

Package ‘rgr’

January 7, 2016

Type Package

Title Applied Geochemistry EDA

Version 1.1.13

Date 2016-01-05

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Depends MASS, fastICA

Suggests akima

Description Geological Survey of Canada (GSC) functions for exploratory data analysis with applied geochemical data, with special application to the estimation of background ranges and identification of outliers, 'anomalies', to support mineral exploration and environmental studies. Additional functions are provided to support analytical data QA/QC, ANOVA for investigations of field sampling and analytical variability, and utility tasks. NOTE: function caplot() for concentration-area plots employs package 'akima', however, 'akima' is only licensed for not-for-profit use. Therefore, not-for-profit users of 'rgr' will have to independently make package 'akima' available through library(...); and use of function caplot() by for-profit users will fail.

License GPL-2

NeedsCompilation no

Repository CRAN

Date/Publication 2016-01-07 15:40:29

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rgr_1.1.13-package *Geological Survey of Canada Applied Geochemistry EDA*

Description

Functions for Exploratory Data Analysis with applied geochemical data.

Details

This package supports the display and analysis of applied geochemical survey data, particularly in the context of estimating the ranges of background variation due to natural phenomena and the identification of outliers that may be due to natural processes or anthropogenic contamination. Functions are provided for use with univariate and multivariate data, in the latter context tools are provided for compositional, constant sum, data. Additional functions support analytical data QA/QC, ANOVA for investigations of field sampling and analytical variability, and utility tasks. NOTE: function cplot for concentration-area plots employs package akima, however, akima is only licensed for

not-for-profit use. Therefore, not-for-profit users of rgr will have to independently make package akima available through library(akima); and use of function caplot by for-profit users will fail.

Package: rgr_1.1.13
Type: Package
Version: 1.0
Date: 2016-01-05
License: GPL-2

Author(s)

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References

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ad.plot1

Plot Results of Analytical Duplicate Analyses

Description

Function to plot the results of analytical duplicate analyses as the percent absolute difference between duplicates relative to their means, or as the Relative Standard Deviation of the individual duplicate. Classical and robust estimates of the arithmetic Relative Standard Deviation (%) and the mean/median to which they apply are displayed based on the pooled estimate of analytical variability from the duplicates. If the duplicate data span more than 1.5 orders of magnitude this estimate is unreliable due to heterogeneity of variance considerations (heteroscedasticity). The x-axis data may either present the duplicates in the order in which they occur in the data, usually a time-series, as the duplicate means, or as the ratio of the duplicate mean to the lower detection limit if it is provided. Optionally the x-axis may be scaled logarithmically if the range of the data requires. If there is a target acceptance level it may be provided and will be displayed as a red dashed line on the plot. For data stored in alternate forms from that expected by this function use [ad.plot2](#).

Usage

```
ad.plot1(x1, x2, xname = deparse(substitute(x1)), if.order = TRUE,  
if.rsds = FALSE, ldl = NULL, ad.tol = NULL, log = FALSE, ...)
```

Arguments

x1	a column vector from a matrix or data frame, $x1[1], \dots, x1[n]$.
x2	another column vector from the matrix or data frame, $x2[1], \dots, x2[n]$. x1 and x2 must be of identical length, n, where x2 is a duplicate measurement of x1.
xname	a title can be displayed with the plot and results, e.g., <code>xname = "Cu (mg/kg)"</code> . If this field is undefined the character string for x is used as a default.
if.order	by default the analytical duplicate results are plotted in the order in which they occur in the data file, this usually corresponds to date of analysis. Alternately, setting <code>if.order = FALSE</code> results in the results being plotted against their means.
if.rsds	by default the absolute difference between the duplicates expressed as a percentage of their mean is plotted on the y-axis. If it is required to plot the individual duplicate relative standard deviations (RSDs), set <code>if.rsds = TRUE</code> .
ldl	by default the x-axis is defined by the measurement units. If it is desired to express the duplicate means as a ratio to the lower detection limit (ldl) of the analytical procedure, then set <code>ldl = 'ldl'</code> in measurement units.
ad.tol	optionally a tolerance level may be provided for the maximum acceptable percent absolute relative difference between duplicates, in which case a red dotted line is added to the plot. When <code>if.rsds = TRUE</code> the value of <code>ad.tol</code> is interpreted to be the maximum acceptable individual duplicate relative standard deviation.
log	optionally the x-axis of the plot employing duplicate means may be plotted with logarithmic scaling, if so, set <code>log = TRUE</code> .
...	any additional arguments to be passed to the <code>plot</code> function for titling, etc.

Details

If the data are as a single concatenated vector from a matrix or data frame as $x1[1], \dots, x1[n]$ followed by $x[n+1], \dots, x[2n]$, or alternated as $x[1]$ and $x[2]$ being a pair through to $x[2*i+1]$ and $x[2*i+2]$ for the i in $1:n$ duplicate pairs use function [ad.plot2](#).

For examples see [ad.plot2](#) as Geological Survey of Canada National Geochemical Reconnaissance survey data are not stored in this format. This function is present as the graphical equivalent to [anova1](#).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

See Also

[ad.plot2](#), [ad.plot3](#), [ltdl.fix.df](#)

ad.plot2

*Plot Results of Analytical Duplicate Analyses, Alternate Input***Description**

Function to prepare data stored in alternate forms from that expected by function [ad.plot1](#) for its use, for further details see `x` in Arguments below. The data will be plotted as the percent absolute difference between duplicates relative to their accession order or their means.

Usage

```
ad.plot2(x, xname = deparse(substitute(x)), if.order = TRUE,
         if.rsds = FALSE, ldl = NULL, ad.tol = NULL, log = FALSE,
         ifalt = FALSE, ...)
```

Arguments

<code>x</code>	a column vector from a matrix or data frame, <code>x[1]</code> , ..., <code>x[2*n]</code> . The default is that the first <code>n</code> members of the vector are the first measurements and the second <code>n</code> members are the duplicate measurements. If the measurements alternate, i.e. duplicate pair 1 measurement 1 followed by measurement 2, etc., set <code>ifalt = TRUE</code> .
<code>xname</code>	a title can be displayed with the plot and results, e.g., <code>xname = "Cu (mg/kg)"</code> . If this field is undefined the character string for <code>x</code> is used as a default.
<code>if.order</code>	by default the analytical duplicate results are plotted in the order in which they occur in the data file, this usually corresponds to date of analysis in a time-series. Alternately, setting <code>if.order = FALSE</code> results in the individual duplicate results being plotted against their means.
<code>if.rsds</code>	by default the absolute difference between the duplicates expressed as a percentage of their mean is plotted on the y-axis. If it is required to plot the relative standard deviations (RSDs), set <code>if.rsds = TRUE</code> .
<code>ldl</code>	by default the x-axis is defined by the measurement units. If it is desired to express the duplicate means as a ratio to the lower detection limit (ldl) of the analytical procedure, then set <code>ldl = 'ldl'</code> in measurement units.
<code>ad.tol</code>	optionally a tolerance level may be provided for the maximum acceptable percent absolute relative difference between duplicates, in which case a red dotted line is added to the plot.
<code>log</code>	optionally the x-axis of the plot employing duplicate means may be plotted with logarithmic scaling, if so, set <code>log = TRUE</code> .
<code>ifalt</code>	set <code>ifalt = TRUE</code> to accommodate alternating sets of paired observations.
...	any additional arguments to be passed to the plot function for titling, etc.

Details

Data may be as a single concatenated vector from a matrix or data frame as $x1[1], \dots, x1[n]$ followed by $x[n+1], \dots, x[2n]$, or alternated as $x[1]$ and $x[2]$ being a pair through to $x[2*i+1]$ and $x[2*i+2]$ for the i in $1:n$ duplicate pairs, see `ifalt`.

If the data are as n duplicate pairs, $x1$ and $x2$, use function `ad.plot1`.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Author(s)

Robert G. Garrett

See Also

`ad.plot1`, `ad.plot4`, `ltdl.fix.df`

Examples

```
## Make test data available
data(ad.test)
attach(ad.test)

## Plot analytical duplicate analyses as a time-series
ad.plot2(Cu, ifalt = TRUE)

## Plot analytical duplicate analyses versus duplicate means,
## annotating more appropriately, with a 20% maximum tolerance
ad.plot2(Cu, "Cu (mg/kg)", if.order = FALSE, ad.tol = 20, ifalt = TRUE)

## Detach test data
detach(ad.test)
```

ad.plot3

Plot Results of Analytical Duplicate Analyses as Ratios

Description

Function to plot the results of analytical duplicate analyses as ratios vs. their means with logarithmic scaling. Classical and robust estimates for the mean/median and Relative Standard Deviations (%) are displayed. In addition to displaying this information the 95% standard error interval on the mean ratio is estimated and used to test that the mean ratio is not significantly different from one. A situation that would indicate severe systematic drift between the analytical duplicates, something that could happen if the duplicates were analysed in a separate batch at a later date, or were analysed as a group at the beginning or end of an analytical batch. The mean ratio and 95% confidence bounds for the duplicate ratios are plotted as dotted lines on the plot. If the lower detection for the

analyses is defined, and it falls within the range of the duplicate means, it will be plotted as a red vertical dotted line. To facilitate comparison between measurements, or measurements by different methods, the range of the ratio display can be defined. Optionally, the locator may be deployed so that the ratios for estimating the 95% confidence bounds on any value may be placed on the plot in a location of the user's choice. Optionally the ratio plot may be supplemented by an CPP of the ratios, in which case they are presented as side-by-side plots. For data stored in alternate forms from that expected by this function use [ad.plot4](#).

Usage

```
ad.plot3(x1, x2, xname = deparse(substitute(x1)), if.order = FALSE, ad.tol = NULL,
        ldl = NULL, maxrat = NULL, if.text = FALSE, if.cpp = FALSE, ...)
```

Arguments

<code>x1</code>	a column vector from a matrix or data frame, <code>x1[1]</code> , ..., <code>x1[n]</code> .
<code>x2</code>	another column vector, <code>x2[1]</code> , ..., <code>x2[n]</code> . <code>x1</code> and <code>x2</code> must be of identical length, where <code>x2</code> is a duplicate measurement of <code>x1</code> .
<code>xname</code>	a title can be displayed with the plot and results, e.g., <code>xname = "Cu (mg/kg)"</code> . If this field is undefined the character string for <code>x</code> is used as a default.
<code>if.order</code>	by default the analytical duplicate results are plotted against their means. Alternately, setting <code>if.order = TRUE</code> causes the results to be plotted in the order in which they occur in the data file, this usually corresponds to date of analysis.
<code>ad.tol</code>	a tolerance level may be provided for the maximum acceptable percent ratio between duplicates when they are plotted sequentially, i.e. <code>if.order = TRUE</code> , in which case red dotted lines are added to the plot.
<code>ldl</code>	if the lower detection limit (<code>ldl</code>) of the analytical procedure is provided and falls within the range of the duplicate means it will be plotted as a vertical red dotted line.
<code>maxrat</code>	optionally the maximum ratio for the plot axis, the minimum is computed as $1/\text{maxrat}$, may be set. This is useful when making comparisons between elements or between measurement methods and ensures similar scaling. Otherwise <code>maxrat</code> is determined from the data.
<code>if.text</code>	if set <code>if.text = TRUE</code> the locator is deployed so that the ratios for estimating the 95% confidence bounds on any value may be placed on the ratio plot.
<code>if.cpp</code>	if set <code>if.cpp = TRUE</code> the ratio plot is displayed side-by-side with an Normal Cumulative Probability Plot (CPP) of the ratios. In this instance <code>if.text</code> is not an option.
...	any additional arguments to be passed to the <code>plot</code> function for titling, etc. For example, setting <code>col = 4</code> will result in the mean ratio and the 95% bounds on the range of the ratios being plotted in blue.

