Package ‘gplots’

January 27, 2019

Title  Various R Programming Tools for Plotting Data

Description Various R programming tools for plotting data, including:
- calculating and plotting locally smoothed summary function as
  (`bandplot', 'wapply'),
- enhanced versions of standard plots (`barplot2', 'boxplot2',
  'heatmap.2', 'smartlegend'),
- manipulating colors (`col2hex', 'colorpanel', 'redgreen',
  'greenred', 'bluered', 'redblue', 'rich.colors'),
- calculating and plotting two-dimensional data summaries (`ci2d',
  'hist2d'),
- enhanced regression diagnostic plots (`lmplot2', 'residplot'),
- formula-enabled interface to 'stats::lowess' function ('lowess'),
- displaying textual data in plots ('textplot', 'sinkplot'),
- plotting a matrix where each cell contains a dot whose size
  reflects the relative magnitude of the elements ('balloonplot'),
- plotting ``Venn'' diagrams ('venn'),
- displaying Open-Office style plots ('ooplot'),
- plotting multiple data on same region, with separate axes
  ('overplot'),
- plotting means and confidence intervals ('plotCI', 'plotmeans'),
- spacing points in an x-y plot so they don't overlap ('space').

Depends  R (>= 3.0)
Imports  gtools, gdata, stats, caTools, KernSmooth
Suggests  grid, MASS
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License  GPL-2
NeedsCompilation  no
angleAxis

Add a Axis to a Plot with Rotated Labels

Description
Add a labeled axis to the current plot with rotated text

Usage
angleAxis(side, labels, at = 1:length(labels), srt = 45, adj, xpd = TRUE, ...)
angleAxis

Arguments

side: an integer specifying which side of the plot the axis is to be drawn on. The axis
is placed as follows: 1=below, 2=left, 3=above and 4=right.

labels: character or expression vector of labels to be placed at the tickpoints.

at: the points at which tick-marks are to be drawn. Non-finite (infinite, NaN or NA)
values are omitted.

srt: The string rotation in degrees. Defaults to 45 degrees (clockwise).

adj: Text justification. A value of 0 produces left-justified text, 0.5 centered text and
1 right-justified text. For side=1 and side=2, the default value is adj=1. For
side=3 and side=4 the default value is adj=0.

xpd: A logical value or NA. If FALSE, labels are clipped to the plot region, if TRUE,
labels are clipped to the figure region, and if NA, labels are clipped to the device
region.

... optional arguments passed to text. Common examples are col, cex.

Details

This function augments the feature of the axis function by allowing the axis labels to be rotated.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

axis

Examples

# create a vector with some values and long labels
values <- sample(1:10)
names(values) <- sapply(letters[1:10],
function(x) paste(rep(x, 10), sep="", collapse="")
)

# barplot labels are too long for the available space, hence some are not plotted
barplot(values)

# to add angled labels, tell barplot not to label the x axis, and store the bar location
at <- barplot(values, xaxt="n")
# then use angleAxis
angleAxis(1, at=at, labels = names(values))

# angle counter-clockwise instead
at <- barplot(values, xaxt="n")
angleAxis(1, at=at, labels = names(values), srt=-45, adj=0)

# put labels at the top
oldpar <- par$mar
par(mar=c(1,4.5,2)+0.1)
at <- barplot(values, xaxt="n")
ageAxis(3, at=at, labels = names(values))
par(oldpar)

# put labels on the left
oldpar <- par$mar
par(mar=c(5,5.3,2)+0.1)
at <- barplot(values, yaxt="n", horiz=TRUE)
ageAxis(2, at=at, labels = names(values))
par(oldpar)

# put labels on the right
oldpar <- par$mar
par(mar=c(2,5.3,5)+0.1)
at <- barplot(values, yaxt="n", horiz=TRUE)
ageAxis(4, at=at, labels = names(values))
par(oldpar)

# specify colors for bars and labels
at <- barplot(values, xaxt="n", col=1:10)
ageAxis(1, at=at, labels = names(values), col=1:10)

balloonplot(x, ..., xlab, ylab, zlab, show.zeros=FALSE,show.margins=TRUE,...)

balloonplot(x,y,z, xlab, ylab, zlab=deparse(substitute(z)),
dotsize=2/max(strwidth(19),strheight(19)),
dotchar=19,
dotcolor="skyblue",
text.size=1,
text.color=par("fg"), main,

balloonplot

Plot a graphical matrix where each cell contains a dot whose size reflects the relative magnitude of the corresponding component.

Description

Plot a graphical matrix where each cell contains a dot whose size reflects the relative magnitude of the corresponding component.

Usage

balloonplot(x, ...)

# Default S3 method:
balloonplot(x,y,z, xlab, ylab, zlab=deparse(substitute(z)),
dotsize=2/max(strwidth(19),strheight(19)),
dotchar=19,
dotcolor="skyblue",
text.size=1,
text.color=par("fg"), main,
Arguments

\texttt{x} \hspace{1em} A table object, or either a vector or a list of several categorical vectors containing grouping variables for the first (x) margin of the plotted matrix.

\texttt{y} \hspace{1em} Vector or list of vectors for grouping variables for the second (y) dimension of the plotted matrix.

\texttt{z} \hspace{1em} Vector of values for the size of the dots in the plotted matrix.

\texttt{xlab} \hspace{1em} Text label for the x dimension. This will be displayed on the x axis and in the plot title.

\texttt{ylab} \hspace{1em} Text label for the y dimension. This will be displayed on the y axis and in the plot title.

\texttt{zlab} \hspace{1em} Text label for the dot size. This will be included in the plot title.

\texttt{dotsize} \hspace{1em} Maximum dot size. You may need to adjust this value for different plot devices and layouts.

\texttt{dotchar} \hspace{1em} Plotting symbol or character used for dots. See the help page for the points function for symbol codes.

\texttt{dotcolor} \hspace{1em} Scalar or vector specifying the color(s) of the dots in the plot.

\texttt{text.size, text.color} \hspace{1em} Character size and color for row and column headers

\texttt{main} \hspace{1em} Plot title text.

\texttt{label} \hspace{1em} Boolean flag indicating whether the actual value of the elements should be shown on the plot.

\texttt{label.digits} \hspace{1em} Number of digits used in formatting value labels.

\texttt{label.size, label.color} \hspace{1em} Character size and color for value labels.

\texttt{scale.method} \hspace{1em} Method of scaling the sizes of the dot, either "volume" or "diameter". See below.
balloonplot

scale.range  Method for scaling original data to compute circle diameter. scale.range="absolute" scales the data relative to 0 (i.e., maps \([0, \max(z)] \rightarrow [0,1]\)), while scale.range="relative" scales the data relative to \(\min(z)\) (i.e., maps \([\min(z), \max(z)] \rightarrow [0,1]\)).

rowsrt, colsrt  Angle of rotation for row and column labels.

rowmar, colmar  Space allocated for row and column labels. Each unit is the width/height of one cell in the table.

show.zeros  boolean. If FALSE, entries containing zero will be left blank in the plotted matrix. If TRUE, zeros will be displayed.

show.margins  boolean. If TRUE, row and column sums are printed in the bottom and right margins, respectively.

cum.margins  boolean. If TRUE, marginal fractions are graphically presented in grey behind the row/column label area.

sorted  boolean. If TRUE, the rows will be arranged in sorted order by using the levels of the first y factor, then the second y factor, etc. The same process is used for the columns, based on the x factors.

label.lines  boolean. If TRUE, borders will be drawn for row and column level headers.

hide.duplicates  boolean. If TRUE, column and row headers will omit duplicates within row/column to reduce clutter. Defaults to TRUE.

fun  function to be used to combine data elements with the same levels of the grouping variables x and y. Defaults to sum.

...  Additional arguments passed to balloonplot.default or plot, as appropriate.

Details

This function plots a visual matrix. In each x,y cell a dot is plotted which reflects the relative size of the corresponding value of z. When scale.method="volume" the volume of the dot is proportional to the relative size of z. When scale.method="diameter", the diameter of the dot is proportional to the relative size of z. The "volume" method is default because the "diameter" method visually exaggerates differences.

Value

Nothing of interest.

Note

z is expected to be non-negative. The function will still operate correctly if there are negative values of z, but the corresponding dots will have 0 size and a warning will be generated.

Author(s)

Gregory R. Warnes <greg@warnes.net>

References

Function inspired by question posed on R-help by Ramon Alonso-Allende <allende@cnb.uam.es>.


### Examples

```r
# Create an example Data Frame Containing Car x Color data
carnames <- c("bmw", "renault", "mercedes", "seat")
carcolors <- c("red", "white", "silver", "green")
datavals <- round(rnorm(16, mean=100, sd=60), 1)
data <- data.frame(Car=rep(carnames,4),
                   Color=rep(carcolors, c(4,4,4,4)),
                   Value=datavals)
# show the data
data

# generate balloon plot with default scaling
balloonplot( data$Car, data$Color, data$Value)

# show margin label rotation & space expansion, using some long labels
levels(data$Car) <- c("BMW: High End, German", "Renault: Medium End, French",
                  "Mercedes: High End, German", "Seat: Imaginary, Unknown Producer")

# generate balloon plot with default scaling
balloonplot( data$Car, data$Color, data$Value, colmar=3, colsrt=90)

# Create an example using table
xnames <- sample( letters[1:3], 50, replace=2)
ynames <- sample( 1:5, 50, replace=2)

tab <- table(xnames, ynames)

balloonplot(tab)

# Example of multiple classification variables using the Titanic data
library(datasets)
data(Titanic)
dframe <- as.data.frame(Titanic) # convert to 1 entry per row format
attach(dframe)
balloonplot(x=Class, y=list(Survived, Age, Sex), z=Freq, sort=TRUE)

# colorize: survivors lightblue, non-survivors: grey
Colors <- Titanic
Colors[, , "Yes"] <- "skyblue"
Colors[, , "No"] <- "grey"
colors <- as.character(as.data.frame(Colors)$Freq)

balloonplot(x=list(Age, Sex),
            y=list(Class=Class,}
```
bandplot

Plot x-y Points with Locally Smoothed Mean and Standard Deviation

Description

Plot x-y points with curves for locally smoothed mean and standard deviation.

Usage

bandplot(x,...)
## S3 method for class 'formula'
bandplot(x, data, subset, na.action, ...
  xlab=NULL, ylab=NULL, add = FALSE, sd = c(-2:2),
  sd.col=c("magenta", "blue", "red", "blue", "magenta"),
  sd.lwd=c(2, 2, 3, 2, 2), sd.lty=c(2, 1, 1, 1, 2),
  method = "frac", width = 1/5, n=50)
## Default S3 method:
bandplot(x, y, ..., add = FALSE, sd = c(-2:2),
  sd.col=c("magenta", "blue", "red", "blue", "magenta"),
  sd.lwd=c(2, 2, 3, 2, 2), sd.lty=c(2, 1, 1, 1, 2),
  method = "frac", width = 1/5, n=50)

Arguments

x
numeric vector of y locations

y
either formula providing a single dependent variable (y) and an single independent variable (x) to use as coordinates in the scatter plot or a numeric vector of x locations

data
an optional data.frame, list, or environment containing the variables used in the model (and in subset). If not found in data, the variables are taken from environment(formula), typically the environment from which lm is called.

subset
an optional vector specifying a subset of observations to be used in the fitting process.

na.action
a function which indicates what should happen when the data contain NAs. The default is set by the na.action setting of options, and is na.fail if that is unset. The factory-fresh default is na.omit. Another possible value is NULL, no action. Value na.exclude can be useful.
bandplot Additional plotting parameters
xlab, ylab x and y axis labels
add Boolean indicating whether the local mean and standard deviation lines should be added to an existing plot. Defaults to FALSE.

sd Vector of multiples of the standard deviation that should be plotted. 0 gives the mean, -1 gives the mean minus one standard deviation, etc. Defaults to -2:2.

sd.col, sd.lwd, sd.lty Color, line width, and line type of each plotted line.
method, width, n Parameters controlling the smoothing. See the help page for wapply for details.

Details
bandplot was created to look for changes in the mean or variance of scatter plots, particularly plots of regression residuals.

The local mean and standard deviation are calculated by calling 'wapply'. By default, bandplot asks wapply to smooth using intervals that include the nearest 1/5 of the data. See the documentation of that function for details on the algorithm.

