frame containing at least 1 marker and the common class labels, coded as 0 and 1 $$
M_string	vector of marker column names
D_string	class label column name

Value

List of data frames containing cutoffs, and estimated true and false positive fractions

calculate_roc

Description

Deprecated, use geom_roc instead

Usage

calculate_roc(M, D, ci = FALSE, alpha = 0.05)

Arguments

М	continuous marker values or predictions of class labels
D	class labels, must be coded as 0 and 1. If not numeric with 0/1, then plotROC assumes the first level in sort order is healthy status, with a warning.
ci	Logical, if true, will calculate exact joint confidence regions for the TPF and FPF
alpha	Confidence level, ignored if ci = FALSE

Details

Confidence intervals for TPF and FPF are calculated using the exact method of Clopper and Pearson (1934) each at the level 1 - sqrt(1 - alpha). Based on result 2.4 from Pepe (2003), the cross-product of these intervals yields a 1 - alpha

Value

A dataframe containing cutoffs, estimated true and false positive fractions, and confidence intervals if ci = TRUE.

calc_auc

Calculate the Area under the ROC curve

Description

Given a ggplot object with a GeomRoc layer, computes the area under the ROC curve for each group

Usage

```
calc_auc(ggroc)
```

Arguments

ggroc

A ggplot object that contains a GeomRoc layer

Value

A data frame with the estimated AUCs for each panel and group

Examples

direct_label Add direct labels to a ROC plot

Description

Add direct labels to a ROC plot

Usage

```
direct_label(
  ggroc_p,
  labels = NULL,
  label.angle = 45,
  nudge_x = 0,
  nudge_y = 0,
  size = 6,
  ...
)
```

Arguments

ggroc_p	A ggplot object that contains a geom_roc layer	
labels,	vector of labels to add directly to the plot next to the curves. If multiple curves, must be in the same order as the grouping factor. If NULL, attempts to determine labels from the ggroc_p object	
label.angle	angle of adjustment for the direct labels	
nudge_x, nudge_y		
	Horizontal and vertical adjustment to nudge labels by. These can be scalars or vectors the same length as the number of labels	
size	Size of labels	
	Other arguments passed to annotate	

export_interactive_roc

Generate svg code for an ROC curve object

Description

Takes a ggplot object that contains a GeomRoc layer and returns a string that contains html suitable for creating a standalone interactive ROC curve plot.

Usage

```
export_interactive_roc(
  ggroc_p,
  add.cis = TRUE,
  hide.points = FALSE,
  prefix = "a",
  width = 6,
  height = 6,
  omit.js = FALSE,
  style = style_roc(theme = theme_grey()),
  ...
)
```

Arguments

ggroc_p	A ggplot object with a GeomRoc layer and optionally a GeomRocci layer as returned by geom_roc and/or geom_rocci. It can be modified with annotations, themes, etc.
add.cis	Logical, if true, removes the current confidence interval layer (if present) and replaces it with a denser layer of confidence regions
hide.points	Logical, if true, hides points layer so that points with cutoff values are only visible when hovering. Recommended for plots containing more than 3 curves.
prefix	A string to assign to the objects within the svg. Enables unique identification by the javascript code
width, height	Width and height in inches of plot
omit.js	Logical. If true, omit inclusion of javascript source in output. Useful for documents with multiple interactive plots
style	A call to the function style_roc
	Other arguments passed to geom_rocci when add.cis = TRUE

Details

If you intend to include more than one of these objects in a single page, use a different prefix string for each one. To use this function in knitr, use the chunk options fig.keep='none' and results = 'asis', then cat() the resulting string to the output. See the vignette for examples. Older browsers (< IE7) are not supported.

Value

A character object containing the html necessary to plot the ROC curve in a web browser

GeomRoc

Empirical Receiver Operating Characteristic Curve

Description

Display the empirical ROC curve. Useful for characterizing the classification accuracy of continuous measurements for predicting binary states

Usage

GeomRoc

```
geom_roc(
 mapping = NULL,
 data = NULL,
 stat = "roc",
  n.cuts = 10,
  arrow = NULL,
  lineend = "butt"
  linejoin = "round",
  linemitre = 1,
  linealpha = 1,
  pointalpha = 1,
  pointsize = 0.5,
  labels = TRUE,
  labelsize = 3.88,
  labelround = 1,
 na.rm = TRUE,
  cutoffs.at = NULL,
  cutoff.labels = NULL,
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  . . .
)
```

Arguments

mapping

Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.