Details

If the data are as a single concatenated vector from a matrix or data frame as `x1[1]`, ..., `x1[n]` followed by `x[n+1]`, ..., `x[2n]`, or alternated as `x[1]` and `x[2]` being a pair through to `x[2*i+1]` and `x[2*i+2]` for the `i` in $1:n$ duplicate pairs use function [ad.plot4](#).

For examples see [ad.plot4](#) as Geological Survey of Canada National Geochemical Reconnaissance survey data are not stored in this format. This function is present as the graphical equivalent to [anova1](#).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

See Also

[ad.plot4](#), [ad.plot2](#), [ltdl.fix.df](#)

ad.plot4	<i>Plot Results of Analytical Duplicate Analyses as Ratios, Alternate Input</i>
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Description

Function to prepare data stored in alternate forms from that expected by function [ad.plot3](#) for its use, for further details see *x* in Arguments below. The data will be plotted as ratios, i.e. dup1/dup2, with logarithmic scaling relative to their means.

Usage

```
ad.plot4(x, xname = deparse(substitute(x)), if.order = FALSE, ad.tol = NULL,
         ifalt = FALSE, ldl = NULL, maxrat = NULL, if.text = FALSE,
         if.cpp = FALSE, ...)
```

Arguments

<i>x</i>	a column vector from a matrix or data frame, $x[1], \dots, x[2*n]$. The default is that the first <i>n</i> members of the vector are the first measurements and the second <i>n</i> members are the duplicate measurements. If the measurements alternate, i.e. duplicate pair 1 measurement 1 followed by measurement 2, etc., set <code>ifalt = TRUE</code> .
<i>xname</i>	a title can be displayed with the plot and results, e.g., <code>xname = "Cu (mg/kg)"</code> . If this field is undefined the character string for <i>x</i> is used as a default.
<i>if.order</i>	by default the analytical duplicate results are plotted in the order in which they occur in the data file, this usually corresponds to date of analysis. Alternately, setting <code>if.order = FALSE</code> results in the results being plotted against their means.

ad.tol	optionally a tolerance level may be provided for the maximum acceptable percent ratio between duplicates, in which case a red dotted line is added to the plot.
ifalt	set <code>ifalt = TRUE</code> to accommodate alternating sets of paired observations.
ldl	if the lower detection limit (<code>ldl</code>) of the analytical procedure is provided and falls within the range of the duplicate means it will be plotted as a vertical black dotted line. If another colour is required, e.g., red, set <code>col = 2</code> . See function display.lty for the range of available colours.
maxrat	optionally the maximum ratio for the plot axis, the minimum is computed as $1/\text{maxrat}$, may be set. This is useful when making comparisons between elements or between measurement methods and ensures similar scaling. Otherwise <code>maxrat</code> is determined from the data.
if.text	if set <code>if.text = TRUE</code> the locator is deployed so that the ratios for estimating the 95% confidence bounds on any value may be placed on the ratio plot.
if.cpp	if set <code>if.cpp = TRUE</code> the ratio plot is displayed side-by-side with an Normal Cumulative Probability Plot (CPP) of the ratios. In this instance <code>if.text</code> is not an option.
...	any additional arguments to be passed to the <code>plot</code> function for titling, etc.

Details

Data may be as a single concatenated vector from a matrix or data frame as `x1[1], ..., x1[n]` followed by `x[n+1], ..., x[2n]`, or alternated as `x[1]` and `x[2]` being a pair through to `x[2*i+1]` and `x[2*i+2]` for the `i` in `1:n` duplicate pairs, see `ifalt`.

If the data are as `n` duplicate pairs, `x1` and `x2`, use function [ad.plot3](#).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

See Also

[ad.plot3](#), [ad.plot1](#), [ltdl.fix.df](#)

Examples

```
## Make test data available
data(ad.test)
attach(ad.test)

## Plot analytical duplicate analyses as ratios in acquisition sequence
ad.plot4(Cu, ifalt = TRUE)
```

```
## Plot analytical duplicate analysis ratios against their means with
## more informative annotation
ad.plot4(Cu, "Cu (mg/kg)", ifalt = TRUE, if.order = FALSE)

## Plot analytical duplicate analysis ratios against their means with a
## CPP of the ratios and more informative annotation
ad.plot4(Cu, "Cu (mg/kg)", ifalt = TRUE, if.order = FALSE, if.cpp = TRUE)

## Detach test data
detach(ad.test)
```

ad.test

National Geochemical Reconnaissance survey QA/QC data

Description

A subset of analytical duplicate data from NGR surveys undertaken in 2000 and 2001.

Usage

```
data(ad.test)
```

Format

A data frame with 160 observations on the following 2 variables. Unique identifiers are present in the data frame, use `dimnames(ad.test)[[1]]` to access or display them.

RS the Replicate Status code.

Cu the copper determinations, mg/kg.

Details

The Replicate Status code indicates the ‘position’ of the geochemical sample in the QA/QC structure. RS = 8 indicates analytical duplicate, RS = 2 indicates the field duplicate, and RS = 1 indicates a routine regional coverage site that was ‘duplicated’. All other routine regional coverage sites are coded RS = 0. In this file the data record for the analytical duplicate is followed by the data record for the physical sample the duplicate was split from. The analytical duplicate may be split from any field sample, but preferably from one of the field duplicates, permitting a more incisive investigation of sampling and analytical variability using function [gx.triples.aov](#).

Source

Internal Geological Survey of Canada files.

alr *Additive Log-Ratio (alr) transformation*

Description

Undertakes an additive log-ratio transformation to remove the effects of closure in a data matrix.

Usage

```
alr(xx, j = NULL, ifclose = FALSE, ifwarn = TRUE)
```

Arguments

xx	a n by p matrix to be additively log ratioed. It is essential that a single unit of measurement is used. Thus it may be required to convert, for example, determinations in percent to ppm (mg/kg) so that all measurements are in ppm prior to executing this function. Natural logarithms are used.
j	the index number of the element in the range [1:p] to be used as the divisor, j, must be defined, there is no default index.
ifclose	if it is required to close a data set prior to transformation set ifclose = TRUE.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out an additive log-ratio transformation all the data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.

Details

Most analytical chemical data for major, minor and trace elements are of a closed form, i.e. for a physical individual sample they sum to a constant, whether it be percent, ppm (mg/kg), or some other units. It does not matter that only some components contributing to the constant sum are present in the matrix, the data are closed. As a result, as some elements increase in concentration others must decrease, this leads to correlation measures and graphical presentations that do not reflect the true underlying relationships. An additive log-ratio is one procedure for removing closure effects, others are centred log-ratios (*clr*) and isometric log-ratios (*ilr*).

Care should be taken in selecting the variable, index = j, for use as the divisor. Variables lacking sufficient significant figures in their quantification, or variables measured at close to their measurement detection limits, should be avoided.

It is worth noting that when the alr transform is undertaken with a geochemically conservative element selected as the divisor and two elements are then displayed in an x-y plot the result is a Pearce Element Ratio plot (Pearce, 1968) with log scaling.

Value

x	a n by (p-1) matrix of additively log-ratioed values, the j-th column of the matrix being dropped.
---	--

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows containing NAs in the data matrix are removed prior to undertaking the transformation.

Author(s)

Robert G. Garrett

References

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Pearce, T.H., 1968. A contribution to the theory of variation diagrams. *Contributions to Mineralogy and Petrology*, 19(2):142-157.

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See Also

[clr](#), [ilr](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
sind.mat <- as.matrix(sind[, -c(1:3)])

## Undertake alr transform, use Pb [j = 6] as the divisor,
## note necessity of converting percent Fe to mg/kg
sind.mat[, 2] <- sind.mat[, 2] * 10000
temp <- alr(sind.mat, 6)
temp

## Clean-up
rm(sind.mat)
rm(temp)
```

`alts2dups`*Create a Matrix of Duplicate Pairs from Sequential Data*

Description

Function to take data stored as stacked records or alternating rows of records for `ndup` duplicate pairs and generate a matrix with `ndup` rows and two columns for the duplicate data, for further details see `x` in Arguments below. The function returns a matrix for processing.

Usage

```
alts2dups(x, ifalt = FALSE)
```

Arguments

<code>x</code>	a column vector from a matrix or data frame, <code>x[1]</code> , ..., <code>x[2*n]</code> . The default is that the first <code>n</code> members of the vector are the first measurements and the second <code>n</code> members are the duplicate measurements. If the measurements alternate, i.e. duplicate pair 1 measurement 1 followed by measurement 2, etc., set <code>ifalt = TRUE</code> .
<code>ifalt</code>	set <code>ifalt = TRUE</code> to accommodate alternating sets of paired observations.

Details

Function provides an easy procedure with function `plot` to present duplicate data as an x-y plot. The matrix generated can be passed directly to `plot` in its call and the matrix column headings synthesized from the input variable name are used as plot axis titles.

Value

<code>xx</code>	an <code>ndup</code> by 2 matrix containing the duplicate pairs. Row names are sequenced <code>1:ndup</code> and the two column names are generated from the input variable name, <code>x</code> with <code>.1</code> or <code>.2</code> appended, respectively.
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Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

Examples

```
## Make test data available
data(ad.test)
attach(ad.test)

## Save matrix of duplicate pairs
test.save <- alts2dups(Cu, ifalt = TRUE)

## Plot analytical duplicate analyses
plot(alts2dups(Cu, ifalt = TRUE), log = "xy")

## Clean-up and detach test data
rm(test.save)
detach(ad.test)
```

anova1	<i>Analysis of Variance (ANOVA)</i>
--------	-------------------------------------

Description

Undertakes a random effects model Analysis of Variance (ANOVA) on a set of duplicate measurements to determine if the analytical, or combined sampling and analytical, (within) variability is significantly smaller than the variability across the duplicates. For data stored in alternate form use [anova2](#)

Usage