Value
Invisibly returns a list containing the x,y points plotted for each line.

Author(s)
Gregory R. Warnes <greg@warnes.net>

See Also
wapply, lowess

Examples

# fixed mean, changing variance
x <- 1:1000
y <- rnorm(1000, mean=1, sd=1 + x/1000 )
bandplot(x,y)
bandplot(y~x)

# fixed variance, changing mean
x <- 1:1000
y <- rnorm(1000, mean=x/1000, sd=1)
bandplot(x,y)

# changing mean and variance
#
x <- abs(rnorm(500))
y <- rnorm(500, mean=2*x, sd=2+2*x)

# the changing mean and dispersion are hard to see with the points alone:
plot(x, y)

# regression picks up the mean trend, but not the change in variance
reg <- lm(y~x)
summary(reg)
abline(reg=reg, col="blue", lwd=2)

# using bandplot on the original data helps to show the mean and
# variance trend
bandplot(y ~ x)

# using bandplot on the residuals helps to see that regression removes
# the mean trend but leaves the trend in variability
bandplot(predict(reg), resid(reg))

---

barplot2

Enhanced Bar Plots

Description

An enhancement of the standard barplot() function. Creates a bar plot with vertical or horizontal bars. Can plot confidence intervals for each bar, a lined grid behind the bars, change plot area color and logarithmic axes may be used.

Usage

## Default S3 method:
barplot2(height, width = 1, space = NULL,
    names.arg = NULL, legend.text = NULL, beside = FALSE,
    horiz = FALSE, density = NULL, angle = 45,
    col = NULL, pcol = NULL, border = par("fg"),
    main = NULL, sub = NULL, xlab = NULL, ylab = NULL,
    xlim = NULL, ylim = NULL, xpd = TRUE, log = "",
    axes = TRUE, axisnames = TRUE,
    cex.axis = par("cex.axis"), cex.names = par("cex.axis"),
    inside = TRUE, plot = TRUE, axis.lty = 0, offset = 0,
    plot.ci = FALSE, ci.l = NULL, ci.u = NULL,
    ci.color = "black", ci.lty = "solid", ci.lwd = 1, ci.width = 0.5,
    plot.grid = FALSE, grid.inc = NULL,
    grid.lty = "dotted", grid.lwd = 1, grid.col = "black",
    add = FALSE, panel.first = NULL, panel.last = NULL, ...)
Arguments

height either a vector or matrix of values describing the bars which make up the plot. If height is a vector, the plot consists of a sequence of rectangular bars with heights given by the values in the vector. If height is a matrix and beside is FALSE then each bar of the plot corresponds to a column of height, with the values in the column giving the heights of stacked “sub-bars” making up the bar. If height is a matrix and beside is TRUE, then the values in each column are juxtaposed rather than stacked.

width optional vector of bar widths. Re-cycled to length the number of bars drawn. Specifying a single value will no visible effect unless xlim is specified.

space the amount of space (as a fraction of the average bar width) left before each bar. May be given as a single number or one number per bar. If height is a matrix and beside is TRUE, space may be specified by two numbers, where the first is the space between bars in the same group, and the second the space between the groups. If not given explicitly, it defaults to c(0,1) if height is a matrix and beside is TRUE, and to 0.2 otherwise.

names.arg a vector of names to be plotted below each bar or group of bars. If this argument is omitted, then the names are taken from the names attribute of height if this is a vector, or the column names if it is a matrix.

legend.text a vector of text used to construct a legend for the plot, or a logical indicating whether a legend should be included. This is only useful when height is a matrix. In that case given legend labels should correspond to the rows of height; if legend.text is true, the row names of height will be used as labels if they are non-null.

beside a logical value. If FALSE, the columns of height are portrayed as stacked bars, and if TRUE the columns are portrayed as juxtaposed bars.

horiz a logical value. If FALSE, the bars are drawn vertically with the first bar to the left. If TRUE, the bars are drawn horizontally with the first at the bottom.

density a vector giving the the density of shading lines, in lines per inch, for the bars or bar components. 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), for the bars or bar components.

col a vector of colors for the bars or bar components. By default, grey is used if height is a vector, and heat.colors(nrow(height)) if height is a matrix.

prcol the color to be used for the plot region.

border the color to be used for the border of the bars.

main, sub overall and sub titles for the plot.

xlab a label for the x axis.

ylab a label for the y axis.

xlim limits for the x axis.

ylim limits for the y axis.

xpd logical. Should bars be allowed to go outside region?
log

a character string which contains "x" if the x axis is to be logarithmic, "y" if the y axis is to be logarithmic and "xy" or "yx" if both axes are to be logarithmic.

axes

logical. If TRUE, a vertical (or horizontal, if horiz is true) axis is drawn.

axisnames

logical. If TRUE, and if there are names.arg (see above), the other axis is drawn (with lty = 0) and labeled.

cex.axis

expansion factor for numeric axis labels.

cex.names

expansion factor for names.

inside

logical. If TRUE, the lines which divide adjacent (non-stacked!) bars will be drawn. Only applies when space = 0 (which it partly is when beside = TRUE).

plot

logical. If FALSE, nothing is plotted.

axis.lty

the graphics parameter lty applied to the axis and tick marks of the categorical (default horizontal) axis. Note that by default the axis is suppressed.

offset

a vector indicating how much the bars should be shifted relative to the x axis.

plot.ci

logical. If TRUE, confidence intervals are plotted over the bars. Note that if a stacked bar plot is generated, confidence intervals will not be plotted even if plot.ci = TRUE

ci.l,ci.u

The confidence intervals (ci.l = lower bound, ci.u = upper bound) to be plotted if plot.ci = TRUE. Values must have the same dim structure as height.

ci.color

the color for the confidence interval line segments

ci.lty

the line type for the confidence interval line segments

ci.lwd

the line width for the confidence interval line segments

ci.width

length of lines used for the "t" at the end of confidence interval line segments, as a multiple of width. Defaults to 0.5.

plot.grid

if TRUE a lined grid will be plotted behind the bars

grid.inc

the number of grid increments to be plotted

grid.lty

the line type for the grid

grid.lwd

the line width for the grid

grid.col

the line color for the grid

add

logical, if TRUE add barplot to current plot.

panel.first

An expression to be evaluated after the plot region coordinates have been set up, but prior to the drawing of the bars and other plot region contents. This can be useful to add additional plot region content behind the bars. This will also work if add = TRUE

panel.last

An expression to be evaluated after the bars have been drawn, but prior to the addition of confidence intervals, a legend and the axis annotation

... further graphical parameters (par) are passed to plot.window(), title() and axis.

Details

This is a generic function, it currently only has a default method. A formula interface may be added eventually.
Value

A numeric vector (or matrix, when beside = TRUE), say mp, giving the coordinates of all the bar midpoints drawn, useful for adding to the graph.

If beside is true, use colMeans(mp) for the midpoints of each group of bars, see example.

Note

Prior to R 1.6.0, barplot behaved as if axis.lty = 1, unintentionally. 0 (zero) and NA values in height will not be plotted if using logarithmic scales. If there are NA values in height and beside = FALSE, values after the NA will not be plotted in stacked bars.

Author(s)

Original barplot() by R-Core. Enhancements by Marc Schwartz <marc\_schwartz@comcast.net>

See Also

plot(..., type = "h"), dotchart, hist.

Examples

tN <- table(Ni <- rpois(100, lambda = 5))
r <- barplot2(tN, col = 'gray')

#- type = "h" plotting *is* `bar'plot
lines(r, tN, type = 'h', col = 'red', lwd = 2)

barplot2(tN, space = 1.5, axisnames = FALSE,
        sub = "barplot2(..., space = 1.5, axisnames = FALSE")

data(VADeaths, package = "datasets")
barplot2(VADeaths, plot = FALSE)
barplot2(VADeaths, plot = FALSE, beside = TRUE)

mp <- barplot2(VADeaths) # default
tot <- colMeans(VADeaths)
text(mp, tot + 3, format(tot), xpd = TRUE, col = "blue")
barplot2(VADeaths, beside = TRUE,
        col = c("lightblue", "mistyrose", "lightcyan",
        "lavender", "cornsilk"),
        legend = rownames(VADeaths), ylim = c(0, 100))
title(main = "Death Rates in Virginia", font.main = 4)

# Example with confidence intervals and grid
hh <- t(VADeaths)[, 5:1]
mybarcol <- "gray20"
ci.l <- hh * 0.85
ci.u <- hh * 1.15
mp <- barplot2(hh, beside = TRUE,
        col = c("lightblue", "mistyrose",
        "lightcyan", "lavender"),
        sub = "barplot2(..., beside = TRUE",
        col = c("lightblue", "mistyrose",
        "lightcyan", "lavender"),
        legend = rownames(hh), ylim = c(0, 100))
title(main = "Death Rates in Virginia", font.main = 4)
boxplot

Produce a Boxplot Annotated with the Number of Observations

Description

This function uses boxplot to produce a boxplot which is then annotated with the number of observations in each group.

Usage

boxplot2(..., top=FALSE, shrink=1, textcolor=NULL)

Arguments

... parameters passed to boxplot.

top logical indicating whether the number of observations should be added to the top or the bottom of the plotting region. Defaults to FALSE.

shrink value to shrink character size (cex) when annotating.

textcolor text color.

Note

This function replaces boxplot.n, which has been deprecated avoid potential problems with S3 method dispatching.
ci2d

Author(s)
Gregory R. Warnes <greg@warnes.net>

See Also
boxplot, text

Examples

data(state)

# n's at bottom
boxplot2( state.area ~ state.region)

# n's at top
boxplot2( state.area ~ state.region, top=TRUE)

# small red text
boxplot2( state.area ~ state.region, shrink=0.8, textcolor="red")

Description
Create 2-dimensional empirical confidence regions from provided data.

Usage

\[
\text{ci2d}(x, y = \text{NULL},
    \text{nbins}=51, \text{method}=c("bkde2D","hist2d"),
    \text{bandwidth}, \text{factor}=1.0,
    \text{ci.levels}=c(0.50,0.75,0.90,0.95,0.975),
    \text{show}=c("filled.contour","contour","image","none"),
    \text{col=topo.colors(length(breaks)-1)},
    \text{show.points} = \text{FALSE},
    \text{pch=par("pch")},
    \text{points.col} = \text{"red"},
    \text{xlab, ylab,}
    \ldots)
\]

## S3 method for class 'ci2d'

print(x, \ldots)
Arguments

x either a vector containing the x coordinates or a matrix with 2 columns.
y a vector containing the y coordinates, not required if ‘x’ is matrix
nbins number of bins in each dimension. May be a scalar or a 2 element vector. Defaults to 51.
method One of "bkde2D" (for KernSmooth::bkde2d) or "hist2d" (for gplots::hist2d) specifying the name of the method to create the 2-d density summarizing the data. Defaults to "bkde2D".
bandwidth Bandwidth to use for KernSmooth::bkde2d. See below for default value.
factor Numeric scaling factor for bandwidth. Useful for exploring effect of changing the bandwidth. Defaults to 1.0.
cl.levels Confidence level(s) to use for plotting data. Defaults to c(0.5, 0.75, 0.9, 0.95, 0.975)
show Plot type to be displayed. One of "filled.contour", "contour", "image", or "none". Defaults to "filled.contour".
show.points Boolean indicating whether original data values should be plotted. Defaults to TRUE.
pch Point type for plots. See points for details.
points.col Point color for plotting original data. Defaults to "red".
col Colors to use for plots.
xlab, ylab Axis labels
... Additional arguments passed to KernSmooth::bkde2d or gplots::hist2d.

Details

This function utilizes either KernSmooth::bkde2d or gplots::hist2d to estimate a 2-dimensional density of the data passed as an argument. This density is then used to create and (optionally) display confidence regions.

When bandwidth is omitted and method="bkde2d", KernSmooth::dpik is applied in x and y dimensions to select the bandwidth.

Value

A ci2d object consisting of a list containing (at least) the following elements:

nobs number of original data points
x x position of each density estimate bin
y y position of each density estimate bin
density Matrix containing the probability density of each bin (count in bin/total count)
cumDensity Matrix where each element contains the cumulative probability density of all elements with the same density (used to create the confidence region plots)
contours List of contours of each confidence region.
call Call used to create this object
Note

Confidence intervals generated by ci2d are approximate, and are subject to biases and/or artifacts induced by the binning or kernel smoothing method, bin locations, bin sizes, and kernel bandwidth.