GeomRoc

data	The data to be displayed in this layer. There are three options: If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot().
	A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.
	A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. \sim head(.x, 10)).
stat	Use to override the default connection between geom_roc and stat_roc.
n.cuts	Number of cutpoints to display along each curve
arrow	Arrow specification, as created by arrow
lineend	Line end style (round, butt, square)
linejoin	Line join style (round, mitre, bevel)
linemitre	Line mitre limit (number greater than 1)
linealpha	Alpha level for the lines, alpha.line is deprecated
pointalpha	Alpha level for the cutoff points, alpha.point is deprecated
pointsize	Size of cutoff points, size.point is deprecated
labels	Logical, display cutoff text labels
labelsize	Size of cutoff text labels
labelround	Integer, number of significant digits to round cutoff labels
na.rm	Remove missing values from curve
cutoffs.at	Vector of user supplied cutoffs to plot as points. If non-NULL, it will override the values of n.cuts and plot the observed cutoffs closest to the user-supplied ones.
cutoff.labels	vector of user-supplied labels for the cutoffs. Must be a character vector of the same length as cutoffs.at.
position	Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().
	Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3. They may also be parameters to the paired geom/stat.

Format

An object of class GeomRoc (inherits from Geom, ggproto, gg) of length 6.

GeomRoc

Computed variables

false_positive_fraction estimate of false positive fraction

true_positive_fraction estimate of true positive fraction

cutoffs values of m at which estimates are calculated

Aesthetics

geom_roc understands the following aesthetics (required aesthetics are in bold):

- x The FPF estimate. This is automatically mapped by stat_roc
- y The TPF estimate. This is automatically mapped by stat_roc smallest level in sort order is assumed to be 0, with a warning
- alpha
- color
- fill
- linetype
- size

See Also

See geom_rocci for displaying rectangular confidence regions for the empirical ROC curve, style_roc for adding guidelines and labels, and direct_label for adding direct labels to the curves. Also export_interactive_roc for creating interactive ROC curve plots for use in a web browser.

Examples

geom_rocci

Description

Display rectangular confidence regions for the empirical ROC curve.

Usage

```
geom_rocci(
 mapping = NULL,
 data = NULL,
 stat = "rocci",
 ci.at = NULL,
  sig.level = 0.05,
  na.rm = TRUE,
  alpha.box = 0.3,
 labels = TRUE,
 labelsize = 3.88,
  labelround = 1,
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  . . .
)
```

GeomRocci

Arguments

mapping	Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot().
	A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.
	A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. \sim head(.x, 10)).
stat	Use to override the default connection between geom_rocci and stat_rocci.
ci.at	Vector of values in the range of the biomarker where confidence regions will be displayed

sig.level	Significance level for the confidence regions
na.rm	If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.
alpha.box	Alpha level for the confidence regions
labels	If TRUE, adds text labels for the cutoffs where the confidence regions are dis- played
labelsize	Size of cutoff text labels
labelround	Integer, number of significant digits to round cutoff labels
position	Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().
	Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3. They may also be parameters to the paired geom/stat.

Format

An object of class GeomRocci (inherits from Geom, ggproto, gg) of length 6.

Aesthetics

geom_rocci understands the following aesthetics (required aesthetics are in bold). stat_rocci automatically maps the estimates to the required aesthetics:

- x The FPF estimate
- y The TPF estimate
- xmin Lower confidence limit for the FPF
- xmax Upper confidence limit for the FPF
- ymin Lower confidence limit for the TPF
- ymax Upper confidence limit for the TPF
- alpha
- color
- fill
- linetype
- size

getD3

See Also

See geom_roc for the empirical ROC curve, style_roc for adding guidelines and labels, and direct_label for adding direct labels to the curves. Also export_interactive_roc for creating interactive ROC curve plots for use in a web browser.