```
anova1(x1, x2, xname = deparse(substitute(x1)), log = FALSE)
```

Arguments

x1	a column vector from a matrix or data frame, x1[1], ..., x1[n].
x2	another column vector from the matrix or data frame, x2[1], ..., x2[n]. x1 and x2 must be of identical length, n, where x2 is a duplicate measurement of x1.
xname	by default the character string for x1 is used for the title. An alternate title can be displayed with xname = "text string", see Examples.
log	if a logarithmic transformation (base 10) of the data is required to meet homogeneity of variance considerations (i.e. severe heteroscedasticity) set log = TRUE. This is also advisable if the range of the observations exceeds 1.5 orders of magnitude.

Details

In field geochemical surveys the combined sampling and analytical variability is more important than analytical variability alone. If the at site (within) variability is not significantly smaller than the between duplicate sites variability it cannot be stated that there are statistically significant spatial patterns in the data, and they are likely not suitable for mapping. This may not mean that the data cannot be used to recognize individuals with above threshold or action level observations. However, under these conditions there also may be above threshold or action level instances that the survey data have failed to detect (Garrett, 1983).

A random effects ANOVA is undertaken, the ANOVA table is displayed, together with estimates of the variance components, i.e. how much of the total variability is between and within the duplicate measurements, and the USGS mapping reliability measures of V and Vm (Miesch et al., 1976). Additionally, the data are investigated through a two-way model following the procedure of Bolviken and Sinding-Larsen (1973).

If the data are as a single concatenated vector from a matrix or data frame as $x1[1], \dots, x1[n]$ followed by $x[n+1], \dots, x[2n]$, or alternated as $x[1]$ and $x[2]$ being a pair through to $x[2*i+1]$ and $x[2*i+2]$, for the i in $1:n$ duplicate pairs use function [anova2](#).

Note

The script does not follow a standard computation of Mean Squares, but is based on a procedure developed after Garrett (1969) for use in the field in the 1970s when pocket calculators first had mean and standard deviation functions.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Duplicate pairs $x1, x2$ containing any NAs are omitted from the calculations.

If a log transformation is undertaken and any less than or equal to zero values occur in the data the function will halt with a warning to that effect.

Author(s)

Robert G. Garrett

References

- Bolviken, B. and Sinding-Larsen, R., 1973. Total error and other criteria in the interpretation of stream sediment data. In *Geochemical Exploration 1972*, Institution of Mining and Metallurgy, London, pp. 285-295.
- Garrett, R.G., 1969. The determination of sampling and analytical errors in exploration geochemistry. *Economic Geology*, 64(4):568-569.
- Garrett, R.G., 1983. Sampling methodology. In Chapter 4 of *Handbook of Exploration Geochemistry*, Vol. 2, Statistics and Data Analysis in Geochemical Prospecting (Ed. R.J. Howarth), Elsevier, pp. 83-110.
- Miesch, A.T. et al., 1976. Geochemical survey of Missouri - methods of sampling, analysis and statistical reduction of data. U.S. Geological Survey Professional Paper 954A, 39 p.

See Also

[anova2, ltdl.fix.df](#)

Examples

```
## Make test data available
data(ms.data1)
attach(ms.data1)

## Undertake an ANOVA for duplicate measurements on rock samples
anova1(MS.1, MS.2, log = TRUE,
xname = "Duplicate measurements of Magnetic Susceptibility")

## Detach test data
detach(ms.data1)
```

anova2

Analysis of Variance (ANOVA), Alternate Input

Description

Function to prepare data stored in alternate forms from that expected by function [anova1](#) for its use. For further details see *x* in Arguments below.

Usage

```
anova2(x, xname = deparse(substitute(x)), log = FALSE, ifalt = FALSE)
```

Arguments

<i>x</i>	a column vector from a matrix or data frame, $x[1], \dots, x[2*n]$. The default is that the first <i>n</i> members of the vector are the first measurements and the second <i>n</i> members are the duplicate measurements. If the measurements alternate, i.e. duplicate pair 1 measurement 1 followed by measurement 2, etc., set <code>ifalt = TRUE</code> .
<i>xname</i>	by default the character string for <i>x</i> is used for the title. An alternate title can be displayed with <code>xname = "text string"</code> , see Examples.
<i>log</i>	if a logarithmic transformation (base 10) of the data is required to meet homogeneity of variance considerations (i.e. severe heteroscedasticity) set <code>log = TRUE</code> . This is also advisable if the range of the observations exceeds 1.5 orders of magnitude.
<i>ifalt</i>	set <code>ifalt = TRUE</code> to accommodate alternating sets of paired observations.

Details

For further details see [anova1](#).

If the data are as *n* duplicate pairs, *x1* and *x2*, use function [anova1](#).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

See Also

[anova1](#), [ltdl.fix.df](#)

Examples

```
## Make test data available
data(ms.data2)
attach(ms.data2)

## Undertake an ANOVA for duplicate measurements on rock samples
anova2(MS, log = TRUE,
xname = "Duplicate measurements of Magnetic Susceptibility")

## Detach test data
detach(ms.data2)

## Make test data available
data(ms.data3)
attach(ms.data3)

## Undertake an ANOVA for duplicate measurements on rock samples
anova2(MS, log = TRUE, ifalt = TRUE,
xname = "Duplicate measurements of Magnetic Susceptibility")

## Detach test data
detach(ms.data3)
```

bwplots

Plot Vertical Box-and-Whisker Plots

Description

Plots a series of vertical box-and-whisker plots (Garrett, 1988) where the individual plots represent the data subdivided by the value of some factor. Optionally the y-axis may be scaled logarithmically (base 10). A variety of other plot options are available, see Details and Note below.

Usage

```
bwplots(x, by, log = FALSE, wend = 0.05, notch = TRUE, xlab = "",
        ylab = deparse(substitute(x)), ylim = NULL, main = "",
        label = NULL, plot.order = NULL, xpos = NA, width,
        space = 0.25, las = 1, cex.axis = 1, adj = 0.5, add = FALSE,
        ssl1 = 1, colr = 8, pch = 3, ...)
```

Arguments

<code>x</code>	name of the variable to be plotted.
<code>by</code>	the name of the factor variable to be used to subdivide the data. See Details below for when <code>by</code> is undefined.
<code>log</code>	to display the data with logarithmic (y-axis) scaling, set <code>log = TRUE</code> .
<code>wend</code>	the locations of the whisker-ends have to be defined. By default these are at the 5th and 95th percentiles of the data. Setting <code>wend = 0.02</code> plots the whisker ends at the 2nd and 98th percentiles.
<code>notch</code>	determines if the boxplots are to be “notched” such that the notches indicate the 95% confidence intervals for the medians. The default is to notch the boxplots, to suppress the notches set <code>notch = FALSE</code> . See Details below.
<code>xlab</code>	a title for the x-axis, by default none is provided. A title may be provided, see Examples.
<code>ylab</code>	by default the character string for <code>x</code> is used for the y-axis title. An alternate title can be displayed with <code>ylab = "text string"</code> , see Examples.
<code>ylim</code>	only for <code>log = FALSE</code> , defines the limits of the y-axis if the default limits based on the range of the data are unsatisfactory. It can be used to ensure the y-axis scaling in multiple sets of boxplots are the same to facilitate visual comparison.
<code>main</code>	a main title may be added optionally above the display by setting <code>main</code> , e.g., <code>main = "Kola Project, 1995"</code> .
<code>label</code>	by default the character strings defining the factors are used to label the boxplots along the x-axis. Alternate labels can be provided with <code>label = c("Alt1", "Alt2", "Alt3")</code> , see Examples.
<code>plot.order</code>	provides an alternate order for the boxplots. Thus, <code>plot.order = c(2, 1, 3)</code> will plot the 2nd ordered factor in the 1st position, the 1st in the 2nd, and the 3rd in its 3rd ordered position, see Details and Examples below.
<code>xpos</code>	the locations along the x-axis for the individual vertical boxplots to be plotted. By default this is set to <code>NA</code> , which causes default equally spaced positions to be used, i.e. boxplot 1 plots at value 1 on the x-axis, boxplot 2 at value 2, etc., up to boxplot “n” at value “n”. See Details below for defining <code>xpos</code> .
<code>width</code>	the width of the boxes, by default this is set to the minimum distance between all adjacent boxplots times the value of <code>space</code> . With the default values of <code>xpos</code> this results in a minimum difference of 1, and with the default of <code>space = 0.25</code> the width is computed as 0.25. To specify different widths for all boxplots use, for example, <code>width = c(0.3)</code> . See Details below for changing individual boxplot widths.

<code>space</code>	the space between the individual boxplots, by default this is 0.25 x-axis units.
<code>las</code>	controls whether the x-axis labels are written parallel to the x-axis, the default <code>las = 1</code> , or are written down from the x-axis by setting <code>las = 2</code> . See also, Details below.
<code>cex.axis</code>	controls the size of the font used for the factor labels plotted along the x-axis. By default this is 1, however, if the labels are long it is sometimes necessary to use a smaller font, for example <code>cex.axis = 0.8</code> results in a font 80% of normal size.
<code>adj</code>	controls justification of the x-axis labels. By default they are centred, <code>adj = 0.5</code> , to left justify them if the labels are written downwards set <code>adj = 0</code> .
<code>add</code>	permits the user to plot additional boxplots into an existing display. It is recommended that this option is left as <code>add = FALSE</code> .
<code>ss11</code>	determines the minimum data subset size for which a subset will be plotted. By default this is set to 1, which leads to only a circle with a median bar being plotted, as the subset size increases additional features of the box-and-whisker plot are displayed. If <code>ss11</code> results in subset boxplots not being plotted, a gap is left and the factor label is still plotted on the x-axis.
<code>colr</code>	by default the boxes are infilled in grey, <code>colr = 8</code> . If no infill is required, set <code>colr = 0</code> . See <code>display.lty</code> for the range of available colours.
<code>pch</code>	by default the plotting symbol for the subset maxima and minima are set to a plus, <code>pch = 3</code> , alternate plotting symbols may be chosen from those displayed by <code>display.marks</code> .
<code>...</code>	further arguments to be passed to methods. For example, the size of the axis titles by setting <code>cex.lab</code> , and the size of the plot title by setting <code>cex.main</code> . For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

There are two ways to execute this function. Firstly by defining `x` and `by`, and secondly by combining the two variables with the `split` function. See the first two examples below. The `split` function can be useful if the factors to use in the boxplot are to be generated at run-time, see the last example below. Note that when the `split` construct is used instead of by the whole `split` statement will be displayed as the default y-axis title. Also note that when using by the subsets are listed in the order that the factors are encountered in the data, but when using `split` the subsets are listed alphabetically. In either case they can be re-ordered using `plot.order`, see Examples.

In a box-and-whisker plot there are two special cases. When `wend = 0` the whiskers extend to the observed minima and maxima that are not plotted with the plus symbol. When `wend = 0.25` no whiskers or the data minima and maxima are plotted, only the medians and boxes representing the span of the middle 50% of the data are displayed.

The `width` option can be used to define different widths for the individual boxplots. For example, the widths could be scaled to be proportional to the subset population sizes as some function of the square root (`const * sqrt(n)`) or logarithm (`const * log10(n)`) of those sizes (`n`). The constant, `const`, would need to be chosen so that on average the width of the individual boxes would be approximately 0.25, see Example below. It may be desirable for cosmetic purposes to adjust the positions of the boxes along the x-axis, this can be achieved by specifying `xpos`.

Long subset (factor) names can lead to display problems, changing the `las` parameter from its default of `las = 1` which plots subset labels parallel to the axis to `las = 2`, to plot perpendicular to the axis, can help. It may also help to use `label` and split the character string into two lines, e.g., by changing the string "Granodiorite" that was supplied to replace the coded factor variable GRDR to "Grano-\ndiorite". If this, or setting `las = 2`, causes a conflict with the x-axis title, if one is needed, the title can be moved down a line by using `xlab = "\nLithological Units"`. In both cases the `\n` forces the following text to be placed on the next lower line.

If there are more than 7 labels (subsets) and no alternate labels are provided `las` is set to 2, otherwise some labels may fail to be displayed.

The notches in the boxplots indicate the 95% confidence intervals for the medians and can extend beyond the upper and lower limits of the boxes indicating the middle 50% of the data when subset population sizes are small. The confidence intervals are estimated using the binomial theorem. It can be argued that for small populations a normal approximation would be better. However, it was decided to remain with a non-parametric estimate to be consistent with the use of non-parametric statistics in this display.

Note

This function is based on a script shared by Doug Nychka on S-News, April 28, 1992.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to preparing the boxplots.

For summary statistics displays to complement the graphics see [gx.summary.groups](#) or [framework.summary](#).

Author(s)

Douglas W. Nychka and Robert G. Garrett

References

Garrett, R.G., 1988. IDEAS - An Interactive Computer Graphics Tool to Assist the Exploration Geochemist. In Current Research Part F, Geological Survey of Canada Paper 88-1F, pp. 1-13 for a description of box-and-whisker plots.

See Also

[cat2list](#), [ltdl.fix.df](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## Display a simple box-and-whisker plot
bwplots(Cu, by = COUNTRY)
bwplots(split(Cu,COUNTRY))
```

```

## Display a more appropriately labelled and scaled box-and-whisker plot
bwplots(Cu, by = COUNTRY, log = TRUE, xlab = "Country",
ylab = "Cu (mg/kg) in <2 mm C-horizon soil")

## Display a west-to-east re-ordered plot using the full country names
bwplots(split(Cu, COUNTRY), log = TRUE,
ylab = "Cu (mg/kg) in <2 mm C-horizon soil",
label = c("Finland", "Norway", "Russia"),
plot.order = c(2, 1, 3))

## Detach test data
detach(kola.c)

## Make test data kola.o available, setting a -9999, indicating a
## missing pH measurement, to NA
data(kola.o)
kola.o.fixed <- ltdl.fix.df(kola.o, coded = -9999)
attach(kola.o.fixed)

## Display relationship between pH in one pH unit intervals and Cu in
## 0-horizon (humus) soil, extending the whiskers to the 2nd and 98th
## percentiles
bwplots(split(Cu, trunc(pH+0.5)), log = TRUE, wend = 0.02,
xlab = "0-horizon soil pH to the nearest pH unit",
ylab = "Cu (mg/kg) in <2 mm Kola 0-horizon soil")

## As above, but demonstrating the use of variable box widths and the
## suppression of 95% confidence interval notches. The box widths are
## computed as (Log10(n)+0.1)/5, the 0.1 is added as one subset has a
## population of 1. Note: paste is used in constructing xlab, below,
## as the label is long and overflows the text line length
table(trunc(pH+0.5))
bwplots(split(Cu, trunc(pH+0.5)), log=TRUE, wend = 0.02, notch = FALSE,
xlab = paste("0-horizon soil pH to the nearest pH unit,",
"\nbox widths proportional to Log(subset_size)"),
ylab = "Cu (mg/kg) in <2 mm Kola 0-horizon soil",
width = c(0.26, 0.58, 0.24, 0.02))

## Detach test data
detach(kola.o.fixed)