The conf2d function in the r2d2 package may create a more accurate confidence region, and reports the actual proportion of points inside the region.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

bkde2D, conf2d, dpik, hist2d

Examples

###
## Basic usage
###
data(geyser, package="MASS")

x <- geyser$duration
y <- geyser$waiting

# 2-d confidence intervals based on binned kernel density estimate
ci2d(x,y)  # filled contour plot
ci2d(x,y, show.points=TRUE)  # show original data

# image plot
ci2d(x,y, show="image")
ci2d(x,y, show="image", show.points=TRUE)

# contour plot
ci2d(x,y, show="contour", col="black")
ci2d(x,y, show="contour", col="black", show.points=TRUE)

###
## Control Axis scales
###
x <- rnorm(2000, sd=4)
y <- rnorm(2000, sd=1)

# 2-d confidence intervals based on binned kernel density estimate
.ci2d(x,y)

# 2-d confidence intervals based on 2d histogram
ci2d(x,y, method="hist2d", nbins=25)

# Require same scale for each axis, this looks oval
ci2d(x,y, range.x=list(c(-20,20), c(-20,20)))
ci2d(x,y, method=\"hist2d\", same.scale=TRUE, nbins=25) # hist2d

###
## Control smoothing and binning
###
x <- rnorm(2000, sd=4)
y <- rnorm(2000, mean=x, sd=2)

# Default 2-d confidence intervals based on binned kernel density estimate

ci2d(x,y)

# change the smoother bandwidth

ci2d(x,y, bandwidth=c(sd(x)/8, sd(y)/8))

# change the smoother number of bins

ci2d(x,y, nbins=10)
ci2d(x,y)
ci2d(x,y, nbins=100)

# Default 2-d confidence intervals based on 2d histogram

ci2d(x,y, method=\"hist2d\", show.points=TRUE)

# change the number of histogram bins

ci2d(x,y, nbins=10, method=\"hist2d\", show.points=TRUE)
ci2d(x,y, nbins=25, method=\"hist2d\", show.points=TRUE)

###
## Perform plotting manually
###
data(geyser, package=\"MASS\")

# let ci2d handle plotting contours...
ci2d(geyser$duration, geyser$waiting, show=\"contour\", col=\"black\")

# call contour() directly, show the 90 percent CI, and the mean point

est <- ci2d(geyser$duration, geyser$waiting, show=\"none\")
contour(est$x, est$y, est$cumDensity,
    xlab=\"duration\", ylab=\"waiting\",
    levels=0.90, lwd=4, lty=2)
points(mean(geyser$duration), mean(geyser$waiting),
    col=\"red\", pch=\"x\")

###
## Extract confidence region values
###
data(geyser, package=\"MASS\")

## Empirical 90 percent confidence limits
quantile( geyser$duration, c(0.05, 0.95) )
quantile( geyser$waiting, c(0.05, 0.95) )
## Bivariate 90 percent confidence region

```r
est <- ci2d(geyser$duration, geyser$waiting, show="none")
names(est$contours) # show available contours

ci.90 <- est$contours[names(est$contours)=="0.9"] # get region(s)
ci.90 <- rbind(ci.90[[1]],NA, ci.90[[2]], NA, ci.90[[3]]) # join them

print(ci.90) # show full contour
range(ci.90$x, na.rm=TRUE) # range for duration
range(ci.90$y, na.rm=TRUE) # range for waiting
```

### Visually compare confidence regions

```r
# Bivariate smoothed 90 percent confidence region
est <- ci2d(geyser$duration, geyser$waiting, show="none")
names(est$contours) # show available contours

ci.90 <- est$contours[names(est$contours)=="0.9"] # get region(s)
ci.90 <- rbind(ci.90[[1]],NA, ci.90[[2]], NA, ci.90[[3]]) # join them

plot( waiting ~ duration, data=geyser, 
     main="Comparison of 90 percent confidence regions")
polygon( ci.90, col="green", border="green", density=10)

# Univariate Normal-Theory 90 percent confidence region
mean.x <- mean(geyser$duration)
mean.y <- mean(geyser$waiting)

dx <- sd(geyser$duration)
dy <- sd(geyser$waiting)

t.value <- qt(c(0.05,0.95), df=gdata::nobs(geyser$duration), lower=TRUE)
ci.x <- mean.x + t.value* dx

ci.y <- mean.y + t.value* dy

plotCI(mean.x, mean.y, 
       li=ci.x[1],
       ui=ci.x[2],
       barcol="blue", col="blue",
       err="x",
       pch="X",
       add=TRUE)

plotCI(mean.x, mean.y, 
       li=ci.y[1],
       ui=ci.y[2],
       barcol="blue", col="blue",
       err="y",
       pch=NA,
       add=TRUE)
```
# rect(ci.x[1], ci.y[1], ci.x[2], ci.y[2], border="blue",
# density=5,
# angle=45,
# col="blue")

## Empirical univariate 90 percent confidence region
box <- cbind( x=quantile( geyser$duration, c(0.05, 0.95 )),
              y=quantile( geyser$waiting, c(0.05, 0.95 )))
rect(box[1,1], box[1,2], box[2,1], box[2,2], border="red",
     density=5,
     angle=-45,
     col="red")

## now a nice legend
legend("topright", legend=c("Region type",
                           "Univariate Normal Theory",
                           "Univariate Empirical",
                           "Smoothed Bivariate"),
       lwd=c(NA,1,1,1),
       col=c("black","blue","red","green"),
       lty=c(NA,1,1,1))

###
### Test with a large number of points
###
### Not run:
x <- rnorm(60000, sd=1)
y <- c( rnorm(40000, mean=x, sd=1),
       rnorm(20000, mean=x+4, sd=1) )
hist2d(x,y)
ct <- ci2d(x,y)
ct

## End(Not run)

---

```r

col2hex(cname)
```

**Description**

Convert color names to hex RGB strings

**Usage**

```r
col2hex(cname)
```
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>cname</code></td>
<td>Color name(s)</td>
</tr>
</tbody>
</table>

Value

Character vector giving the hex color code translation of the provided color names.

Author(s)

Gregory R. Warnes

See Also

col2rgb, colors.rgb

Examples

col2hex(c("red","yellow","lightgrey"))

colorpanel  Generate a smoothly varying set of colors

Description

colorpanel generate a set of colors that varies smoothly. redgreen, greenred, bluered, and redblue generate red-black-green, green-black-red, red-white-blue, and blue-white-red colorbars, respectively. colors

Usage

colorpanel(n, low, mid, high)
redgreen(n)
greenred(n)
bluered(n)
redblue(n)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>n</code></td>
<td>Desired number of color elements in the panel.</td>
</tr>
<tr>
<td><code>low</code>, <code>mid</code>, <code>high</code></td>
<td>Colors to use for the Lowest, middle, and highest values. mid may be omitted.</td>
</tr>
</tbody>
</table>

Details

The values for `low`, `mid`, `high` can be given as color names ("red"), plot color index (2=red), and HTML-style RGB, ("#FF0000"=red).

If `mid` is supplied, then the returned color panel will consist of `n - floor(n/2)` HTML-style RGB elements which vary smoothly between `low` and `mid`, then between `mid` and `high`. Note that if `n` is even, the color `mid` will occur twice at the center of the sequence.

If `mid` is omitted, the color panel will vary smoothly between `low` and `high`. 
Value

Vector of HTML-style RGB colors.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

colors

Examples

```r
showpanel <- function(col)
{
  image(z=matrix(1:100, ncol=1), col=col, xaxt="n", yaxt="n")
}

par(mfrow=c(3,3))

# two colors only:
showpanel(colorpanel(8, low="red", high="green"))

# three colors
showpanel(colorpanel(8, "red", "black", "green"))
# note the duplicatation of black at the center, using an odd
# number of elements resolves this:
showpanel(colorpanel(9, "red", "black", "green"))

showpanel(greenred(64))
showpanel(redgreen(64))
showpanel(bluered(64))
showpanel(redblue(64))
```

---

gplots-defunct

Defunct functions

Description

These functions are defunct and have been removed from the gplots package.

Usage

```r
boxplot(..., top=FALSE, shrink=1, textcolor=NULL)
plot.lm2(
    x,
    which = 1:5,
```
caption = c("Residuals vs Fitted", "Normal Q-Q plot", "Scale-Location plot", "Cook's distance plot"),
panel = panel.smooth,
sub.caption = deparse(x$call),
main = "",
ask,
..., id.n = 3,
labels.id = names(residuals(x)),
cex.id = 0.75,
band=TRUE, rug=TRUE, width=1/10, max.n=5000
)
smartlegend(x = c("left", "center", "right"), y = c("top", "center", "bottom"),
..., inset = 0.05)

Arguments

ask, band, caption, cex.id, id.n, inset, labels.id, main, max.n, panel, rug, shrink, sub.caption, textcolor, top, which, width, x, y

see man page for the corresponding replacement function

Details

These functions are no longer available. Please refer to the manual page for the replacement function:

- boxplot.n has been replaced by boxplot2
- plot.lm2 has been replaced by lmplot2
- smartlegend is no longer needed because relative positioning has been implemented in legend.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

boxplot2, lmplot2, legend, Defunct
gplots-deprecated  Deprecated functions

Description

These functions have been deprecated and will be removed in future releases of gplots.

Usage

## No deprecated functions at this time ##

Details

These functions have been deprecated. Please refer to the manual page for the replacement function:

- (No deprecated functions at this time)

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

- Deprecated

heatmap.2  Enhanced Heat Map

Description

A heat map is a false color image (basically \texttt{image}(t(x))) with a dendrogram added to the left side and/or to the top. Typically, reordering of the rows and columns according to some set of values (row or column means) within the restrictions imposed by the dendrogram is carried out.

This heatmap provides a number of extensions to the standard R \texttt{ heatmap} function.

Usage

\begin{verbatim}
heatmap.2 (x,
    # dendrogram control
    Rowv = TRUE,
    Colv=if(symm)"Rowv" else TRUE,
    distfun = dist,
    hclustfun = hclust,
    dendrogram = c("both","row","column","none"),
    reorderfun = function(d, w) reorder(d, w),
\end{verbatim}
symm = FALSE,

# data scaling
scale = c("none", "row", "column"),
na.rm=TRUE,

# image plot
revC = identical(Colv, "Rowv"),
add.expr,

# mapping data to colors
breaks,
symbreaks=any(x < 0, na.rm=TRUE) || scale!="none",

# colors
col="heat.colors",

# block sepration
colsep,
rowsep,
seppcolor="white",
sepwidth=c(0.05, 0.05),

# cell labeling
cellnote,
notecex=1.0,
notecol="cyan",
na.color=par("bg"),

# level trace
trace=c("column", "row", "both", "none"),
tracecol="cyan",
hline=median(breaks),
vline=median(breaks),
linecol=tracecol,

# Row/Column Labeling
margins = c(5, 5),
ColSideColors,
RowSideColors,
cexRow = 0.2 + 1/log10(nr),
cexCol = 0.2 + 1/log10(nc),
labRow = NULL,
labCol = NULL,
srtRow = NULL,
srtCol = NULL,
adjRow = c(0, NA),
adjCol = c(NA, 0),
offsetRow = 0.5,
offsetCol = 0.5,
colRow = NULL,
colCol = NULL,

# color key + density info
density.info=c("histogram","density","none"),
denscol=tracecol,
symkey = any(x < 0, na.rm=TRUE) || symbreaks,
densadj = 0.25,
key.title = NULL,
key.xlab = NULL,
key.ylab = NULL,
key.xtickfun = NULL,
key.ytickfun = NULL,
key.par=list(),

# plot labels
denscol = tracecol,
symkey = any(x < 0, na.rm=TRUE) || symbreaks,
densadj = 0.25,
key.title = NULL,
key.xlab = NULL,
key.ylab = NULL,
key.xtickfun = NULL,
key.ytickfun = NULL,
key.par=list(),

# plot layout
lmat = NULL,
llhei = NULL,
lwid = NULL,

# extras
extrafun=NULL,
...
)

Arguments

x numeric matrix of the values to be plotted.