Examples

getD3	Reads included JavaScript functions and returns them as a string for
	pasting into a webpage

Description

Reads included JavaScript functions and returns them as a string for pasting into a webpage

Usage

getD3()

ggroc

Plot an ROC curve

Description

Deprecated, use geom_roc instead

Usage

```
ggroc(
  rocdata,
  fpf_string = "FPF",
  tpf_string = "TPF",
  c_string = "c",
  ci = FALSE,
  label = NULL,
  label.adj.x = 0,
  label.adj.y = 0,
  label.angle = 45,
  plotmath = FALSE,
  xlabel = "False positive fraction",
  ylabel = "True positive fraction"
)
```

Arguments

rocdata	Data frame containing true and false positive fractions, and cutoff values
fpf_string	Column name identifying false positive fraction column
tpf_string	Column name identifying true positive fraction column
c_string	Column name identifying cutoff values
ci	Logical, not supported
label	Not supported
label.adj.x	Not supported
label.adj.y	Not supported
label.angle	Not supported
plotmath	Not supported
xlabel	Defaults to "False positive fraction"
ylabel	Defaults to "True positive fraction"

Value

A ggplot object

melt_roc

Transform biomarkers stored as wide to long

Description

Multiple biomarkers measured on the same subjects are often stored as multiple columns in a data frame. This is a convenience function that transforms the data into long format, suitable for use with ggplot and geom_roc

multi_ggroc

Usage

melt_roc(data, d, m, names = NULL)

Arguments

data	Data frame containing disease status and biomarkers stored in columns
d	Column containing binary disease status. Can be a column name or index
m	Vector of column names or indices identifying biomarkers
names	Optional vector of names to assign to the biomarkers. If NULL, names will be
	taken from the column names

Value

A data frame in long format with three columns: D = binary disease status, M = biomarker value, and name = biomarker name

Examples

```
D.ex <- rbinom(50, 1, .5)
widedata <- data.frame(D = D.ex, M1 = rnorm(50, mean = D.ex, sd = 1),
    M2 = rnorm(50, mean = D.ex, sd = .5))
longdata <- melt_roc(widedata, "D", c("M1", "M2"))
ggplot(longdata, aes(d = D, m = M, color = name)) + geom_roc()</pre>
```

multi_ggroc	Plot multiple ROC curves

Description

Given a list of results computed by calculate_roc, plot the curve using ggplot with sensible defaults. Pass the resulting object and data to export_interactive_roc, plot_interactive_roc, or plot_journal_roc.

Usage

```
multi_ggroc(
    datalist,
    fpf_string = rep("FPF", length(datalist)),
    tpf_string = rep("TPF", length(datalist)),
    c_string = rep("c", length(datalist)),
    label = NULL,
    legend = TRUE,
    label.adj.x = rep(0, length(datalist)),
    label.adj.y = rep(0, length(datalist)),
    label.angle = rep(45, length(datalist)),
    plotmath = FALSE,
    xlabel = "False positive fraction",
    ylabel = "True positive fraction"
```

Arguments

datalist	List of data frames each containing true and false positive fractions and cutoffs
fpf_string	Column names identifying false positive fraction
tpf_string	Column names identifying true positive fraction
c_string	Column names identifying cutoff values
label	Not supported.
legend	If true, draws legend
label.adj.x	Not supported.
label.adj.y	Not supported.
label.angle	Not supported.
plotmath	Logical. Not supported.
xlabel	Defaults to "False positive fraction"
ylabel	Defaults to "True positive fraction"

Value

A ggplot object

plotROC

Tools for plotting ROC Curves

Description

Generate Useful ROC Curve Charts for Print and Interactive Use. This defines a set of stats and geoms for use with ggplot2. In addition, ggplot objects created with these geoms can be exported and turned into interactive plots for use on the web. The interactive features include hovering to display hidden labels, and clicking to reveal confidence regions.

Details

To get started, see geom_roc, geom_rocci, or the examples below. For transforming data, we also provide the convenience function melt_roc.