```

bwplots.by.var
Plot Vertical Box-and-Whisker Plots for Variables

Description

Plots a series of vertical box-and-whisker plots (Garrett, 1988) where the individual plots represent the data subdivided by variables. Optionally the y-axis may be scaled logarithmically (base 10). A variety of other plot options are available, see Details and Note below.

Usage

```
bwplots.by.var(xmat, log = FALSE, wend = 0.05, notch = FALSE,
  xlab = "Measured Variables", ylab = "Reported Values",
  main = "", label = NULL, plot.order = NULL, xpos = NA,
  las = 1, cex.axis = 1, adj = 0.5, colr = 8, pch = 3, ...)
```

Arguments

xmat	the data matrix or data frame containing the data (variables).
log	to display the data with logarithmic (y-axis) scaling, set log = TRUE.
wend	the locations of the whisker-ends has to be defined. By default these are at the 5th and 95th percentiles of the data. Setting wend = 0.02 plots the whisker ends at the 2nd and 98th percentiles. See Details below.
notch	determines if the boxplots are to be “notched” such that the notches indicate the 95% confidence intervals for the medians. The default is not to notch the boxplots, to have notches set notch = TRUE.
xlab	a title for the x-axis, by default xlab = “Measured Variables”.
ylab	a title for the y-axis, by default ylab = “Reported Values”, alternate titling may be provided, see Examples.
main	a main title may be added optionally above the display by setting main, e.g., main = “Kola Project, 1995”.
label	by default the character strings defining the variables are used to label the boxplots along the x-axis. Alternate labels can be provided with label = c(“Alt1”, “Alt2”, “Alt3”), see Examples.
plot.order	provides an alternate order for the boxplots. By default the boxplots are plotted in alphabetical order of the factor variables. Thus, plot.order = c(2, 1, 3) will plot the 2nd alphabetically ordered factor in the 1st position, the 1st in the 2nd, and the 3rd in its alphabetically 3rd ordered position.
xpos	the locations along the x-axis for the individual vertical boxplots to be plotted. By default this is set to NA, which causes default equally spaced positions to be used, i.e. boxplot 1 plots at value 1 on the x-axis, boxplot 2 at value 2, etc., up to boxplot “n” at value “n”. See Details below for defining xpos.
las	controls whether the x-axis labels are written parallel to the x-axis, the default las = 1, or are written down from the x-axis by setting las = 2. See also, Details below.
cex.axis	controls the size of the font used for the factor labels plotted along the x-axis. By default this is 1, however, if the labels are long it is sometimes necessary to use a smaller font, for example cex.axis = 0.8 results in a font 80% of normal size.
adj	controls justification of the x-axis labels. By default they are centred, adj = 0.5, to left justify them if the labels are written downwards set adj = 0.
colr	by default the boxes are infilled in grey, colr = 8 . If no infill is required, set colr = 0. See display.lty for the range of available colours.

`pch` by default the plotting symbol for the subset maxima and minima are set to a plus, `pch = 3`, alternate plotting symbols may be chosen from those displayed by `display.marks`.

`...` further arguments to be passed to methods. For example, the size of the axis titles by setting `cex.lab`, and the size of the plot title by setting `cex.main`. For example, if it is required to make the plot title smaller, add `cex.main = 0.9` to reduce the font size by 10%.

Details

There are two ways to provide data to this function. Firstly, if all the variables in a data frame are to be displayed, and there are no factor variables, the data frame name can be entered for `xmat`. However, if there are factor variables, or only a subset of the variables are to be displayed, the data are entered via the `cbind` construct, see Examples below.

In a box-and-whisker plot there are two special cases. When `wend = 0` the whiskers extend to the observed minima and maxima that are not plotted with the plus symbol. When `wend = 0.25` no whiskers or the data minima and maxima are plotted, only the medians and boxes representing the span of the middle 50% of the data are displayed.

Long variable names can lead to display problems, changing the `las` parameter from its default of `las = 1` which plots subset labels parallel to the axis to `las = 2`, to plot perpendicular to the axis, can help. It may also help to use `label` and split the character string into two lines, e.g., by changing the string "Specific Conductivity" that was supplied to replace the variable name `SC` to "Specific\nConductivity". If this, or setting `las = 2`, causes a conflict with the x-axis title, if one is needed, the title can be moved down a line by using `xlab = "\nPhysical soil properties"`. In both cases the `\n` forces the following text to be placed on the next lower line.

If there are more than 7 labels (variables) and no alternate labels are provided `las` is set to 2, otherwise some variable names may fail to be displayed.

The notches in the boxplots indicate the 95% confidence intervals for the medians and can extend beyond the upper and lower limits of the boxes indicating the middle 50% of the data when subset population sizes are small. The confidence intervals are estimated using the binomial theorem. It can be argued that for small populations a normal approximation would be better. However, it was decided to remain with a non-parametric estimate to be consistent with the use of non-parametric statistics in this display.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Any NAs in the data vectors are removed prior to preparing the boxplots.

For a summary statistics display to complement the graphics see `gx.summary.mat`.

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1988. IDEAS - An Interactive Computer Graphics Tool to Assist the Exploration Geochemist. In Current Research Part F, Geological Survey of Canada Paper 88-1F, pp. 1-13 for a description of box-and-whisker plots.

See Also

[bwplots](#), [var2fact](#), [ltdl.fix.df](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## Display a simple box-and-whisker plot for measured variables
bwplots.by.var(cbind(Co,Cu,Ni))

## Display a more appropriately labelled and scaled box-and-whisker plot
bwplots.by.var(cbind(Co,Cu,Ni), log = TRUE,
ylab = "Levels (mg/kg) in <2 mm Kola C-horizon soil")

## Detach test data
detach(kola.c)

## Make test data available
data(ms.data1)
attach(ms.data1)

## Display variables in a data frame extending the whiskers to the
## 2nd and 98th percentiles of the data, remembering to omit the
## sample IDs
bwplots.by.var(ms.data1[, -1], log = TRUE, wend = 0.02)

## Detach test data
detach(ms.data1)
```

bxplot

Plot a Horizontal Boxplot or Box-and-Whisker Plot

Description

Plots a single horizontal boxplot as part of the multi-panel display provided by function [shape](#), the default is a Tukey boxplot, alternately a box-and-whisker plot (Garrett, 1988) may be displayed. Optionally the x-axis may be scaled logarithmically (base 10).

Usage

```
bxplot(xx, xlab = deparse(substitute(xx)), log = FALSE,
       ifbw = FALSE, wend = 0.05, xlim = NULL, main = "", ifn = TRUE,
       colr = 8, cex = 1, ...)
```

Arguments

<code>xx</code>	name of the variable to be plotted.
<code>xlab</code>	by default the character string for <code>xx</code> is used for the x-axis title. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
<code>log</code>	to display the data with logarithmic (x-axis) scaling, set <code>log = TRUE</code> .
<code>ifbw</code>	the default is to plot a horizontal Tukey boxplot, if a box-and-whisker plot is required set <code>ifbw = TRUE</code> .
<code>wend</code>	if <code>ifbw = TRUE</code> the locations of the whisker-ends have to be defined. By default these are at the 5th and 95th percentiles of the data. Setting <code>wend = 0.02</code> plots the whisker ends at the 2nd and 98th percentiles.
<code>xlim</code>	when used in the <code>shape</code> function, <code>xlim</code> is determined by <code>gx.hist</code> and used to ensure all four panels in <code>shape</code> have the same x-axis scaling. However when used stand-alone the limits may be user-defined by setting <code>xlim</code> , see Note below.
<code>main</code>	when used stand-alone a title may be added optionally above the plot by setting <code>main</code> , e.g., <code>main = "Kola Project, 1995"</code> .
<code>ifn</code>	an internal 'switch' set <code>FALSE</code> to suppress the addition of the sample size to the plot.
<code>colr</code>	by default the box is infilled in grey, <code>colr = 8</code> . If no infill is required, set <code>colr = 0</code> . See <code>display.lty</code> for the range of available colours.
<code>cex</code>	by default the size of the text for data set size, <code>N</code> , is set to 80%, i.e. <code>cex = 0.8</code> , and may be changed if required.
<code>...</code>	further arguments to be passed to methods. For example, the size of the axis scale annotation can be change by setting <code>cex.axis</code> , the size of the axis titles by setting <code>cex.lab</code> , and the size of the plot title by setting <code>cex.main</code> . For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

The function can be used stand-alone, but as Tukey boxplots and box-and-whisker plots are usually used to compare the distributions of data subsets the functions `tbplots` (Tukey boxplots) and `bwplots` (box-and-whisker plots) are required for that purpose.

When the boxplot is displayed on a logarithmically scaled x-axis, the data are log transformed prior to the computation of the positions of the fences used in the Tukey boxplot to identify near and far outliers, plotted as plusses and circles, respectively.

In a box-and-whisker plot there are two special cases. When `wend = 0` the whiskers extend to the observed minima and maxima that are not plotted with the plus symbol. When `wend = 0.25` no whiskers or the data minimum and maximum are plotted, only the median and box representing the span of the middle 50 percent of the data are displayed.

Note

Any less than detection limit values represented by negative values, or zeros or numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plot.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range a truncated plot will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or `xx` may be defined in the function call as, for example, `Cu[Cu < some.value]` which would remove the influence of one or more outliers having values greater than `some.value`. In this case the number of data values displayed will be the number that are `<some.value`.

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1988. IDEAS - An Interactive Computer Graphics Tool to Assist the Exploration Geochemist. In Current Research Part F, Geological Survey of Canada Paper 88-1F, pp. 1-13 for a description of box-and-whisker plots.

See Also

[shape](#), [display.lty](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Display a simple boxplot
bxplot(Cu)

## Display a more appropriately labelled and scaled boxplot
bxplot(Cu, xlab = "Cu (mg/kg) in <2 mm Kola 0-horizon soil", log = TRUE)

## Display a box-and-whisker plot with whiskers ending at the 2nd and
## 98th percentiles
bxplot(Cu, xlab = "Cu (mg/kg) in <2 mm Kola 0-horizon soil", ifbw = TRUE,
wend = 0.02, log = TRUE)

## Detach test data
detach(kola.o)
```

caplot

*Prepare a Concentration-Area (C-A) Plot***Description**

Displays a concentration-area (C-A) plot to assess whether the data are spatially multi-fractal (Cheng et al., 1994; Cheng and Agterberg, 1995) as a part of a four panel display. This procedure is useful for determining if multiple populations that are spatially dependent are present in a data set. It can be used to determine the practical limits, upper or lower bounds, of the influence of the biogeochemical processes behind the spatial distribution of the data. Optionally the data may be logarithmically transformed (base 10) prior to interpolation, the points may be ‘jittered’ (see Arguments below), the size of the interpolated grid may be modified, and alternate colour schemes can be chosen for display of the interpolated data.