Rowv determines if and how the row dendrogram should be reordered. By default, it is TRUE, which implies dendrogram is computed and reordered based on row means. If NULL or FALSE, then no dendrogram is computed and no reordering is done. If a dendrogram, then it is used "as-is", i.e. without any reordering. If a vector of integers, then dendrogram is computed and reordered based on the order of the vector.

Colv determines if and how the column dendrogram should be reordered. Has the options as the Rowv argument above and additionally when x is a square matrix, Colv="Rowv" means that columns should be treated identically to the rows.
**distfun** function used to compute the distance (dissimilarity) between both rows and columns. Defaults to `dist`.

**hclustfun** function used to compute the hierarchical clustering when `Rowv` or `Colv` are not dendrograms. Defaults to `hclust`.

**dendrogram** character string indicating whether to draw 'none', 'row', 'column' or 'both' dendrograms. Defaults to 'both'. However, if `Rowv` (or `Colv`) is FALSE or NULL and dendrogram is 'both', then a warning is issued and `Rowv` (or `Colv`) arguments are honoured.

**reorderfun** function of dendrogram and weights for reordering the row and column dendrograms. The default uses `stats(reorder.dendrogram)`.

**symm** logical indicating if `x` should be treated symmetrically; can only be true when `x` is a square matrix.

**scale** character indicating if the values should be centered and scaled in either the row direction or the column direction, or none. The default is "none".

**na.rm** logical indicating whether NA’s should be removed.

**revC** logical indicating if the column order should be reversed for plotting, such that e.g., for the symmetric case, the symmetry axis is as usual.

**add.expr** expression that will be evaluated after the call to `image`. Can be used to add components to the plot.

**breaks** (optional) Either a numeric vector indicating the splitting points for binning `x` into colors, or a integer number of break points to be used, in which case the break points will be spaced equally between `min(x)` and `max(x)`.

**symbreaks** Boolean indicating whether breaks should be made symmetric about 0. Defaults to TRUE if the data includes negative values, and to FALSE otherwise.

**col** colors used for the image. Defaults to heat colors (`heat.colors`).

**colsep**, **rowsep**, **sepcolor** (optional) vector of integers indicating which columns or rows should be separated from the preceding columns or rows by a narrow space of color `sepcolor`.

**sepwidth** (optional) Vector of length 2 giving the width (colsep) or height (rowsep) the separator box drawn by `colsep` and `rowsep` as a function of the width (colsep) or height (rowsep) of a cell. Defaults to `c(0.05, 0.05)`.

**cellnote** (optional) matrix of character strings which will be placed within each color cell, e.g. p-value symbols.

**notecex** (optional) numeric scaling factor for `cellnote` items.

**notecol** (optional) character string specifying the color for `cellnote` text. Defaults to "cyan".

**na.color** Color to use for missing value (NA). Defaults to the plot background color.

**trace** character string indicating whether a solid "trace" line should be drawn across 'row's or down 'column's, 'both' or 'none'. The distance of the line from the center of each color-cell is proportional to the size of the measurement. Defaults to 'column'.

**tracecol** character string giving the color for "trace" line. Defaults to "cyan".
hline, vline, linecol

Vector of values within cells where a horizontal or vertical dotted line should
be drawn. The color of the line is controlled by linecol. Horizontal lines
are only plotted if trace is 'row' or 'both'. Vertical lines are only drawn if
trace 'column' or 'both'. hline and vline default to the median of the breaks,
linecol defaults to the value of tracecol.

margins
numeric vector of length 2 containing the margins (see par(mar= *)) for col-
umn and row names, respectively.

ColSideColors (optional) character vector of length ncol(x) containing the color names for a
horizontal side bar that may be used to annotate the columns of x.

RowSideColors (optional) character vector of length nrow(x) containing the color names for a
vertical side bar that may be used to annotate the rows of x.

cexRow, cexCol positive numbers, used as cex.axis in for the row or column axis labeling. The
defaults currently only use number of rows or columns, respectively.

labRow, labCol character vectors with row and column labels to use; these default to rownames(x)
or colnames(x), respectively.

srtRow, srtCol angle of row/column labels, in degrees from horizontal

adjRow, adjCol 2-element vector giving the (left-right, top-bottom) justification of row/column
labels (relative to the text orientation).

offsetRow, offsetCol

Number of character-width spaces to place between row/column labels and the
edge of the plotting region.

colRow, colCol color of row/column labels, either a scalar to set the color of all labels the same,
or a vector providing the colors of each label item

key logical indicating whether a color-key should be shown.

keysize numeric value indicating the size of the key

density.info character string indicating whether to superimpose a 'histogram', a 'density'
plot, or no plot ('none') on the color-key.

denscol character string giving the color for the density display specified by density.info,
defaults to the same value as tracecol.

symkey Boolean indicating whether the color key should be made symmetric about 0.
Defaults to TRUE if the data includes negative values, and to FALSE otherwise.

densadj Numeric scaling value for tuning the kernel width when a density plot is drawn
on the color key. (See the adjust parameter for the density function for de-
tails.) Defaults to 0.25.

key.title main title of the color key. If set to NA no title will be plotted.

density.info x axis label of the color key. If set to NA no label will be plotted.
density.info y axis label of the color key. If set to NA no label will be plotted.
density.info function computing tick location and labels for the xaxis of the color key. Re-
turns a named list containing parameters that can be passed to axis. See exam-
density.info

density.info function computing tick location and labels for the y axis of the color key. Re-
turns a named list containing parameters that can be passed to axis. See exam-
density.info

key.par  
graphical parameters for the color key. Named list that can be passed to par.

main, xlab, ylab  
main, x- and y-axis titles; defaults to none.

lmat, lhei, lwid  
visual layout: position matrix, column height, column width. See below for details

extrafun  
A function to be called after all other work. See examples.

...  
additional arguments passed on to image

Details

If either Rowv or Colv are dendrograms they are honored (and not reordered). Otherwise, dendrograms are computed as dd <- as.dendrogram(hclustfun(distfun(X))) where X is either x or t(x).

If either is a vector (of “weights”) then the appropriate dendrogram is reordered according to the supplied values subject to the constraints imposed by the dendrogram, by reorder(dd, Rowv), in the row case.

If either is missing, as by default, then the ordering of the corresponding dendrogram is by the mean value of the rows/columns, i.e., in the case of rows, Rowv <- rowMeans(x, na.rm=na.rm).

If either is NULL, no reordering will be done for the corresponding side.

If scale="row" (or scale="col") the rows (columns) are scaled to have mean zero and standard deviation one. There is some empirical evidence from genomic plotting that this is useful.

The default colors range from red to white (heat.colors) and are not pretty. Consider using enhancements such as the RColorBrewer package, https://cran.r-project.org/package=RColorBrewer to select better colors.

By default four components will be displayed in the plot. At the top left is the color key, top right is the column dendrogram, bottom left is the row dendrogram, bottom right is the image plot. When RowSideColor or ColSideColor are provided, an additional row or column is inserted in the appropriate location. This layout can be overridden by specifying appropriate values for lmat, lwid, and lhei. lmat controls the relative position of each element, while lwid controls the column width, and lhei controls the row height. See the help page for layout for details on how to use these arguments.

Value

Invisibly, a list with components

rowInd  row index permutation vector as returned by order.dendrogram.

colInd  column index permutation vector.

call  the matched call

rowMeans, rowSDs  mean and standard deviation of each row: only present if scale="row"

colMeans, colSDs  mean and standard deviation of each column: only present if scale="column"
carpet  reordered and scaled 'x' values used generate the main 'carpet'
rowDendrogram  row dendrogram, if present
colDendrogram  column dendrogram, if present
breaks  values used for color break points
col  colors used
vline  center-line value used for column trace, present only if trace="both" or trace="column"
hline  center-line value used for row trace, present only if trace="both" or trace="row"
colorTable  A three-column data frame providing the lower and upper bound and color for each bin
layout  A named list containing the values used for lmat, lhei, and lwid.

Note
The original rows and columns are reordered to match the dendrograms Rowv and Colv (if present).
heatmap.2() uses layout to arrange the plot elements. Consequently, it can not be used in a multi column/row layout using layout(...), par(mfrow=...) or (mfcol=...).

Author(s)
Andy Liaw, original; R. Gentleman, M. Maechler, W. Huber, G. Warnes, revisions.

See Also
image, hclust

Examples

data(mtcars)
x <- as.matrix(mtcars)
rc <- rainbow(nrow(x), start=0, end=.3)
cc <- rainbow(ncol(x), start=0, end=.3)

##
## demonstrate the effect of row and column dendrogram options
##
heatmap.2(x)  ## default - dendrogram plotted and reordering done.
heatmap.2(x, dendrogram="none")  ## no dendrogram plotted, but reordering done.
heatmap.2(x, dendrogram="row")  ## row dendrogram plotted and row reordering done.
heatmap.2(x, dendrogram="col")  ## col dendrogram plotted and col reordering done.
heatmap.2(x, keysize=2)  ## default - dendrogram plotted and reordering done.

heatmap.2(x, Rowv=FALSE, dendrogram="both")  ## generates a warning!
heatmap.2(x, Rowv=NULL, dendrogram="both")  ## generates a warning!
heatmap.2(x, Colv=FALSE, dendrogram="both")  ## generates a warning!

## Reorder dendrogram by branch means rather than sums
heatmap.2(x, reorderfun=function(d, w) reorder(d, w, agglo.FUN = mean) )
## plot a sub-cluster using the same color coding as for the full heatmap

```r
full <- heatmap.2(x)
heatmap.2(x, Colv=full$colDendrogram[[2]], breaks=full$breaks)  # column subset
heatmap.2(x, Rowv=full$rowDendrogram[[1]], breaks=full$breaks)  # row subset
heatmap.2(x, Colv=full$colDendrogram[[2]], Rowv=full$rowDendrogram[[1]], breaks=full$breaks)  # both
```

## Show effect of row and column label rotation

```r
heatmap.2(x, srtCol=NULL)
heatmap.2(x, srtCol=0, adjCol = c(0.5,1))
heatmap.2(x, srtCol=45, adjCol = c(1,1))
heatmap.2(x, srtCol=135, adjCol = c(1,0))
heatmap.2(x, srtCol=180, adjCol = c(0.5,0))
heatmap.2(x, srtCol=225, adjCol = c(0,0))  ## not very useful
heatmap.2(x, srtCol=270, adjCol = c(0,0.5))
heatmap.2(x, srtCol=315, adjCol = c(0,1))
heatmap.2(x, srtCol=360, adjCol = c(0.5,1))

heatmap.2(x, srtRow=45, adjRow=c(0, 1))
heatmap.2(x, srtRow=45, adjRow=c(0, 1), srtCol=45, adjCol=c(1,1))
heatmap.2(x, srtRow=45, adjRow=c(0, 1), srtCol=270, adjCol=c(0,0.5))
```

## Show effect of `offsetRow`/`offsetCol` (only works when `srtRow/srtCol` is not also present)

```r
heatmap.2(x, offsetRow=0, offsetCol=0)
heatmap.2(x, offsetRow=1, offsetCol=1)
heatmap.2(x, offsetRow=2, offsetCol=2)
heatmap.2(x, offsetRow=-1, offsetCol=-1)

heatmap.2(x, srtRow=0, srtCol=90, offsetRow=0, offsetCol=0)
heatmap.2(x, srtRow=0, srtCol=90, offsetRow=1, offsetCol=1)
heatmap.2(x, srtRow=0, srtCol=90, offsetRow=2, offsetCol=2)
heatmap.2(x, srtRow=0, srtCol=90, offsetRow=-1, offsetCol=-1)
```

## Show how to use `extrafun` to replace the `key` with a scatterplot

```r
lmat <- rbind(c(5,3,4), c(2,1,4))
lhei <- c(1.5, 4)
lwid <- c(1.5, 4, 0.75)

myplot <- function() {
  oldpar <- par("mar")
  par(mar=c(5.1, 4.1, 0.5, 0.5))
  plot(mpg ~ hp, data=x)
}

heatmap.2(x, lmat=lmat, lhei=lhei, lwid=lwid, key=FALSE, extrafun=myplot)
```