The vignette contains some examples, along with explanations of the results. To view, run vignette("examples", package = "plotROC")

Author(s)

Michael Sachs (@sachsmc)

See Also

Useful links:

- https://sachsmc.github.io/plotROC/
- Report bugs at https://github.com/sachsmc/plotROC/issues/

plot_interactive_roc

Examples

<pre>plot_interactive_roc</pre>	Generate a standalone html document displaying an interactive ROC
	curve

Description

Generate a standalone html document displaying an interactive ROC curve

Usage

```
plot_interactive_roc(ggroc, file = NULL, ...)
```

Arguments

ggroc	An object as returned by ggroc or multi_ggroc. It can be modified with annota- tions, themes, etc.
file	A path to save the result to. If NULL, will save to a temporary directory
	arguments passed to export_interactive_roc

Value

NULL opens an interactive document in Rstudio or the default web browser

plot_journal_roc *Plot an ROC curve for use in print*

Description

Deprecated, use style_roc instead

Usage

```
plot_journal_roc(
  ggroc_p,
  font.size = 3,
  n.cuts = 20,
  ci.at = NULL,
  opacity = 0.3,
  lty = NULL,
  color = NULL,
  lwd = NULL,
  legend = FALSE
)
```

Arguments

ggroc_p	An object as returned by ggroc or multi_ggroc. It can be modified with annota- tions, themes, etc.
font.size	Not supported
n.cuts	Not supported
ci.at	Not supported
opacity	Not supported
lty	Not supported
color	Not supported
lwd	Not supported
legend	Not supported

Value

A ggplot object

roc_key

Description

Key for ROC geom

Usage

roc_key(data, params, size)

Arguments

data	Data created by stat
params	parameters
size	Size

shiny_plotROC Start the plotROC Shiny app

Description

A convenience function to easily start the shiny application. It will open in Rstudio, or in the default web browser.

Usage

shiny_plotROC()

StatRoc

Calculate the empirical Receiver Operating Characteristic curve

Description

Given a binary outcome d and continuous measurement m, computes the empirical ROC curve for assessing the classification accuracy of m

Usage

StatRoc

```
stat_roc(
  mapping = NULL,
  data = NULL,
  geom = "roc",
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  na.rm = TRUE,
  max.num.points = 1000,
  increasing = TRUE,
  ...
)
```

Arguments

mapping	Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options:
	If NULL, the default, the data is inherited from the plot data as specified in the call to $ggplot()$.
	A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.
	A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. \sim head(.x, 10)).
geom	The geometric object to use to display the data, either as a ggproto Geom sub- class or as a string naming the geom stripped of the geom_ prefix (e.g. "point" rather than "geom_point")
position	Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().
na.rm	Remove missing observations
max.num.points	maximum number of points to plot
increasing	TRUE (default) if M is positively associated with $Pr(D = 1)$, if FALSE, assumes M is negatively associated with $Pr(D = 1)$

StatRocci

... Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3. They may also be parameters to the paired geom/stat.

Format

An object of class StatRoc (inherits from Stat, ggproto, gg) of length 6.

Aesthetics

stat_roc understands the following aesthetics (required aesthetics are in bold):

- m The continuous biomarker/predictor
- d The binary outcome, if not coded as 0/1, the smallest level in sort order is assumed to be 0, with a warning
- alpha Controls the label alpha, see also linealpha and pointalpha
- color
- linetype
- size Controls the line weight, see also pointsize and labelsize

Computed variables

false_positive_fraction estimate of false positive fraction

true_positive_fraction estimate of true positive fraction

cutoffs values of m at which estimates are calculated

Examples

ggplot(rocdata, aes(m = M, d = D)) + stat_roc()

StatRocci

Calculate confidence regions for the empirical ROC curve

Description

Confidence intervals for TPF and FPF are calculated using the exact method of Clopper and Pearson (1934) each at the level 1 - sqrt(1 - alpha). Based on result 2.4 from Pepe (2003), the cross-product of these intervals yields a 1 - alpha

Usage

StatRocci

```
stat_rocci(
  mapping = NULL,
  data = NULL,
  geom = "rocci",
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  ci.at = NULL,
  sig.level = 0.05,
  na.rm = TRUE,
  ...
)
```

Arguments

mapping	Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options:
	If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot().
	A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.
	A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. \sim head(.x, 10)).
geom	The geometric object to use to display the data, either as a ggproto Geom sub- class or as a string naming the geom stripped of the geom_prefix (e.g. "point" rather than "geom_point")
position	Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().
ci.at	Vector of cutoffs at which to display confidence regions. If NULL, will auto- matically choose 3 evenly spaced points to display the regions
sig.level	Significance level for the confidence regions

StatRocci

na.rm	Remove missing observations
	Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3. They may also be parameters to the paired geom/stat.

Format

An object of class StatRocci (inherits from Stat, ggproto, gg) of length 6.