Usage

```
caplot(x, y, z, zname = deparse(substitute(z)), caname = NULL,
       log = TRUE, ifjit = FALSE, ifrev = FALSE, ngrid = 100,
       colr = topo.colors(16), xcoord = "Easting", ycoord = "Northing")
```

Arguments

x	name of the x-axis spatial coordinate, the eastings.
y	name of the y-axis spatial coordinate, the northings.
z	name of the variable to be processed and plotted.
zname	by default the character string for z is used for the titles of the x-axes of the CPP (Cumulative Normal Percentage Probability) and C-A plot panels. An alternate title can be displayed with <code>zname = "text string"</code> , see Examples.
caname	a title for the image of the interpolated data, by default the character string for z, suitably modified if the z data are log10 transformed. It is often desirable to replace the default title of the input variable name text string with a more informative title, e.g., <code>caname = "Cu (mg/kg) in <2 mm 0-horizon soil"</code> . For no title, set <code>caname = ""</code> .
log	the default is set to <code>log = TRUE</code> as in most cases this function is used with data from a ‘closed’ composition or are positively skewed, where a logarithmic data transform is appropriate. If it is required to undertake the C-A plot interpolation without a logarithmic data transformation, set <code>log = FALSE</code> . This also results in the accompanying probability (CPP) plots being arithmically scaled (x-axes).
ifjit	if there is a possibility that the data set contains multiple measurements at an identical spatial (x,y) location set <code>ifjit = TRUE</code> . The presence of multiple data at an identical location will cause the Akima (1996) interpolation function to fail.
ifrev	by default the empirical C-A function is plotted from highest value to lowest, <code>ifrev = FALSE</code> . As the C-A plot is a log-log display this provides greater

detail for the highest values. The direction of accumulation can be key in detecting multi-fractal patterns, it is usually informative to also prepare a plot with `ifrev = TRUE`, i.e. accumulation from lowest to highest values. To see a dramatic example of this, run the Examples below.

<code>ngrid</code>	by default <code>ngrid = 100</code> , this results in the data being interpolated into a 100 x 100 grid that extends between the data set's spatial extremes determined for the (x,y) spatial coordinates for the data. See Details below.
<code>colr</code>	by default the <code>topo.colors(16)</code> palette is used to render the interpolated grid as an image. For alternative palettes see colors , and see Details below.
<code>xcoord</code>	a title for the x-axis, defaults to <code>Easting</code> .
<code>ycoord</code>	a title for the y-axis, defaults to <code>Northing</code> .

Details

The function creates a four panel display. The percentage cumulative probability (CPP) plot of the data in the upper left, and the CPP plot of the interpolated data to be used in the C-A plot in the upper right. The lower left panel contains an image of the interpolated data, and the lower right the C-A plot.

Akima's (1978, 1996) interpolation function is used to obtain a linear interpolation between the spatial data values. If the data are from a 'closed' composition or are positively skewed the use of a logarithmic data transformation, `log = TRUE`, is highly recommended, as noted above this is commonly the case and is the default. Following generation of the interpolated grid and prior to further processing the interpolated grid values are clipped by the convex-hull of the spatial locations, therefore there is no interpolation beyond the spatial extent, support, of the data is displayed.

The use of the `topo.colors(16)` palette to display the image of the interpolated values leads to low values being plotted in blue, and as the interpolated values increase they take on green, yellow and orange colors. For a grey-scale display for black-and-white use set `colr = grey(0:8/8)`. This leads to lowest interpolated values being plotted in black and the highest in white, using `colr = grey(8:0/8)` reverses this, with the lowest values being plotted in white and the highest in black. In either case, if the values plotted in white occur at the study area boundary, i.e. at the convex hull, the difference between no data and white cannot be discerned.

For preparation of the C-A plot the ordered vector of interpolated values is used as a surrogate for the measurement of area greater than, or less than, a stated interpolated value. The cumulative percentage count of the interpolated values being plotted on the y-axis of the C-A plot. As noted above, it is both informative and important to display the C-A plot accumulated both upwards and downwards.

Note

This wrapper function was developed from a S-Plus function to prepare C-A plots using Akima's (1978, 1996) interpolation procedure written by Graeme Bonham-Carter, Geological Survey of Canada, in April 2004.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any sites with NAs in their (x,y,z) data are removed prior to spatial interpolation and preparation of the C-A plot.

In some R installations the generation of multi-panel displays and the use of function `eqscplot` from package `MASS` causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering options (`warn = -1`) on the R command line, or that line may be included in a first function prepared by the user for loading the 'rgr' package, etc.

Author(s)

Graham F. Bonham-Carter and Robert G. Garrett

References

Akima, H. (1978). A Method of Bivariate Interpolation and Smooth Surface Fitting for Irregularly Distributed Data Points. *ACM Transactions on Mathematical Software*, 4(2):148-164.

Akima, H. (1996). Algorithm 761: scattered-data surface fitting that has the accuracy of a cubic polynomial. *ACM Transactions on Mathematical Software*, 22(3):362-371.

Cheng, Q. and Agterberg, F.P., 1995. Multifractal modeling and spatial point processes. *Mathematical Geology*, 27(7):831-845.

Cheng, Q., Agterberg, F.P. and Ballantyne, S.B., 1994. The separation of geochemical anomalies from background by fractal methods. *Journal of Geochemical Exploration*, 51(2):109-130.

See Also

[cnpplt](#), [interp](#), [gx.mf](#), [shape.alt](#), [colors](#), [ltdl.fix.df](#)

Examples

```
## The following examples are commented out as package akima is not
## automatically made available as it is only a suggest, not a depends,
## and therefore caplot fails when the examples are run during package
## checking and building. Functions shape.alt and/or gx.mf also be used
## to study mulifractality, and these have no restrictions

## Make test data available
## data(kola.o)
## attach(kola.o)

## A default (uninformative) C-A plot
## caplot(UTME/1000, UTMN/1000, Cu)

## Plot a more appropriately scaled (log transformed data) and
## titled display
## caplot(UTME/1000, UTMN/1000, Cu, log = TRUE,
## zname = "Cu (mg/kg) in\n<2 mm 0-horizon soil",
## caname = "Kola Project, 1995\nCu (mg/kg) in <2 mm 0-horizon soil")

## Plot as above but with the C-A plot accumulation reversed
## caplot(UTME/1000, UTMN/1000, Cu, log = TRUE, ifrev = TRUE,
## zname = "Cu (mg/kg) in\n<2 mm 0-horizon soil",
```



```
## caname = "Kola Project, 1995\nCu (mg/kg) in <2 mm 0-horizon soil")  
  
## Detach test data  
## detach(kola.o)
```

cat2list

Divides Data into Subsets by Factor

Description

Converts data into a list form where data are grouped together by factor. Achieves the same objective as the base function [split](#).

Usage

```
cat2list(x, a)
```

Arguments

x name of the data variable to be processed.
a name of the factor variable by which the data are to be split.

Value

data a list containing factors as columns and the values for those factors as rows. The order of the resulting groups, subsets, is the order in which the factor variable names were encountered in parameter a passed to the function.

Note

This function is called by functions [tbplots](#) and [bwplots](#) to prepare Tukey boxplots and box-and-whisker plots, respectively. It is an integral part of the script shared by Doug Nychka on S-News, April 28, 1992. As such it may pre-date the time that [split](#) was added to the S-Plus library.

If by is undefined in the calling functions, [tbplots](#) and [bwplots](#), the same result may be achieved by using the `split(x, a)` construct instead of stating x as the variable to be displayed as boxplots. In which case the data are grouped, subsetted, in alphabetical order of factor variable names.

Author(s)

Douglas W. Nychka

clr *Centred Log-Ratio (clr) transformation*

Description

Undertakes a centred log-ratio transformation to remove the effects of closure in a data matrix.

Usage

```
clr(xx, ifclose = FALSE, ifwarn = TRUE)
```

Arguments

xx	a n by p matrix to be log centred. It is essential that a single unit of measurement is used. Thus it may be required to convert, for example, determinations in percent to ppm (mg/kg) so that all measurements are in ppm prior to executing this function. Natural logarithms are used.
ifclose	if it is required to close a data set prior to transformation set ifclose = TRUE.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out a centred log-ratio transformation all the data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.

Details

Most analytical chemical data for major, minor and trace elements are of a closed form, i.e. for a physical individual sample they sum to a constant, whether it be percent, ppm (mg/kg), or some other units. It does not matter that only some components contributing to the constant sum are present in the matrix, the data are closed. As a result, as some elements increase in concentration others must decrease, this leads to correlation measures and graphical presentations that do not reflect the true underlying relationships. A centred log-ratio is one procedure for removing closure effects, others are additive log-ratios ([alr](#)) and isometric log-ratios ([ilr](#)).

Value

x a n by p matrix of log-centred values.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows containing NAs in the data matrix are removed prior to undertaking the transformation.

The clr transform is suitable for the study of correlation coefficients and subsequent multivariate data analyses. However, for the calculation of Mahalanobis distances, which require matrix inversion, [ilr](#) should be used. Furthermore, in some cases it is preferable to use an [ilr](#) transform prior to undertaking a Principal Component or Factor Analysis, however, a [clr](#) transform is often sufficient.

The `ifclose` option can be useful if a petrochemical ternary system is under investigation. A data subset for a ternary system may be closed and transformed for investigation in x-y plots and comparison with the inferences that may be drawn from a classical ternary diagram display. Ternary plots are not included in this release of 'rgr', their use is discouraged as they do not reveal the true inter-component relationships. However, their use as classification tools is acknowledged where a user's data may be compared to data for known rock types and processes, etc. R users interested in ternary and classification diagrams rather than exploratory data analysis should investigate GCDkit (ver 2.3, R 2.7.0 2008/05/11) by Janousek, Farrow, Erban and Smid. See also Janousek et al. (2006).