## Show how to customize the color key

```r
heatmap.2(x,
  key.title=NA, # no title
```
key.xlab=NA, # no xlab
key.par=list(mgp=c(1.5, 0.5, 0),
            mar=c(2.5, 2.5, 1, 0)),
key.tickfun=function() {
  breaks <- parent.frame()$breaks
  return(list(
    at=parent.frame()$scale01(c(breaks[1],
                                breaks[length(breaks)])),
    labels=c(as.character(breaks[1]),
             as.character(breaks[length(breaks)]))
  ))
}

heatmap.2(x,
          breaks=256,
          key.title=NA,
          key.xlab=NA,
          key.par=list(mgp=c(1.5, 0.5, 0),
                        mar=c(1, 2.5, 1, 0)),
          key.tickfun=function() {
            cex <- par("cex")*par("cex.axis")
            side <- 1
            line <- 0
            col <- par("col.axis")
            font <- par("font.axis")
            mtext("low", side=side, at=0, adj=0,
                   line=line, cex=cex, col=col, font=font)
            mtext("high", side=side, at=1, adj=1,
                   line=line, cex=cex, col=col, font=font)
            return(list(labels=FALSE, tick=FALSE))
          })

##
## Show effect of z-score scaling within columns, blue-red color scale
##
hv <- heatmap.2(x, col=bluered, scale="column", tracecol="#303030")

###
### Look at the return values
###
names(hv)

## Show the mapping of z-score values to color bins
hv$colorTable

## Extract the range associated with white
hv$colorTable[hv$colorTable[,"color"]=="#FFFFFF",]

## Determine the original data values that map to white
whiteBin <- unlist(hv$colorTable[hv$colorTable[,"color"]=="#FFFFFF",1:2])
rbind(whiteBin[1] * hv$colSDs + hv$colMeans,
      whiteBin[2] * hv$colSDs + hv$colMeans )
## A more decorative heatmap, with z-score scaling along columns

```r
hv <- heatmap.2(x, col=cm.colors(255), scale="column",
                  RowSideColors=rc, ColSideColors=cc, margin=c(5, 10),
                  xlab="specification variables", ylab="Car Models",
                  main="heatmap(<Mtcars data>, ...., scale="column")",
                  tracecol="green", density="density")
```

## Note that the breakpoints are now symmetric about 0

```r
hv <- heatmap.2(x, col=cm.colors(255), scale="column",
                  RowSideColors=rc, ColSideColors=cc, margin=c(5, 10),
                  xlab="specification variables", ylab="Car Models",
                  main="heatmap(<Mtcars data>, ...., scale="column")",
                  tracecol="green", density="density", colRow=rc, colCol=cc,
                  srtCol=45, adjCol=c(0.5,1))
```

```r
data(attitude)
round(Ca <- cor(attitude), 2)
symnum(Ca) # simple graphic
```

# with reorder
```r
heatmap.2(Ca, symm=TRUE, margin=c(6, 6), trace="none")
```

# without reorder
```r
heatmap.2(Ca, Rowv=FALSE, symm=TRUE, margin=c(6, 6), trace="none")
```

## Place the color key below the image plot
```r
heatmap.2(x, lmat=rbind( c(0, 3), c(2,1), c(0,4) ), lhei=c(1.5, 4, 2))
```

## Place the color key to the top right of the image plot
```r
heatmap.2(x, lmat=rbind( c(0, 3, 4), c(2,1,0) ), lwid=c(1.5, 4, 2))
```

## For variable clustering, rather use distance based on cor():
```r
data(USJudgeRatings)
symnum(cU <- cor(USJudgeRatings))
```

```r
hU <- heatmap.2(cU, Rowv=FALSE, symm=TRUE, col=topo.colors(16),
                distfun=function(c) as.dist(1 - c), trace="none")
```

## The Correlation matrix with same reordering:
```r
hM <- format(round(cU, 2))
hM
```

# now with the correlation matrix on the plot itself
```r
heatmap.2(cU, Rowv=FALSE, symm=TRUE, col=rev(heat.colors(16)),
          distfun=function(c) as.dist(1 - c), trace="none",
          Rowv=FALSE, symm=TRUE, col=topo.colors(16),
          distfun=function(c) as.dist(1 - c), trace="none")
```
## hist2d

**Compute and Plot a 2-Dimensional Histogram**

### Description

Compute and plot a 2-dimensional histogram.

### Usage

```r
hist2d(x, y=NULL, nbins=200, same.scale=FALSE, na.rm=TRUE, show=TRUE,
       col=c("black", heat.colors(12)), FUN=base::length, xlab, ylab,
       ...)  
```

### Arguments

- **x**: either a vector containing the x coordinates or a matrix with 2 columns.
- **y**: a vector containing the y coordinates, not required if `x` is matrix.
- **nbins**: number of bins in each dimension. May be a scalar or a 2 element vector. Defaults to 200.
- **same.scale**: use the same range for x and y. Defaults to FALSE.
- **na.rm**: Indicates whether missing values should be removed. Defaults to TRUE.
- **show**: Indicates whether the histogram be displayed using `image` once it has been computed. Defaults to TRUE.
**hist2d**

**col**
Colors for the histogram. Defaults to "black" for bins containing no elements, a set of 16 heat colors for other bins.

**FUN**
Function used to summarize bin contents. Defaults to `base::length`. Use, e.g., `mean` to calculate means for each bin instead of counts.

**xlab, ylab**
(Optional) x and y axis labels

... Parameters passed to the `image` function.

**Details**
This function creates a 2-dimensional histogram by cutting the x and y dimensions into nbins sections. A 2-dimensional matrix is then constructed which holds the counts of the number of observed (x,y) pairs that fall into each bin. If show=TRUE, this matrix is then then passed to `image` for display.

**Value**
A list containing 5 elements:
- **counts** Matrix containing the number of points falling into each bin
- **x.b breaks, y.b breaks** Lower and upper limits of each bin
- **x,y** midpoints of each bin

**Author(s)**
Gregory R. Warnes <greg@warnes.net>

**See Also**
`image`, `persp`, `hist`, `freq2d`

**Examples**
```r
## example data, bivariate normal, no correlation
x <- rnorm(2000, sd=4)
y <- rnorm(2000, sd=1)

## separate scales for each axis, this looks circular
hist2d(x,y)

## same scale for each axis, this looks oval
hist2d(x,y, same.scale=TRUE)

## use different ## bins in each dimension
hist2d(x,y, same.scale=TRUE, nbins=c(100,200) )

## use the hist2d function to create an h2d object
h2d <- hist2d(x,y,show=FALSE, same.scale=TRUE, nbins=c(20,30))

## show object summary
```
h2d

## object contents
str(h2d)

## perspective plot
persp( h2d$x, h2d$y, h2d$counts,
       ticktype="detailed", theta=30, phi=30,
       expand=0.5, shade=0.5, col="cyan", ltheta=-30)

## for contour (line) plot ...
contour( h2d$x, h2d$y, h2d$counts, nlevels=4 )

## for a filled contour plot ...
filled.contour( h2d$x, h2d$y, h2d$counts, nlevels=4,
                col=gray((4:0)/4 )

---

**lmplot2**

Plots to assess the goodness of fit for the linear model objects

**Description**

Plots to assess the goodness of fit for the linear model objects

**Usage**

```r
lmplot2(
  x,
  which = 1:5,
  caption = c("Residuals vs Fitted", "Normal Q-Q plot",
            "Scale-Location plot", "Cook's distance plot"),
  panel = panel.smooth,
  sub.caption = deparse(x$call),
  main = "",
  ask = interactive() && nb.fig < length(which)
  && .Device != "postscript",
  ...,
  id.n = 3,
  labels.id = names(residuals(x)),
  cex.id = 0.75,
  band=TRUE,
  rug=TRUE,
  width=1/10,
  max.n=5000
)
```
Arguments

- **x**: lm object
- **which**: Numerical values between 1 and 5, indicating which plots to be shown. The codes are:
  - 1: Fitted vs residuals
  - 2: Normal Q-Q
  - 3: Scale-Location
  - 4: Cook’s distance
  - 5: Residuals vs. predictor
- **caption**: Caption for each type of plot
- **panel**: function to draw on the existing plot
- **sub.caption**: SubCaption for the plots
- **main**: Main title of the plot
- **ask**: whether interactive graphics
- **...**: parameters passed to lmplot2.
- **id.n**: integer value, less than or equal to residuals of lm object
- **labels.id**: Names of the residuals of the lm object
- **cex.id**: Parameter to control the height of text strings
- **band**: logical vector indicating whether bandplot should also be plotted
- **rug**: logical vector indicating whether rug should be added to the existing plot
- **width**: Fraction of the data to use for plot smooths
- **max.n**: Maximum number of points to display in plots

Note

This function replaces plot.lm2, which has been deprecated to avoid potential problems with S3 method dispatching.

Author(s)

Gregory R. Warnes <greg@warnes.net> and Nitin Jain <nitin.jain@pfizer.com>

See Also

plot.lm

Examples

```r
cnt1 <- rnorm(100, 4)
cnt2 <- rnorm(100, 4.5)
group <- gl(2,100,200, labels=c("Ctl","Trt"))
weight <- c(cnt1, cnt2)
wt.err <- rnorm(length(weight), mean=weight, sd=1/2)
x <- lm(weight ~ group + wt.err)
```
lowess

Scatter Plot Smoothing

Description

The `lowess` function performs the computations for the LOWESS smoother (see the reference below). `lowess` returns a an object containing components `x` and `y` which give the coordinates of the smooth. The smooth can then be added to a plot of the original points with the function `lines`. Alternatively, `plot` can be called directly on the object returned from `lowess` and the 'lowess' method for `plot` will generate a scatterplot of the original data with a `lowess` line superimposed. Finally, the `plotlowess` function both calculates the `lowess` smooth and plots the original data with a `lowess` smooth.

Usage

lowess(x, ...)

## Default S3 method:
lowess(x, y=NULL, f=2/3, iter=3L, delta=0.01 *
diff(range(x)), ...)

## S3 method for class 'formula'
lowess(formula, data=parent.frame(), ..., subset, f=2/3,
iter=3L, delta=.01*diff(range(mf[-response])))

## S3 method for class 'lowess'
plot(x, y, ..., col.lowess="red", lty.lowess=2)

plotlowess(formula, data=parent.frame(), ..., subset=parent.frame(),
col.lowess="red", lty.lowess=2 )

Arguments

formula      formula providing a single dependent variable (y) and an single independent variable (x) to use as coordinates in the scatter plot.
data         a data.frame (or list) from which the variables in 'formula' should be taken.
subset       an optional vector specifying a subset of observations to be used in the fitting process.
x, y         vectors giving the coordinates of the points in the scatter plot. Alternatively a single plotting structure can be specified.
the smoother span. This gives the proportion of points in the plot which influence the smooth at each value. Larger values give more smoothness.

iter the number of robustifying iterations which should be performed. Using smaller values of iter will make lowess run faster.

delta values of x which lie within delta of each other replaced by a single value in the output from lowess.

... parameters for methods.

col.lowess, lty.lowess
color and line type for plotted line

References


See Also

loess (in package {modreg}), a newer formula based version of lowess (with different defaults!).

Examples

data(cars)

# # x, y method

plot(cars$speed, cars$dist, main="lowess(cars)"
lines(lowess(cars$speed, cars$dist), col=2)
lines(lowess(cars$speed, cars$dist, f=.2), col=3)
legend(5, 120, c(paste("f="", c("2/3", ","))), lty=1, col=2:3)

# # formula method: plot, then calculate the lowess smoother, # then add smooth to the plot

plot(dist ~ speed, data=cars, main="lowess(cars)"
lines(lowess(dist ~ speed, data=cars), col=2, lty=2)
lines(lowess(dist ~ speed, data=cars, f=.2), col=3) # smaller bandwidth
legend(5, 120, c(paste("f="", c("2/3", ","))), lty=1, col=2:3)

# # formula method: calculate lowess() smoother, then call plot() # on the lowess object

lw <- lowess(dist ~ speed, data=cars)
plot(lw, main="lowess(cars)"

#
# formula method: calculate and plot in a single command

```r
plotLowess(dist ~ speed, data=cars, main="lowess(cars)")
```

## ooplot.default

*Create an OpenOffice style plot*

### Description

An extension of barplot2. Creates bar- and line-plots mimicking the style of OpenOffice plots. This utility can plot the values next to each point or bar as well as confidence intervals.