Aesthetics

stat_rocci understands the following aesthetics (required aesthetics are in bold):

- m The continuous biomarker/predictor
- d The binary outcome, if not coded as 0/1, the smallest level in sort order is assumed to be 0, with a warning
- alpha
- color
- fill
- linetype
- size

Computed variables

- FPF estimate of false positive fraction
- TPF estimate of true positive fraction

cutoffs values of m at which estimates are calculated

- FPFL lower bound of confidence region for FPF
- FPFU upper bound of confidence region for FPF
- TPFL lower bound of confidence region for TPF
- TPFU upper bound of confidence region for TPF

References

- Clopper, C. J., and Egon S. Pearson. "The use of confidence or fiducial limits illustrated in the case of the binomial." Biometrika (1934): 404-413.
- Pepe, M.S. "The Statistical Evaluation of Medical Tests for Classification and Prediction." Oxford (2003).

Examples

style_roc Add guides and annotations to a ROC p	style_roc	Add guides and annotations to a ROC plot
---	-----------	--

Description

Adds a diagonal guideline, minor grid lines, and optionally direct labels to ggplot objects containing a geom_roc layer.

Usage

```
style_roc(
  major.breaks = c(0, 0.1, 0.25, 0.5, 0.75, 0.9, 1),
  minor.breaks = c(seq(0, 0.1, by = 0.01), seq(0.9, 1, by = 0.01)),
  guide = TRUE,
  xlab = "False positive fraction",
  ylab = "True positive fraction",
  theme = theme_bw
)
```

Arguments

major.breaks	vector of breakpoints for the major gridlines and axes
minor.breaks	vector of breakpoints for the minor gridlines and axes
guide	logical, if TRUE draws diagonal guideline
xlab	X-axis label
ylab	Y-axis label
theme	Theme function compatible with ggplot2

Examples

```
D.ex <- rbinom(50, 1, .5)
fakedata <- data.frame(M1 = rnorm(50, mean = D.ex),
    D = D.ex)
ggplot(fakedata, aes(m = M1, d = D)) + geom_roc() + style_roc()
ggplot(fakedata, aes(m = M1, d = D)) + geom_roc() + style_roc(xlab = "1 - Specificity")
ggplot(fakedata, aes(m = M1, d = D)) + geom_roc() + style_roc(theme = theme_grey)</pre>
```

verify_d

Description

Checks for two classes and gives a warning message indicating which level is assumed to be 0/1. Throws an error if more than two levels appear in D.

Usage

verify_d(D)

Arguments

D

Vector that will be checked for 2-class labels

Value

A vector the same length as D that takes values 0, indicating no disease or 1 indicating disease.

Examples

```
verify_d(c(1, 0, 1))
## Not run:
verify_d(c(TRUE, FALSE, TRUE)) #warning
verify_d(c("Dead", "Alive", "Dead")) #warning
verify_d(c("Disease", "Healthy", "Missing")) #error
```

End(Not run)

Index

* Internal getD3, 11 roc_key, 17 * datasets geom_rocci,9 GeomRoc, 6 StatRoc, 17 StatRocci, 19 aes(), 6, 9, 18, 20 annotate, 4 arrow, 7 borders(), 7, 10, 18, 20 calc_auc, 3 calculate_multi_roc, 2 calculate_roc, 3, 13 direct_label, 4, 8, 11 export_interactive_roc, 5, 8, 11, 13, 15 fortify(), 7, 9, 18, 20 geom_roc, 2-5, 11, 12, 14 geom_roc (GeomRoc), 6 geom_rocci, 5, 8, 9, 14 GeomRoc, 6 GeomRocci (geom_rocci), 9 getD3, 11 ggplot(), 7, 9, 18, 20 ggroc, 11, 15, 16 layer(), 7, 10, 19, 21 melt_roc, 12, 14 multi_ggroc, 13, 15, 16 plot_interactive_roc, 13, 15 plot_journal_roc, 13, 16

plotROC, 14 plotROC-package (plotROC), 14 roc_key, 17 shiny_plotROC, 17 stat_roc, 8 stat_roc (StatRoc), 17 stat_rocci (StatRocci), 19 StatRoc, 17 StatRocci, 19 style_roc, 5, 8, 11, 16, 22 verify_d, 23