Author(s)

Robert G. Garrett

References

- Aitchison, J., 1984. The statistical analysis of geochemical compositions. *Mathematical Geology*, 16(6):531-564.
- Aitchison, J., 1986. *The Statistical Analysis of Compositional data*. Chapman and Hall, London, U.K., 416 p.
- Aitchison, J. and Egozcue, J.J., 2005. Compositional data analysis; where are we and where should we be heading. *Mathematical Geology*, 37(7):829-850.
- Buccianti, A., Mateu-Figueras, G, and Pawlowsky-Glahn, V. (eds.), 2006. *Compositional data analysis in the geosciences: from theory to practice*. The Geological Society Publishing House, Bath, U.K. Special Publication 264, 224 p.
- Janousek, V., Farrow, C.M. and Erban, V., 2006. Interpretation of whole-rock geochemical data in igneous geochemistry introducing Geochemical Data Toolkit (GCDkit). *Journal of Petrology*, 47(6):1255-1259.
- Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. *Statistical Data Analysis Explained: Applied Environmental Statistics with R*. Wiley, 362 p.

See Also

[clr](#), [ilr](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
sind.mat <- as.matrix(sind[, -c(1:3)])

## Undertake clr transform, note necessity
## of converting percent Fe to mg/kg
sind.mat[, 2] <- sind.mat[, 2] * 10000
temp <- clr(sind.mat)
temp

## Clean-up and detach test data
rm(sind.mat)
rm(temp)
```

cnpplt

*Cumulative Normal Percentage Probability (CPP) Plot***Description**

Displays a cumulative normal percentage probability (CPP) plot, equivalent to a Q-Q plot, as has been traditionally used by physical scientists and engineers.

Usage

```
cnpplt(xx, xlab = deparse(substitute(xx)),
       ylab = "% Cumulative Probability", log = FALSE, xlim = NULL,
       main = "", ifqs = FALSE, ifshape = FALSE, pch = 3,
       cex = 0.8, cexp = 1, cex.axis = 0.8, ...)
```

Arguments

<code>xx</code>	name of the variable to be plotted.
<code>xlab</code>	by default the character string for <code>xx</code> is used for the x-axis title. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
<code>ylab</code>	a title for the y-axis, defaults to "% Cumulative Probability".
<code>log</code>	to display the data with logarithmic (x-axis) scaling, set <code>log = TRUE</code> .
<code>xlim</code>	when used in the <code>shape</code> function, <code>xlim</code> is determined by function <code>gx.hist</code> and used to ensure all four panels in <code>shape</code> have the same x-axis scaling. However, when used stand-alone the limits may be user-defined by setting <code>xlim</code> , see Details below.
<code>main</code>	when used stand-alone a title may be added optionally above the plot by setting <code>main</code> , e.g., <code>main = "Kola Ecogeochemistry Project, 1995"</code> .
<code>ifqs</code>	setting <code>ifqs = TRUE</code> results in horizontal and vertical dotted lines being plotted at the three central quartiles and their values, respectively.
<code>ifshape</code>	when used with function <code>shape</code> or <code>caplot</code> to plot into a panel set <code>ifshape = TRUE</code> to ensure only essential probability scale axis labels are displayed to avoid overplotting on the reduced size panel plot.
<code>pch</code>	by default the plotting symbol is set to a plus, <code>pch = 3</code> , alternate plotting symbols may be chosen from those displayed by <code>display.marks</code> , see also Note below.
<code>cex</code>	by default the size of the text for data set size, <code>N</code> , is set to 80%, i.e. <code>cex = 0.8</code> , and may be changed if required.
<code>cexp</code>	by default the size of the plotting symbol, <code>pch</code> , is set to 100%, and may be changed if required.
<code>cex.axis</code>	if overplotting occurs in the y-axis labelling the size of the y-axis labels may be reduced by setting <code>cex.axis</code> to a number smaller than the default of <code>cex.axis = 0.8</code> .

... further arguments to be passed to methods. For example, the size of the axis titles by setting `cex.lab`, and the size of the plot title by setting `cex.main`. For example, if it is required to make the plot title smaller, add `cex.main = 0.9` to reduce the font size by 10%. The colour of the plotting symbols may be changed from the default black, e.e., `col = 2` for red.

Details

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range a truncated plot will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or `xx` may be defined in the function call as, for example, `Cu[Cu < some.value]` which would remove the influence of one or more outliers having values greater than `some.value`. In this case the number of data values displayed will be the number that are `<some.value`.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plot.

The available symbols are:

`pch`: 0 = square, 1 = circle, 2 = triangle, 3 = plus, 4 = X,
 5 = diamond, 6 = upside-down triangle, 7 = square with X,
 8 = asterisk, 9 = diamond with plus, 10 = circle with plus,
 11 = double triangles, 12 = square with plus,
 13 = circle with X, 14 = square with upside-down triangle.
 Symbols 15 to 18 are solid in the colour specified:
 15 = square, 16 = circle, 17 = triangle, 18 = diamond.

Author(s)

Robert G. Garrett

See Also

[display.marks](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## A stand-alone cumulative normal percentage probability plot
cnpplt(Cu)
```

```
## A more appropriately labelled and scaled cumulative normal percentage
## probability plot using a cross/x rather than a plus
cnpplt(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE,
pch = 4)

## Detach test data
detach(kola.o)
```

 crm.plot

Plot Results of Control Reference Material (CRM) Analyses

Description

Function to plot the results of Control Reference Material (CRM) analyses in the order in which they occur in a file, assuming that this order is a time-series, so that the presence of ‘drift’ may be recognized, in addition to the presence of gross outliers reflecting ‘analytical problems’. The data are plotted as either values, when the associated standard deviation of the CRM is provided, or percent absolute relative difference from the ‘recommended’ value when a target tolerance level is provided (see below). The expected ‘recommended’ value (long-term mean) for the CRM being displayed must be supplied, together with its associated standard deviation, or a target tolerance level for percent absolute relative difference. By default the CRM recommended value and standard deviation are used to plot red dashed lines at the recommended value ± 2 standard deviations, and a green line for the recommended value, alternate standard deviation multiples may be provided. This display is also known as a Shewart plot.

Usage

```
crm.plot(xx, xname = deparse(substitute(xx)), crm.mean = NULL,
crm.sd = NULL, n.sd = 2, crm.tol = NULL, ...)
```

Arguments

xx	a column vector of determinations from a data frame or matrix for a measured parameter on a CRM.
xname	a title can be displayed with the plot and results, e.g., xname = "Cu (mg/kg)". If this field is undefined the character string for xx is used as a default.
crm.mean	the recommended value for the CRM. A value must be provided, otherwise the function will terminate.
crm.sd	the standard deviation associated with the recommended value for the CRM. Appropriate red dotted control lines are plotted above and below the mean.
n.sd	by default 2 standard deviation limits are used on the Shewart plot, alternate values may be supplied.
crm.tol	optionally a percentage tolerance level may be provided for the maximum acceptable absolute relative percent difference from the CRM recommended value, in which case a red dotted control line is added to the plot.
...	any additional arguments to be passed to the plot function for titling, etc.

Details

Either a standard deviation for the CRM analyses or an upper limit tolerance level must be provided, otherwise the function will fail. If both are provided an percentage absolute relative difference plot is displayed.

Where the input data file contains determinations for more than one CRM, either a subset for the CRM of interest must be created, e.g., with `gx.subset`, or the R construct `Cu[CRM=="X"]` must be used to pass the data to the function.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#)

Examples

```
## Make test data available
data(crm.test)
attach(crm.test)

## Plot CRM analyses time-series for CRM-X using the CRM SD estimate
## and the default 2 SD tolerance bounds
crm.plot(Cu[CRM=="X"], "Cu(mg/kg) in CRM-X", crm.mean = 34.5, crm.sd = 2.19)

## Plot CRM analyses time-series for CRM-X using the CRM SD estimate
## and 2.5 SD tolerance bounds
crm.plot(Cu[CRM=="X"], "Cu(mg/kg) in CRM-X", crm.mean = 34.5, crm.sd = 2.19,
n.sd = 2.5)

## Plot CRM analyses time-series for CRM-X using a maximum acceptable
## percentage tolerance bound
crm.plot(Cu[CRM=="X"], "Cu(mg/kg) in CRM-X", crm.mean = 34.5, crm.tol = 15)

## Detach test data
detach(crm.test)
```

crm.plot.new	<i>Plot Results of Control Reference Material (CRM) Analyses Adding New Data</i>
--------------	--

Description

Function to plot new CRM data with historic CRM data. Summary statistics are estimated for the historic data, and if an established mean and SD are not provided these are used to estimate the tolerance bounds for the new data to be plotted. The user may define the number of SDs for the tolerance bounds, the default is two. The user must provide the number of new CRM determinations to be plotted so that the x-axis may be extended appropriately. The inclusive range of the new data to be plotted must be provided to ensure all the data and tolerance bounds are plotted. The new data are plotted into the display once the function has executed using the R `plot` command, see Examples.

Usage

```
crm.plot.new(xx, xname = deparse(substitute(xx)), crm.mean = NULL,
  crm.sd = NULL, n.sd = 2, crm.new = 0, ylim = NULL, ...)
```

Arguments

xx	a column vector of determinations from a data frame or matrix for a measured parameter on a CRM.
xname	a title can be displayed with the plot and results, e.g., xname = "Cu (mg/kg)". If this field is undefined the character string for xx is used as a default.
crm.mean	the recommended value for the CRM. If not defined the mean will be estimated from the input (historic) data.
crm.sd	the standard deviation associated with the recommended value for the CRM. If not defined the SD will be estimated from the input(historic) data. Appropriate red dotted control lines are plotted above and below the mean.
n.sd	by default 2 standard deviation limits are used on the Shewart plot, alternate values may be supplied.
crm.new	the number of new CRM determinations to be added to the plot of historic data. The new data may be in several subsets, crm.new has to be the total of all new determinations to be added.
ylim	the inclusive range of the new determinations to be added. The limits for the y-axis will be determined internally taking into account the limits of the historic data and the estimated 2 standard deviation limits.
...	any additional arguments to be passed to the plot function for titling, etc.

Details

A mean and standard deviation for the historic CRM analyses should be provided. If they are not, estimates are made from the input (historic) data.

Following the completion of the plot and the addition of the new data, the plot may be annotated using the R text function. In the example below, to the right of the cursor position, in red and at 80% font size:

```
text(locator(1), "New determinations as red asterisks", adj = 0, col = 2, cex = 0.8)
```

If `crm.new` is undefined the historic data will be plotted using either the external, or internal, estimates of the historic mean and standard deviation.

Where the input data file contains determinations for more than one CRM, either a subset for the CRM of interest must be created, e.g., with `gx.subset`, or the R construct `Cu[CRM=="X"]` must be used to pass the data to the function.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Author(s)

Robert G. Garrett

See Also

`crm.plot`, `ltdl.fix.df`

Examples

```
## Make test data available
data(crm.test)
attach(crm.test)
data(crm.test.new)

## Determine the range of the new determinations to be plotted
range(crm.test.new$Cu)

## Plot CRM analyses time-series for 25 CRM-X determinations using
## a historic CRM mean and SD estimate, and the default 2 SD
## tolerance bounds
crm.plot.new(Cu[CRM=="X"], "Cu(mg/kg) in CRM-X", crm.mean = 34.5, crm.sd = 2.19,
crm.new = 10, ylim = c(29, 39))

## Add the 10 new CRM-X determinations to the plot as red asterisks
points(seq(26,35), crm.test.new$Cu, pch = 8, col = 2)

## Plot CRM analyses time-series for 25 CRM-X determinations and
## base the CRM mean, SD estimate and confidence bounds on those
## data, using the default 2 SD tolerance bounds
crm.plot.new(Cu[CRM=="X"], "Cu(mg/kg) in CRM-X", crm.new = 10, ylim = c(29, 39))
```

```
## Add the 10 new CRM-X determinations to the plot as red asterisks
points(seq(26,35), crm.test.new$Cu, pch = 8, col = 2)

## Detach test data
detach(crm.test)
```

crm.test

National Geochemical Reconnaissance survey QA/QC data

Description

A subset of Control Reference Material (CRM) data from NGR surveys undertaken in 2000 and 2001.