### Usage

```r
ooplot(data, ...)
```

### Arguments

- **data**: a matrix of values describing the values that make up the plot. The first column of `data` is taken as the axis against which all the other values are plotted. The first column of `data` may not be sparse.
- **width**: optional vector of barwidths. Recycled to the number of bars drawn. A single value will have no visible effect.
space

the amount of space left before each bar. May be given as a single number or one
number per bar. If type is stackbar, space may be specified by two numbers,
where the first is the space between bars in the same group, and the second the
space between groups. Defaults to c(0, 1) if type is a stackbar, and to 0.2
otherwise.

names.arg

a vector of names to be plotted below each bar or group of bars. If this argument
is omitted, then the names are taken from the row names of data.

legend.text

a vector of text used to construct a legend for the plot, or a logical indicating
whether a legend should be included; if legend.text is true, the row names of
data will be used as labels if they are non-null.

horiz

a logical value. If FALSE, the bars are drawn vertically with the first bar to the
left. If TRUE, the bars are drawn horizontally with the first at the bottom.

density

a vector giving the the density of shading lines, in lines per inch, for the bars
or bar components. 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), for
the bars or bar components.

kmg

the set of SI units to convert, defaults to "fpnumkMGTP". See below for details.

kmglim

logical. If FALSE the conversion to SI units is not performed. Default is TRUE.

type

a string indicating the preferred format of the plot, choices are: xyplot : plot
where y is plotted against the x-value. linear : plot where y values are plotted
against equidistant x-values. barplot : plot where y values are represented as
bars against equidistant x-values. stackplot : plot where y values are stacked for
identical x-values and bars are equidistant.

col

a vector of colors for the bars or bar components.

prcol

the color to be used for the plot region.

border

the color to be used for the border of the bars.

main, sub

overall and sub titles for the plot.

xlab

a label for the x axis.

ylab

a label for the y axis.

xlim

limits for the x axis.

ylim

limits for the y axis.

xpd

logical. Should bars be allowed to go outside region?

log

a character string which contains "x" if the x axis is to be logarithmic, "y"
if the y axis is to be logarithmic and "xy" or "yx" if both axes are to be
logarithmic.

axes

logical. If TRUE, a vertical (or horizontal, if horiz is true) axis is drawn.

axisnames

logical. If TRUE, and if there are names.arg (see above), the other axis is drawn
(with lty=0) and labeled.

prval

logical. If TRUE, then values are plotted above all points and bars.

lm

logical. If TRUE, the linear fit is plotted.
cex.axis, cex.names, cex.values
character scaling factor for numeric axis labels, names, and displayed values, respectively.

inside logical. If TRUE, the lines which divide adjacent (non-stacked!) bars will be drawn. Only applies when space = 0 (which it partly is when beside = TRUE).

plot logical. If FALSE, nothing is plotted.

axis.lty the graphics parameter lty applied to the axis and tick marks of the categorical (default horizontal) axis. Note that by default the axis is suppressed.

plot.ci logical. If TRUE, confidence intervals are plotted over the bars. Note that if a stacked bar plot is generated, confidence intervals will not be plotted even if plot.ci = TRUE

ci.l, ci.u The confidence intervals (ci.l = lower bound, ci.u = upper bound) to be plotted if plot.ci = TRUE. Values must have the same dim structure as height.

ci.color the color for the confidence interval line segments

ci.lty the line type for the confidence interval line segments

plot.grid the line width for the confidence interval line segments

grid.inc if TRUE a lined grid will be plotted behind the bars

grid.lty the number of grid increments to be plotted

grid.lwd the line type for the grid

grid.lwd the line width for the grid

grid.col the line color for the grid

add logical, if TRUE add barplot to current plot.

by.row Logical value. If TRUE the data matrix is organized with variables along rows rather than down columns.

... further graphical parameters (par) are passed to plot.window(), title() and axis.

Details

Plot units are automatically scaled to SI units based on the maximum value present, according to the set of units specified by characters in the kmg parameter. These letters are interpreted as

P  peta = 1E15
T  tera = 1E12
G  giga = 1E09
M  mega = 1E06
k  kilo = 1E03
m  milli = 1E-03
u  micro = 1E-06
n  nano = 1E-09
p  pico = 1E-12
f  femto = 1E-15

with the default being "fpnumkMGTP" (all of these units). For example, if the largest value plotted is 1243000, it would be presented as 1.234M.
Value

A numeric vector (or matrix, when beside = TRUE), say mp, giving the coordinates of all the bar midpoints drawn, useful for adding to the graph.

If beside is true, use colMeans(mp) for the midpoints of each group of bars, see example.

Author(s)

Lodewijk Bonebakker <bonebakker@comcast.net> with modifications by Gregory R. Warnes <greg@warnes.net>. Based on barplot2().

See Also

plot, boxplot

Examples

data(VADeaths, package = "datasets")
VADeaths <- cbind( Age=c(50,55,60,65,70), VADeaths)

mp <- ooplot(VADeaths) # default
mp <- ooplot(VADeaths, type="xyplot") # same as default
mp <- ooplot(VADeaths, type="linear") # linear scale
mp <- ooplot(VADeaths, type="linear", log="y") # log scale on y axis
mp <- ooplot(VADeaths, type="barplot") # barplot
mp <- ooplot(VADeaths, type="stackbar") # stacked

tot <- colMeans(VADeaths[,1])
ooplot(VADeaths,
    col = c("lightblue", "mistyrose", "lightcyan", "lavender"),
    legend = colnames(VADeaths)[-1], ylim = c(0, 100),
    type="barplot", cex.values=0.75)
title(main = "Death Rates in Virginia", font.main = 4)

##
## Capability demo
##
## examples for the ooplot routine
##
## create some test data
test1 <- data.frame(x=c(0,1,2,3,4), y=c(0,1,2,3,4))
test2 <- data.frame(x=c(0,1,2,3,4), par=c(0,1,4,9,16))
test3 <- data.frame(x=c(-2,-1,0,1,2),y2=c(4,1,0,1,4))
## single line test example
test1f <- test1
## two column example
test2F <- merge(test1,test2,by.x="x",all=TRUE,sort=TRUE)
## three column example
test3f <- merge(test2F,test3,by.x="x",all=TRUE,sort=TRUE)
## overplot

**Plot multiple variables on the same region, with appropriate axes**

**Description**

overplot graphs a set of variables defined on the same x-range but which have varying y-ranges on the same plotting area. For each set of y-values it uses a different color and line-type and and draws a correspondingly colored and line-typed axis. panel.overplot is used by overplot to draw the individual graphs.

**Usage**

```r
overplot(formula, data = parent.frame(), same.scale = FALSE, xlab, ylab,
         xlim, ylim, min.y, max.y, log = "", panel = "panel.overplot",
         subset, plot = TRUE, groups, main, f = 2/3, ...)```

overplot

Arguments

- **formula**: Formula describing the x and y variables. It should be of the form x ~ y|z. The conditioning variable (z) should be a factor.
- **same.scale**: Logical value indicating whether the plot region should have the same range for all plots. Defaults to FALSE.
- **xlab, ylab, xlim, ylim, main**: Standard plotting parameters. See `plot` for details.
- **min.y, max.y**: Scalar or vector values used to specify the y plotting limits for individual plots. If a single scalar value is provided, it will be used for all plots. These parameters can be used specify one end of the individual plot ranges, while allowing the other end to vary with the data. EG, to force 0 to always be within the plot region.
- **log**: character string "", 'x', 'y', or 'xy', indicating which axes should be plotted on a log scale. Defaults to " (neither).
- **panel**: a plotting function to be called to draw the individual plots. Defaults to `overplot.panel`, which plots the points and a lowess smooth.
- **plot**: Logical value indicating whether to draw the plot.
- **groups**: (optional) character vector giving the names of levels of the conditioning variable to plot. Defaults to all levels of the conditioning variable.
- **f**: Smoothing parameter for lowess
- **data, subset, ...**: parameters passed to `model.frame` to obtain the data to be plotted from the formula.

Details

This function essentially performs