Usage

```
data(crm.test)
```

Format

A data frame with 97 observations on the following 2 variables. Unique identifiers are present in the data frame, use `dimnames(crm.test)[[1]]` to access or display them.

CRM a code indicating the particular CRM analysed. A factor variable with levels: STSD-1, STSD-2, STSD-3, STSD-4, TILL-4, X, Y, and Z.

Cu the copper determinations, mg/kg.

Details

The 'value' of CRM is used to select the CRM data to be displayed, either by creating a specific subset, e.g., using `gx.subset`, or using the R construct `Cu[CRM=="X"]` in the call to function `crm.plot`.

Source

Internal Geological Survey of Canada files.

 crm.test.new

Simulated National Geochemical Reconnaissance survey QA/QC data

Description

A subset, $N = 10$, of simulated Control Reference Material (CRM) data for copper with mean 34.5 and SD 2.19.

Usage

```
data(crm.test.new)
```

Format

A data frame with 10 observations on the following 2 variables. Unique identifiers are present in the data frame, use `dimnames(crm.test)[[1]]` to access or display them.

CRM a code indicating the particular CRM simulated. A factor variable with a single level: X.

Cu the simulated copper determinations, mg/kg.

Details

A set of simulated copper data for demonstrating function `crm.plot.new`, for details of usage see [crm.plot.new](#).

Source

```
rnorm(10, 34.5, 2.19)
```

 cutter

Function to Identify in Which Interval a Value Falls

Description

Function to identify in which interval of a set of cut points, `cuts`, a value `x` falls within or beyond. The number of intervals is equal to the number of cut points plus 1. Values of `x` have to exceed the value of the cut point to be allocated to the higher interval.

Usage

```
cutter(x, cuts)
```

Arguments

<code>x</code>	name of the vector to be processed.
<code>cuts</code>	the vector of cut points.

Value

`xi` a vector of the same length as `x` containing an integer between 1 and the number of cut points plus 1 indicating in which interval each value of `x` fell. Values `<cut[1]` have `xi` set to 1, and values `>cut[highest]` have `xi` set to the number of cut points plus 1.

Author(s)

Robert G. Garrett

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## Cut the data into quartiles
xi <- cutter(Cu, quantile(Cu, probs = c(0.25, 0.5, 0.75)))

## Detach test data
detach(kola.c)
```

df.test

Check for the Existence of a Data Frame

Description

A utility function to determine if a data frame is attached, or exists in the working directory. If the data frame exists the names of the variables are displayed together with the data frame dimensions.

Usage

```
df.test(dfname)
```

Arguments

`dfname` name of a data frame.

Details

Based on a function shared on S-News.

Author(s)

Unkown

See Also

[search](#), [ls](#)

Examples

```
## Make test data available
data(kola.o)

## Check that the data frame kola.o is available
df.test(kola.o)
```

display.ascii.o	<i>Display the Windows Latin 1 Font Octal Table</i>
-----------------	---

Description

A utility function to display the octal numbers corresponding to the Windows Latin 1 Font.

Usage

```
display.ascii.o()
```

Details

The ASCII octal ‘escape codes’ are used to insert special characters in text strings for axis labelling, and titles etc., in graphical displays. For example the escape string `\265` results in the Greek letter mu being displayed.

Based on a function shared on S-News.

Author(s)

Unknown

Examples

```
display.ascii.o()
```

display.lty	<i>Display Available Line Styles and Colour Codes</i>
-------------	---

Description

Displays the line styles and colours corresponding to `lty = 1 to 9` and `colr = 1 to 9`, respectively.

Usage

```
display.lty()
```

Note

All 'rgr' functions that plot boxes or polygons have their default infill colour, `colr`, set to grey, `colr = 8`. This may be changed to an alternate colour, `colr = 1` to `7` or `9`, for no infill colour, set `colr = 0`.

Author(s)

Robert G. Garrett

`display.marks` *Display Available Plotting Marks*

Description

Displays the available plotting marks. Where specified, the 'rgr' functions use a plus sign, `pch = 3`, as the plotting symbol, alternate plotting marks may be selected from this display. For example, `pch = 1` results in an open circle, the R default, and `pch = 4` results in a cross, 'x'. For additional symbols available only in R (`pch = 19:25`) see [points](#).

Usage

```
display.marks()
```

Note

Function to display `pch` codes based on a script originally shared on S-News by Bill Venables, 1996/07, and modified by Shawn Boles, 1996/07/31.

Author(s)

Various, see Note

`display.rainbow` *Display the Colours of the Rainbow(36) Palette*

Description

Displays the available colours in the `rainbow(36)` palette to support the selection of alternate colour schemes.

Usage

```
display.rainbow()
```

Author(s)

Robert G. Garrett

expit	<i>Inverse-logit transformation(s)</i>
-------	--

Description

Undertakes an inverse-logit transformation for a vector or single value.

Usage

```
expit(z)
```

Arguments

`z` the value(s) to be inverse-logit transformed. Natural logarithms are used.

Details

Most analytical chemical data for major, minor and trace elements are of a closed form, i.e. for a sample they sum to a constant, whether it be percent, ppm (mg/kg), or some other units. It does not matter that only some components contributing to the constant sum are present in the matrix, the data are closed. As a result, as some elements increase in concentration others must decrease, this leads to statistics and graphical presentations that do not reflect the true underlying situation even in situations of univariate data analysis and display. The [logit](#) transformation provides an appropriate transformation for univariate compositional data. Procedures for removing closure effects for multivariate data are additive log-ratios ([alr](#)), centred log-ratios ([clr](#)), and isometric log-ratios ([ilr](#)).

Value

`p` the proportion(s) corresponding to the logit transformed value(s), `z`, passed to the function.

Note

This function is provided so that summary statistics generated by 'rgr' functions can be back-transformed to the original units following computations using logit transformed data, see [logit](#).

Author(s)

Robert G. Garrett

References

Filzmoser, P., Hron, K. and Reimann, C., 2009. Univariate statistical analysis of environmental (compositional) data: Problems and possibilities. *Science of the Total Environment*, 407(1/3):6100-6108.

See Also

[logit](#), [alr](#), [clr](#), [ilr](#)

Examples

```
## Generate test data
z <- c(1.6, 0, -2.3)

## Undertake and display inverse-logit transformation(s)
p <- expit(z)
p

## Clean-up
rm(z)
rm(p)
```

fences

Generate and Display Fence Values

Description

Function to generate fence values to support the selection of the upper and lower bounds of background variability, i.e. threshold(s) or action levels, when an obvious graphical solution is not visually recognizable.

Usage

```
fences(xx, units = "ppm", display = TRUE)
```

Arguments

xx	name of the variable to be processed.
units	the units of measurement, options are: "pct", "ppm", "ppb", "ppt". The default is "ppm".
display	the default is to display the tabular output on the current device, i.e. display = TRUE. However, when the function is used by fences.summary and in order to suppress output to the current device display = FALSE as the displayed results will be saved to a text file for subsequent use/editing and reference.

Details

The fence values are computed by several procedures both with and without a logarithmic data transformation and with a logit transformation, together with the 98th percentile of the data for display. Fences are computed following Tukey's boxplot procedure, as median $\pm 2 * MAD$ (Median Absolute Deviation), and mean $\pm 2 * SD$ (Standard Deviation), see Reimann et al. (2005). It is essential that these estimates be viewed in the context of the graphical distributional displays, e.g., [shape](#) and its graphical components, [gx.hist](#), [gx.ecdf](#), [cnpplt](#) and [bxplot](#), and if spatial coordinates for the

sample sites are available [map.eda7](#), [map.eda8](#) and [caplot](#). The final selection of a range for background or the selection of a threshold level needs to take the statistical and spatial distributions of the data into account. It is also necessary to be aware that it might be appropriate to have more than one background range/threshold in a study or survey (Reimann and Garrett, 2005). The presence of relevant information in the data frame may permit the data to be subset on the basis of that information for display with the [tbplots](#), [bwplots](#) and [gx.cnplots](#) functions. If these indicate that the medians and middle 50% of the data are visibly different, multiple background ranges may be advisable.

Note

The logit transformation requires that the input value be in the range zero to one. This transformation takes into consideration the closed, constant sum, nature of geochemical analytical data (Filzmoser et al., 2009). Therefore the measurement units must be defined so that the value can be divided by the appropriate constant. The default is “ppm”, and other acceptable units are “pct”, “ppb” and “ppt”. However, it should be noted that at trace element levels the differences between fences computed with logarithmic and logit transformations are small, and in most applied geochemical applications the logarithmic transformation will suffice. This is not the case for concentrations at major element levels, where the data are more ‘normally’ distributed and fences will be markedly different between untransformed and logit based estimates.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to computing the fences.

Author(s)

Robert G. Garrett

References

- Filzmoser, P., Hron, K. and Reimann, C., 2009. Univariate statistical analysis of environmental (compositional) data: Problems and possibilities. *Science of the Total Environment*, 407(1/3):6100-6108.
- Reimann, C. and Garrett, R.G., 2005. Geochemical background - Concept and reality. *Science of the Total Environment*, 350(1-3):12-27.
- Reimann, C., Filzmoser, P. and Garrett, R.G., 2005. Background and threshold: critical comparison of methods of determination. *Science of the Total Environment*, 346(1-3):1-16.
- Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. *Statistical Data Analysis Explained: Applied Environmental Statistics with R*. John Wiley & Sons, Ltd., 362 p.

See Also

[fences.summary](#), [ltdl.fix.df](#), [remove.na](#), [logit](#), [expit](#)

Examples

```
## Make test data available
data(kola.o)
```

```
attach(kola.o)

## Display the fences computed for Cu
fences(Cu)

## Detach test data
detach(kola.o)
```

fences.summary

Generate and Save Fence Values for Data Subsets

Description

Function to generate fences and save the displayed table of values, by default in the R Working Directory, for subsets of the data for a variable when the data can be subdivided by some criterion (factor) such as EcoRegion, Province, physical sample parent material, etc. The function supports the selection of the upper and lower bounds of background variability, and threshold(s) or action levels, when obvious graphical solutions are not visually recognizable.

Usage