```r
tmp <- split(data, z)
for(i in levels(z))
  plot( x ~ y, data=tmp[[z]] )
```

except that all of the plots are shown on the same plotting region and varying scales for each value of z are handled nicely.

Value

A copy of the data split by the conditioning variable.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

`interaction.plot, coplot` for alternative visualizations of 3-way data.
Examples

# Example teratogenicity rtPCR data
data(rtpcr)

dataHrtpcr
C same scale
overplotH rq ~ Conc..ug.ml. | Test.Substance,
data=rtpcr,
subset=Detector=="ProbeType 1" & Conc..ug.ml. > 0,
same.scale=TRUE,
log="xy",
f=3/4,
main="Detector=ProbeType 1",
xlab="Concentration (ug/ml)",
ylab="Relative Gene Quantification"
)

# different scales, but force lower limit to 0.01
overplotH rq ~ Conc..ug.ml. | Test.Substance,
data=rtpcr,
subset=Detector=="ProbeType 8" & Conc..ug.ml. > 0,
log="xy",
f=3/4,
main="Detector=ProbeType 8",
xlab="Concentration (ug/ml)",
ylab="Relative Gene Quantification",
min.y=0.01
)

plotCI

Plot Error Bars and Confidence Intervals

Description

Given a set of x and y values and interval width or upper and lower bounds, plot the points with error bars. This can be a useful tool for visualizing confidence intervals.

Usage

plotCI(x, y = NULL, uiw = uiw, liw = uiw, ui, li, err='y', ylim=NULL, xlim=NULL, type="p", col=par("col"), barcol=col,
pt.bg = par("bg"), sfrac = 0.01, gap=1, lwd=par("lwd"),
lty=par("lty"), labels=FALSE, add=FALSE, xlab, ylab, minbar,
maxbar, ...)
### Arguments

- **x, y**: coordinates for the center of error bars. `y` defaults to `1:n`.
- **uiw**: width of the upper or right error bar. Set to NULL or NA to omit upper bars.
- **liw**: width of the lower or left error bar. Defaults to same value as `uiw`. Set to NULL or NA to omit lower bars.
- **ui**: upper end of error bars. Defaults to `y + uiw` or `x + uiw` depending on `err`. Set to NULL or NA to omit upper bars.
- **li**: lower end of error bars. Defaults to `y - liw` or `x - liw` depending on `err`. Set to NULL or NA to omit lower bars.
- **err**: direction for error bars. Set to "y" for vertical bars. Set to "x" for horizontal bars. Defaults to "y".
- **col**: color of plotting character used center marker of error bars. Default is "black".
- **xlim, ylim**: range of `x/y` values to include in the plotting area.
- **type**: point/line type; passed to `points`.
- **barcol**: color of the error bars. Defaults to the same value as `col`.
- **pt.bg**: background color of points (use `pch=21, pt.bg=par("bg")` to get open points superimposed on error bars).
- **sfrac**: width of "crossbar" at the end of error bar as a fraction of the `x` plotting region. Defaults to 0.01.
- **gap**: space left between the center of the error bar and the lines marking the error bar in units of the height (width) of the letter "O". Defaults to 1.0
- **lwd**: width of bar lines.
- **lty**: line type of bar lines.
- **labels**: either a logical value indicating whether the circles representing the `x` values should be replaced with text giving the actual values or a vector containing labels to use instead. Defaults to `FALSE`.
- **add**: logical indicating whether error bars should be added to the current plot. If `FALSE` (the default), a new plot will be created and symbols/labels for the `x` values will be plotted before drawing error bars.
- **minbar**: minimum allowed value for bar ends. If specified, values smaller than `minbar` will be replaced with `minbar`.
- **maxbar**: maximum allowed value for bar ends. If specified, values larger than `maxbar` will be replaced with `maxbar`.
- **xlab**: label for `x` axis.
- **ylab**: label for `y` axis.

### Author(s)

Original version by Bill Venables `<w venable@attunga.stats.adelaide.edu.au>` posted to `r-help` on Sep. 20, 1997. Enhanced version posted to `r-help` by Ben Bolker `<ben@zoo.ufl.edu>` on Apr. 16, 2001. This version was modified and extended by Gregory R. Warnes `<greg@warnes.net>`.

Additional changes suggested by Martin Maechler `<maechler@stat.math.ethz.ch>` integrated on July 29, 2004.
See Also

plotmeans provides an enhanced wrapper to plotCI.

Examples

# plot means and
data(state)
tmp <- split(state.area, state.region)
means <- sapply(tmp, mean)
stdev <- sqrt(sapply(tmp, var))
n <- sapply(tmp, length)

ciw <- qt(0.975, n) * stdev / sqrt(n)

# plain
plotCI(x=means, uiw=ciw)

# prettier
plotCI(x=means, uiw=ciw, col="black", barcol="blue", lwd=1)

# give mean values
plotCI(x=means, uiw=ciw, col="black", barcol="blue",
labels=round(means, -3), xaxt="n", xlim=c(0, 5) )
axis(side=1, at=1:4, labels=names(tmp), cex=0.7)

# better yet, just use plotmeans ... #
plotmeans( state.area ~ state.region )

---

plotmeans  
Plot Group Means and Confidence Intervals

Description

Plot group means and confidence intervals.

Usage

plotmeans(formula, data=NULL, subset, na.action,
bars=TRUE, p=0.95, minsd=0, minbar, maxbar,
xlab=names(mf)[2], ylab=names(mf)[1], mean.labels=FALSE,
ci.label=FALSE, n.label=TRUE, text.n.label="n",
digits=getOption("digits"), col="black", barwidth=1,
barcol="blue", connect=TRUE, ccol=
col, legends=names(means), xaxt, use.t=TRUE,
lwd=par("lwd"), ...)
**plotmeans**  

**Arguments**

- **formula**: symbolic expression specifying the outcome (continuous) and grouping variable (factor). See `lm()` for details.
- **data**: optional data frame containing the variables in the model.
- **subset**: an optional vector specifying a subset of observations to be used in the fitting process.
- **na.action**: a function which indicates what should happen when the data contain ‘NA’s. See `lm()` for details.
- **bars**: a logical value indicating whether confidence interval bars should be plotted. Defaults to TRUE.
- **p**: confidence level for error bars. Defaults to 0.95.
- **minsd**: minimum permitted value for the standard deviation within each factor level. Any standard deviation estimates smaller than `minsd` will be replaced with `minsd`. Defaults to 0.
- **minbar**: minimum allowed value for bar ends. If specified, values smaller than `minbar` will be replaced with `minbar`.
- **maxbar**: maximum allowed value for bar ends. If specified, values larger than `maxbar` will be replaced with `maxbar`.
- **xlab**: x-axis label.
- **ylab**: y-axis label.
- **mean.labels**: either a logical value indicating whether the circles representing the group means should be replaced with text giving the actual mean values or a vector containing labels to use instead. Defaults to FALSE.
- **ci.label**: a logical value indicating whether text giving the actual interval end values should be placed at the end of each confidence interval bar. Defaults to FALSE.
- **n.label**: a logical value indicating whether text giving the number of observations in each group should be added to the plot.
- **text.n.label**: prefix text for labeling observation counts. Defaults to "n=".
- **digits**: number of significant digits to use when displaying mean or confidence limit values.
- **col**: color of circles marking group means. Default is "black".
- **barwidth**: linewidth of interval bars and end marks. Default is 1.
- **barcol**: color of interval bars and end marks. Default is "blue".
- **connect**: either a logical value indicating whether the means of each group should be connected by a line, or a list of vectors giving the index of bars that should be connected by a line. Defaults to TRUE.
- **ccol**: color of lines used to connect means. Defaults to the same color as "col".
- **legends**: vector containing strings used to label groups along the x axis. Defaults to group names.
- **xaxt**: a character which specifies the axis type. Specifying "n" causes an axis to be set up, but not plotted.
use.t

A logical value indicating whether the t distribution should be used to compute confidence intervals. If TRUE, the default, a t distribution will the correct number of degrees of freedom for each group be used. If FALSE, the a normal distribution will be used.

lwd

Width of connecting lines

...:

Optional plotting parameters.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

plotCI, boxplot

Examples

# show comparison with boxplot
data(state)
plotmeans(state.area ~ state.region)

# show some color and mean labels
plotmeans(state.area ~ state.region,
          mean.labels=TRUE, digits=3,
          col="red", connect=FALSE)

# show how to specify which means should be connected
plotmeans(state.area ~ state.region, connect=list(1:2, 3:4),
          ccol="red", pch=7)

# more complicated example showing how to show an interaction
data(esoph)
par(las=2,
    las=2, mar=c(10.1,4.1,4.1,2.1),
    mgp=c(8.1,0)
)
plotmeans(ncases/ncontrols ~ interaction(agegp, alcgp, sep=" "),
          connect=list(1:6,7:12,13:18,19:24),
          barwidth=2,
          col="dark green",
          data=esoph,
          xlab="Age Group and Alcohol Consumption",
          ylab="# Cases / # Controls",
          main="("Fraction of Cases for by Age and Alcohol Consumption",
          "Ile-et-Vilaine Esophageal Cancer Study")"
)
abline(v=c(6.5, 12.5, 18.5), lty=2)
qqnorm.aov

Makes a half or full normal plot for the effects from an aov model

Description

Makes a half or full normal plot for the effects from a model inheriting from class aov. One can interactively label the points in the plot.

Usage

```r
## S3 method for class 'aov'
qqnorm(y, full=FALSE, label=FALSE, omit=NULL,
       xlab=paste(if (full) "" else "Half", " Normal plot"),
       ylab="Effects", ...)
```

Arguments

- `y` A model object inheriting from aov
- `full` Full or half normal plot (half is default)
- `label` If TRUE, function allows interactive labelling of points in plot, using the mouse
- `omit` Numeric or character vector of effects to omit, the intercept is always omitted
- `xlab` Horizontal axix label
- `ylab` Vertical axis label
- `...` Further arguments to be given to the plot function

Details

Produces a (half) normal plot of the effects from an AOV model. The idea behind the plot is that most effects will be small or null, and this effects can be used as a basis for estimation of the experimental variance. This small effects will show up in the plot as a straight line, other effects can be judged against this as a background. Heavily used by Box, Hunter \& Hunter, which attributes the idea to Daniel.

This is a simpler implementation than the one in S-Plus.

Value

If `label=TRUE`, the vector of points identified, else nothing of interest.

Author(s)

Kjetil Halvorsen <kjetil@entelnet.bo>
References

Box, Hunter and Hunter: Statistics for Experimenters. An Introduction to Design, Data Analysis and Model Building. Wiley.
Daniel, C (1959): Use of half-normal plot in interpreting factorial two-level experiments. Technometrics 1, 149.

Examples

```r
library(MASS)
data(npk)
npk.aov <- aov(yield ~ block + N*P*K, npk)
qqnorm(npk.aov)

## interactive labeling of points. Click mouse on points to show label.
if (dev.interactive()) qqnorm(npk.aov, omit=2:6, label=TRUE)
```

---

residplot

Undocumented functions

Description

These functions are undocumented. Some are internal and not intended for direct use. Others simply haven’t been documented yet.

Usage

```r
residplot(model, formula, ...)
```

Arguments

- `model` Undocumented
- `formula` Undocumented
- `...` arguments to be passed to `fun`

Details

These functions are undocumented. Some are internal and not intended for direct use. Others simply haven’t been documented yet.

Author(s)

Gregory R. Warnes <greg@warnes.net>
Description

Create a vector of n colors that are perceptually equidistant and in an order that is easy to interpret.

Usage

rich.colors(n, palette="temperature", alpha=1.0, rgb=FALSE, plot=FALSE)

Arguments

n  number of colors to generate.
palette  palette to use: "temperature" contains blue-green-yellow-red, and "blues" contains black-blue-white.
alpha  alpha transparency, from 0 (fully transparent) to 1 (opaque).
rgb  if TRUE then a matrix of RGBA values is included as an attribute.
plot  whether to plot a descriptive color diagram.

Value

A character vector of color codes.

Author(s)

Arniragnusson.

See Also

rgb, rainbow, heat.colors.

Examples

m <- absmatrix(1:120+runorm(120), nrow=15, ncol=8))
opar <- par(bg="gray", mfrow=c(1,2))
matplot(m, type="l", lty=1, lwd=3, col=rich.colors(8))
matplot(m, type="l", lty=1, lwd=3, col=rich.colors(8,"blues"))
par(opar)

barplot(rep(1,100), col=rich.colors(100), space=0, border=0, axes=FALSE)
barplot(rep(1,20), col=rich.colors(40)[11:30]) # choose subset

plot(m, rev(m), ylim=c(120,0), pch=16, cex=2,
col=rich.colors(200,"blues",alpha=0.6)[1:120]) # semitransparent

rich.colors(100, plot=TRUE) # describe rgb recipe
rtPCR

Teratogenesis rtPCR data

Description

rtPCR data for experiments investigating a variety of markers for characterizing teratogenicity.

Usage

data(rtPCR)

Format

A data frame with 1672 observations on the following 21 variables.

PlateID a factor with levels A0027002 through A0054019
Test.Substance a factor with levels Compound A through Compound H
Teratogenicity.in.vivo a factor with levels Non Strong Weak / Moderate
Sample a factor with levels Sample 1 - Sample 152
Rep. a factor with levels Rep 1 - Rep 21
Label a factor with levels Ctrl, Neg. Ctrl P1 - P9, No Vehicle Ctrl, and Pos. Ctrl
Conc..ug.ml. a numeric vector
Detector a factor with levels ProbeType 1 - ProbeType 17
Avg.delta.Ct a numeric vector
delta.Ct.SD a numeric vector
delta.delta.Ct a numeric vector
RQ a numeric vector
X..RQ a numeric vector
X100..Custom.. a numeric vector
X100..Custom.. a numeric vector
Custom.. a numeric vector
Custom...1 a numeric vector
RQ.Min a numeric vector
RQ.Max a numeric vector
Threshold a numeric vector
sinkplot

Details

TBA

Source

Anonymized data.

Examples

data(rtPCR)

# same scale
overplot( RQ ~ Conc..ug.ml. | Test.Substance,
  data=rtPCR,
  subset=Detector=="ProbeType 7" & Conc..ug.ml. > 0,
  same.scale=TRUE,
  log="xy",
  f=3/4,
  main="Detector=ProbeType 7",
  xlab="Concentration (ug/ml)",
  ylab="Relative Gene Quantification"
)

# different scales, but force lower limit to 0.01
overplot( RQ ~ Conc..ug.ml. | Test.Substance,
  data=rtPCR,
  subset=Detector=="ProbeType 7" & Conc..ug.ml. > 0,
  log="xy",
  f=3/4,
  main="Detector=ProbeType 7",
  xlab="Concentration (ug/ml)",
  ylab="Relative Gene Quantification",
  min.y=0.01
)

sinkplot  

Send textual R output to a graphics device

Description

Divert R's standard text output to a graphics device.

Usage

sinkplot(operation = c("start", "plot", "cancel"), ...)

Arguments

operation  See below
...  Plot arguments. (Ignored unless operation="plot").
sinkplot

Details

This function allows the printed output of R commands to be captured and displayed on a graphics device.

The capture process is started by calling sinkplot("start"). Now R commands can be executed and all printed output (except errors) will be captured. When the desired text has been captured sinkplot("plot") can be called to actually display the output. sinkplot("cancel") can be used to abort the output capture without plotting.

The current implementation does not allow sinkplot to be nested.

Value

Invisibly returns a character vector containing one element for each line of the captured output.

Author(s)

Gregory R. Warnes <greg@warnes.net>

References


See Also

capture.output, textplot

Examples

set.seed(12456)
x <- factor(sample( LETTERS[1:5], 50, replace=TRUE))
y <- rnorm(50, mean=as.numeric(x), sd=1)

# construct a figure showing a box plot of the data, followed by an
# analysis of variance table for the data
layout(cbind(1:2), heights=c(2,1))

boxplot(y~x, col="darkgreen")
sinkplot()
anova(lm(y~x))
sinkplot("plot", col="darkgreen")
space

Space points in an x-y plot so they don’t overlap.

Usage

space(x, y, s=1/50, na.rm=TRUE, direction="x")

Arguments

x numeric vector of x coordinates.

y numeric vector of x coordinates.

s either a single numeric value or 2 element vector specifying the minimum distance between points in the x and y dimensions as a fraction of the x and y range. Defaults to 1/50.

na.rm logical indicating whether pairs where one or both elements are missing should be removed. Defaults to TRUE.

direction "x" or "y", indicating which direction points should be moved to accomplish spacing.

Details

In an x-y plot where at least one variable has discrete levels several points may be plotted at or very near the same coordinates. This makes it difficult to gauge the number of points in a specific region. A common method of resolving this problem is to ‘jitter’ the points by adding random noise.

This function takes a different approach to the same problem.

When there are two or more points with the same (x,y) value (or within x+s[1] and x+s[2]), it spaces these out in the x direction so that the points are separated by at least distance s.

Another method for dealing with overplotting is available in the sunflowerplot function.

Value

list with two components

x (modified) x location for each input point

y y location of each input point

Author(s)

Gregory R. Warnes <greg@warnes.net>
textplot

Display text information in a graphics plot.

Description

This function displays text output in a graphics window. It is the equivalent of 'print' except that the output is displayed as a plot.

Usage

textplot(object, halign="center", valign="center", cex, ...)  
## Default S3 method:  
textplot(object, halign=c("center", "left", "right"),  
       valign=c("center", "top", "bottom"), cex, ...)  
## S3 method for class 'character'
textplot(object, halign = c("center", "left", "right"),
  valign = c("center", "top", "bottom"), cex, fixed.width=TRUE,
  cspace=1, lspace=1, mar=c(0, 0, 3, 0) + 0.1,
  tab.width = 8, ...)
## S3 method for class 'data.frame'
textplot(object, halign = c("center", "left", "right"),
  valign = c("center", "top", "bottom"), cex, ...)
## S3 method for class 'matrix'
textplot(object, halign = c("center", "left", "right"),
  valign = c("center", "top", "bottom"), cex, cmar = 2,
  rmar = 0.5, show.rownames = TRUE, show.colnames = TRUE,
  hadj = 1, vadj = 1, mar = c(1, 1, 4, 1) + 0.1,
  col.data = par("col"), col.rownames = par("col"),
  col.colnames = par("col"), ...)

Arguments

object Object to be displayed.
halign Alignment in the x direction, one of "center", "left", or "right".
valign Alignment in the y direction, one of "center", "top", or "bottom"
cex Character size, see par for details. If unset, the code will attempt to use the
  largest value which allows the entire object to be displayed.
fixed.width Logical value indicating whether to emulate a fixed-width font by aligning charac-
  ters in each row of text. This is usually necessary for text-formatted tables
display properly. Defaults to 'TRUE'.
cspace Space between characters as a multiple of the width of the letter 'W'. This only
  applies when fixed.width==TRUE.
lspace Line spacing. This only applies when fixed.width==TRUE.
mar Figure margins, see the documentation for par.
rmar, cmar Space between rows or columns, in fractions of the size of the letter 'M'.
show.rownames, show.colnames Logical value indicating whether row or column names will be displayed.

adj, vadj Vertical and horizontal location of elements within matrix cells. These have the
  same meaning as the adj graphics parameter (see par).
col.data Colors for data elements. If a single value is provided, all data elements will
  be the same color. If a matrix matching the dimensions of the data is provided,
  each data element will receive the specified color.
col.rownames, col.colnames Colors for row names and column names, respectively. Either may be specified
  as a scalar or a vector of appropriate length.
tab.width Width of a single tab stop, in characters
...
Optional arguments passed to the text plotting command or specialied object
  methods
Details

A new plot is created and the object is displayed using the largest font that will fit on in the plotting region. The halign and valign parameters can be used to control the location of the string within the plotting region.

For matrices and vectors a specialized textplot function is available, which plots each of the cells individually, with column widths set according to the sizes of the column elements. If present, row and column labels will be displayed in a bold font.

Value

The character scaling factor (cex) used.

Author(s)

Gregory R. Warnes <greg@warnes.net>

See Also

plot, text, capture.output

Examples

```r
## Not run:
### simple examples
# show R version information
textplot(version)

# show the alphabet as a single string
textplot( paste(letters[1:26], collapse=" ") )

# show the alphabet as a matrix
textplot( matrix(letters[1:26], ncol=2) )

### Make a nice 4 way display with two plots and two text summaries

data(iris)
par(mfrow=c(2,2))
plot( Sepal.Length ~ Species, data=iris, border="blue", col="cyan",
     main="Boxplot of Sepal Length by Species")
plotmeans( Sepal.Length ~ Species, data=iris, barwidth=2, connect=FALSE,
          main="Means and 95% Confidence Intervals\nof Sepal Length by Species")

info <- sapply( split(iris$Sepal.Length, iris$Species),
             function(x) round(c(Mean=mean(x), SD=sd(x), N=n.data::nobs(x)),2) )

textplot( info, valign="top" )
title("Sepal Length by Species")

reg <- lm( Sepal.Length ~ Species, data=iris )
textplot( capture.output(summary(reg)), valign="top")
title("Regression of Sepal Length by Species")
```
par(mfrow=c(1,1))

### Show how to control text color
cols <- c("red", "green", "magenta", "forestgreen")
mat <- cbind(name=cols, t(col2rgb(cols)), hex=col2hex(cols))
textplot(mat,
    col.data=matrix(cols, nrow=length(cols), byrow=FALSE, ncol=5),
)

### Show how to manually tune the character size
data(iris)
reg <- lm( Sepal.Length ~ Species, data=iris )
text <- capture.output(summary(reg))

# do the plot and capture the character size used
textplot(text, valign="top")

# see what size was used
cex

# now redo the plot at 80% size
textplot( text, valign="top", cex=cex*0.80)

## End(Not run)

---

**venn**

*Plot a Venn diagram*

---

**Description**

Plot a Venn diagrams for up to 5 sets

**Usage**

```r
venn(data, universe=NA, small=0.7, showSetLogicLabel=FALSE,
     simplify=FALSE, show.plot=TRUE, intersections=TRUE, names,
     ...)
```

## S3 method for class 'venn'

```r
plot(x, y, ..., small=0.7, showSetLogicLabel=FALSE,
     simplify=FALSE)
```
Arguments

data,x Either a list list containing vectors of names or indices of group intersections, or
a data frame containing boolean indicators of group intersectionship (see below)
universe Subset of valid name/index elements. Values ignore values in codedata not in
this list will be ignored. Use NA to use all elements of data (the default).
small Character scaling of the smallest group counts
showSetLogicLabel Logical flag indicating whether the internal group label should be displayed
simplify Logical flag indicating whether unobserved groups should be omitted.
show.plot Logical flag indicating whether the plot should be displayed. If false, simply
returns the group count matrix.
intersections Logical flag indicating if the returned object should have the attribute "individuals.in.intersections" featuring for every set a list of individuals that are assigned
to it.
y Ignored
... Optional graphical parameters.
names Optional vector of group names.

Details

data should be either a named list of vectors containing character string names ("GeneAABBB",
"GeneBBBCY", ... "GeneXXZZ") or indexes of group intersections (1, 2, ... N), or a data frame
containing indicator variables (TRUE, FALSE, TRUE, ..) for group intersectionship. Group names
will be taken from the component list element or column names.

Value

Invisibly returns an object of class "venn", containing:

- A matrix of all possible sets of groups, and the observed count of items belonging to each The
  fist column contains observed counts, subsequent columns contain 0-1 indicators of group
  intersectionship.
- If intersections=TRUE, the attribute intersections will be a list of vectors containing the
  names of the elements belonging to each subset.

Author(s)

Steffen Moeller <steffen\_moeller@gmx.de>, with cleanup and packaging by Gregory R. Warnes
<greg@warnes.net>.

Examples

##
## Example using a list of item names belonging to the
## specified group.
##


## Construct some fake gene names...

```r
oneName <- function() paste(sample(LETTERS, 5), collapse = "")
geneNames <- replicate(1000, oneName())
```

## Example using a list of item indexes belonging to the specified group.

```r
GroupA <- sample(geneNames, 400, replace = FALSE)
GroupB <- sample(geneNames, 750, replace = FALSE)
GroupC <- sample(geneNames, 250, replace = FALSE)
GroupD <- sample(geneNames, 300, replace = FALSE)
input <- list(GroupA, GroupB, GroupC, GroupD)
input

tmp <- venn(input)
attr(tmp, "intersections")
```

## Example using a data frame of indicator ('f'lag) columns

```r
GroupA.f <- geneNames %in% GroupA
GroupB.f <- geneNames %in% GroupB
GroupC.f <- geneNames %in% GroupC
GroupD.f <- geneNames %in% GroupD
input.df <- data.frame(A = GroupA.f, B = GroupB.f, C = GroupC.f, D = GroupD.f)
head(input.df)
venn(input.df)
```

## Smaller set to create empty groupings

```r
csmall <- input.df[1:20,]
```

```r
venn(csmall, simplify = FALSE) # with empty groupings
venn(csmall, simplify = TRUE) # without empty groupings
```

## Capture group counts, but don't plot

```r
tmp <- venn(input, show.plot = FALSE)
tmp
```

## Show internal binary group labels

```r
venn(input, showSetLogicLabel = TRUE)
```
## wapply

### Compute the Value of a Function Over a Local Region Of An X-Y Plot

#### Description

This function applies the specified function to the sets of y values that are defined by overlapping "windows" in the x-dimension. For example, setting \( \text{fun} = \text{mean} \) returns local means, while setting \( \text{fun} = \text{function}(x) \cdot \sqrt{\text{var}(x)} \) returns local estimates of the standard deviation.

#### Usage

\[
\text{wapply}(x, y, \text{fun} = \text{mean}, \text{method} = \text{"range"}, \text{width} = 50, \text{drop.na} = \text{TRUE}, \text{pts}, \ldots)
\]
Arguments

x  vector of x values for (x,y) pairs
y  vector of y values for (x,y) pairs
fun  function to be applied
method  method of defining an x-neighborhood. One of "width", "nobs", "range", or "fraction". See details.
width  width of an x-neighborhood. See details.
n  Number of equally spaced points at which to compute local estimates. See details.
drop.Na  should points which result in missing values NA be omitted from the return value. Defaults to true.
pts  x locations at which to compute the local mean when using the "width" or "range" methods. Ignored otherwise.
...  arguments to be passed to fun

Details

Two basic techniques are available for determining what points fall within the same x-neighborhood. The first technique uses a window with a fixed width in the x-dimension and is is selected by setting method="width" or method="range". For method="width" the width argument is an absolute distance in the x-dimension. For method="range", the width is expressed as a fraction of the x-range. In both cases, pts specifies the points at which evaluation of fun occurs. When pts is omitted, n x values equally spaced along the x range are used.

The second technique uses windows containing k neighboring points. The (x,y) pairs are sorted by the x-values and the nearest k/2 points with higher x values and the k/2 nearest points with lower x values are included in the window. When method="nobs", k equals width (actually 2*floor(width/2)). When method="fraction", width specifies what fraction of the total number of points should be included. The actual number of points included in each window will be floor(n*frac/2)*2. Regardless of the value of pts, the function fun will be evaluated at all x locations.

Value

Returns a list with components

x  x location’

y  Result of applying fun to the window about each x location

Author(s)

Gregory R. Warnes <greg@warnes.net>
Examples

```r
# show local mean and inner 2-sd interval to help diagnose changing mean
# or variance structure
x <- 1:1000
y <- rnorm(1000, mean=1, sd=1 + x/1000)

plot(x,y)
lines(wapply(x,y,mean),col="red")

CL <- function(x,sd) mean(x)+sd*sqrt(var(x))
lines(wapply(x,y,CL,sd=1),col="blue")
lines(wapply(x,y,CL,sd=1),col="blue")
lines(wapply(x,y,CL,sd=2),col="green")
lines(wapply(x,y,CL,sd=2),col="green")

# show local mean and inner 2-sd interval to help diagnose changing mean
# or variance structure
x <- 1:1000
y <- rnorm(1000, mean=x/1000, sd=1)

plot(x,y)
lines(wapply(x,y,mean),col="red")

CL <- function(x,sd) mean(x)+sd*sqrt(var(x))
lines(wapply(x,y,CL,sd=1,method="fraction",width=1/20),col="blue")
lines(wapply(x,y,CL,sd=1,method="fraction",width=1/20),col="blue")
lines(wapply(x,y,CL,sd=2,method="nobs",width=250),col="green")
lines(wapply(x,y,CL,sd=2,method="nobs",width=250),col="green")
```
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