```
fences.summary(group, x, file = NULL, units = "ppm")
```

Arguments

group	the name of the factor variable by which the data are to be subset.
x	name of the variable to be processed.
file	the name of the folder where the file is to be saved, a '/' is appended before the synthesized file name, details of which are displayed on the current device. If no file is specified, the files are saved to the Working Directory, see Details below.
units	the units of measurement, options are: "pct", "ppm", "ppb", "ppt". The default is "ppm".

Details

The fence values are computed by several procedures both with and without a logarithmic data transformation and with a logistic transformation, together with the 98th percentile of the data for display. These computations are based on results returned from function `gx.stats`. Fences are computed following Tukey's boxplot procedure, as median $\pm 2 * MAD$ (Median Absolute Deviation), and mean $\pm 2 * SD$ (Standard Deviation), see Reimann et al. (2005). It is essential that these estimates are viewed in the context of the graphical distributional displays, e.g., `shape` and its graphical components, `gx.hist`, `gx.ecdf`, `cnplot` and `bxplot`, and if spatial coordinates for the sample sites are available `map.eda7`, `map.eda8` and `caplot`. The final selection of a range for background or the selection of a threshold level needs to take the statistical and spatial distributions of the data into account. It is also necessary to be aware that it might be appropriate to have more than one background range/threshold in an area (Reimann and Garrett, 2005). The presence of relevant information in the data frame may permit the data to be subset on the basis of that information for

display with the `tbplots`, `bwplots` and `gx.cnplots` functions. If these indicate that the medians and middle 50% of the data are visibly different, multiple background ranges may be advisable.

A default file name is generated by concatenating the data frame name (see Note below), group and variable, `x`, names, separated by `_s` and terminating in `_fences.txt`. If `file` contains text it is used as the first part of the file name identifying the data source for the file to be saved in the specified folder, for example, `file = "D://R_work//Project3"`, to which the synthesized file name is concatenated. Otherwise the file is saved in the Working Directory.

Output to the current device is suppressed. The output file is formatted as a tab delimited file to be read with a spread sheet program. It can be inspected with a text viewer, and column spacings edited for cosmetic purposes with an ASCII editor of the user's choice.

Note

The synthesis of the file name uses the data frame name which it is assumed is located in `search()` position `[[2]]`.

The logit transformation requires that the input value be in the range zero to one. This transformation takes into consideration the closed, constant sum, nature of geochemical analytical data (Filzmoser et al., 2009). Therefore the measurement units must be defined so that the value can be divided by the appropriate constant. The default is "ppm", and other acceptable units are "pct", "ppb" and "ppt". However, it should be noted that at trace element levels the differences between fences computed with logarithmic and logit transformations are small, and in most applied geochemical applications the logarithmic transformation will suffice. This is not the case for concentrations at major element levels, where the data are more 'normally' distributed and fences will be markedly different between untransformed and logit based estimates.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Any NAs in the data vector are removed prior to computing the fences.

The function `fences` is employed to compute the statistical fence estimates.

Author(s)

Robert G. Garrett

References

- Filzmoser, P., Hron, K. and Reimann, C., 2009. Univariate statistical analysis of environmental (compositional) data: Problems and possibilities. *Science of the Total Environment*, 407(1/3):6100-6108.
- Reimann, C. and Garrett, R.G., 2005. Geochemical background - Concept and reality. *Science of the Total Environment*, 350(1-3):12-27.
- Reimann, C., Filzmoser, P. and Garrett, R.G., 2005. Background and threshold: critical comparison of methods of determination. *Science of the Total Environment*, 346(1-3):1-16.
- Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. *Statistical Data Analysis Explained: Applied Environmental Statistics with R*. John Wiley & Sons, Ltd., 362 p.

See Also

[fences](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## Saves the file kola_c_COUNTRY_Cu_fences.txt for later use
## in the R Working Directory.
fences.summary(COUNTRY, Cu)

## Detach test data
detach(kola.c)
```

fix.test

Test Data for Function ltdl.fix.df

Description

A set of test data to demonstrate how negative values are changed to half their positive value. Optionally numeric coded values representing missing data and/or zero values may be replaced by NAs.

The .csv file was read without deleting ID, the row (observation) identifier in the first column, from the header record. Therefore the character row ID is saved as a factor variable. If ID had been deleted from the header record the row ID would have been stored as `dimnames(fix.test)[[1]]`.

Usage

```
fix.test
```

Format

A data frame containing 15 rows and 5 columns (3 factors, one is ID, and 2 numeric).

See Also

[fix.test.asis](#)

`fix.test.asis`*Test Data for Function ldl.fix.df*

Description

A set of test data to demonstrate how negative values are changed to half their positive value. Optionally numeric coded values representing missing data and/or zero values may be replaced by NAs.

The .csv file was read without deleting ID, the row (observation) identifier in the first column, from the header record, and with `as.is` set to `as.is = c(1)`. Therefore the character row ID is saved as a character variable, not as a factor variable. If ID had been deleted from the header record the row ID would have been stored as `dimnames(fix.test)[[1]]`.

Usage

```
fix.test
```

Format

A data frame containing 15 rows and 5 columns (1 character, 2 factors, and 2 numeric).

See Also

[fix.test](#)

`framework.stats`*Compile Framework/Subset Summary Statistics*

Description

Function to compile summary statistics for use with function [framework.summary](#) from the ‘output’ of function `gx.stats`.

Usage

```
framework.stats(xx)
```

Arguments

`xx` name of the variable to be processed.

Details

The function compiles summary statistics consisting of the count of valid data, the number of NAs, the minimum, 2nd, 5th, 10th, 25th (Q1), 50th (median), 75th (Q3), 90th, 95th and 98th percentiles and the maximum. The 95% confidence interval for the median is computed via the binomial theorem. In addition the Median Absolute Deviation (MAD) and Inter-Quartile Standard Deviation (IQSD) are computed as robust estimates of the standard deviation. Finally, the mean, standard deviation and coefficient of variation as a percentage are computed.

Value

table	a 20-element table is returned, see below:
[1]	the data/subset (sample) size, N.
[2]	number of NAs encountered in the input vector, NNA.
[3:13]	the data minimum, 2nd, 5th, 10th, 25th (Q1), 50th (median), 75th (Q3), 90th, 95th and 98th percentiles and the maximum.
[14:15]	the lower and upper 95% confidence bounds for the median.
[16]	the Median Absolute Deviation (MAD).
[17]	the Inter-Quartile Standard Deviation (IQSD).
[18]	the data (sample) Mean.
[19]	the data (sample) Standard Deviation (SD).
[20]	the Coefficient of Variation as a percentage (CV%).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are counted and then removed prior to computing the summary statistics.

Author(s)

Robert G. Garrett

See Also

[gx.stats](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## Computes summary statistics for the Cu data
fs <- framework.stats(Cu)
fs

## Computes summary statistics for Finnish subset of the Cu data
```

```
fs <- framework.stats(Cu[COUNTRY == "FIN"])
fs

## Clean-up and detach test data
rm(fs)
detach(kola.c)
```

framework.summary *Generate and Save Framework/Subset Summary Statistics*

Description

Function to generate ‘framework’ or subset summary statistics and save them as a ‘.csv’ file in the R working directory. The file can be directly imported into a spreadsheet, e.g., MS Excel, for inspection, or into other software, e.g., a Geographical Information System (GIS) where the spatial information concerning the ‘framework’ units is available, e.g., ecoclassification units.

Usage

```
framework.summary(group, x, file = NULL)
```

Arguments

group	the name of the factor variable by which the data are to be subset.
x	name of the variable to be processed.
file	the name of the folder where the file is to be saved, a ‘/’ is appended before the synthesized file name, details of which are displayed on the current device. If no file is specified, the files are saved to the Working Directory, see Details below.

Details

The file name is synthesized by concatenating the data frame name, group and variable, x, names, separated by `_s`, and `.csv`. If a folder name has been provided in `file`, for example, `file = "D://R_work//Project3"`, a ‘/’ is appended prior to the synthesized file name.

The file name for the `.csv` file is displayed on the current device, and all other output is suppressed. The output file can be inspected with spread sheet software or a viewer of the user’s choice.

Note

The synthesis of the file name uses the data frame name which it is assumed is located in `search()` position `[[2]]`.

To set the R working directory, if it has not already been set in a first function, use at the R command line, for example, `setwd("C:\R\WDn")`, where ‘n’ is some number, which will result in all saved output being placed in that folder. The folder must be created before the R session commences.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are counted and then removed prior to computing the summary statistics.

The function [framework.stats](#) is employed to compute the summary statistics.

Author(s)

Robert G. Garrett

See Also

[framework.stats](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## Saves the file kola_c_COUNTRY_Cu_summary.csv for later use
## in the R Working Directory.
framework.summary(COUNTRY, Cu)

## Detach test data
detach(kola.c)
```

gx.2dproj

Function to Compute and Display 2-d Projections for Data Matrices

Description

Function computes and displays 2-d projections of data matrices using either Sammon Non-linear Mapping (default), Multidimensional Scaling, Kruskal's non-metric Multidimensional Scaling (see Venables and Ripley (2001) and Cox and Cox (2001)). The original S-Plus implementation also computed the Minimum Spanning Tree plane projection (Friedman and Rafsky, 1981) as it was available in the Venables and Ripley MASS library for S-Plus. However, the R implementation of the MASS library does not include Minimum Spanning Trees. In the R implementation, Projection Pursuit has been added using the fastICA procedure of Hyvarinen and Oja (2000). Provision is made to optionally trim individuals (rows) from the input data matrix.

Usage

```
gx.2dproj(xx, proc = "sam", ifilr = FALSE, log = FALSE, rsnd = FALSE, snd = FALSE,
range = FALSE, main = "", setseed = FALSE, row.omits = NULL, ...)
```


Arguments

xx	then by p matrix for which the 2-d projection is required.
proc	the 2-d projection procedure required, the default is <code>proc = "sam"</code> for Sammon Non-Linear Mapping. For Classic (metric) Multidimensional Scaling use <code>proc = "mds"</code> , for Kruskal's non-metric Multidimensional Scaling use <code>"iso"</code> , and for Projection Pursuit use <code>"ica"</code> .
ifilr	optional isometric log-ratio transformation, the default is no transformation. Recommended for closed compositionl, geochemical, data, when <code>ifilr = TRUE</code> all other transformations are ignored.
log	optional (natural) log transformation of the data, the default is no log transformation. For a log transformation set <code>log = TRUE</code> .
rsnd	optional robust normalization of the data with matrix column medians and MADs, the default is no transformation. For a robust normalization set <code>rsnd = TRUE</code> .
snd	optional normalization of the data with matrix column means and standard deviations, the default is no transformation. For a normalization set <code>snd = TRUE</code> . If <code>rsnd = TRUE</code> , then <code>snd</code> will be set to <code>FALSE</code> .
range	optional range transformation for the matrix columns, the data values being scaled to between zero and one for, respectively, the minimum and maximum column values. If the data are range transformed, other normalization transformation requests will be ignored.
main	an alternative plot title, see Details below.
row.omits	permits rows, individuals, to be trimmed from the input matrix, the default <code>row.omits = NULL</code> is for no trimming. To trim individuals enter their row numbers as a concatenated string, e.g. <code>row.omits = c(13,15,16)</code> . The list may be extended by adding additional row numbers so as to display the 2-d structure of the remaining core data and whether further multivariate outliers are present.
setseed	sets the random number seed for fastICA so that all runs result in the same projection, and that projection is generally similar to the Sammon projection on the <code>ilr</code> transformed Howarth - Sinding-Larsen data set.
...	further arguments to be passed to methods concerning the generated plots. For example, if smaller plotting characters are required, specify <code>cex = 0.8</code> ; or if some colour other than black is required for the plotting characters, specify <code>col = 2</code> to obtain red (see display.lty for the default colour palette). If it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

If `main` is undefined a default plot title is generated by appending the input matrix name to the text string "2-d Projection for: ". If no plot title is required set `main = ""`, or if a user defined plot title is required it should be defined in `main`, e.g., `main = "Plot Title Text"`.

Firstly, it is strongly recommended that if the input data matrix is for data from a closed compositional, geochemical, data matrix that an `ilr` transform be applied to the data, `ifilr = TRUE`. This

has the effect of reducing the dimension of the data matrix from p to $(p-1)$. Otherwise, it is desirable to normalize, centre and scale, or undertake a range transformation on the data to ensure the variables have equal ‘weight’ in the projections. If no transformation is requested a warning message is displayed.

The x- and y-axis labels are set appropriately to indicated the type of 2-d projection in the display.

A measure of the ‘stress’ in generating the 2-d projection is estimated and displayed, low stress indicates the projection faithfully represents the relative ‘positions’ of the data in the original p -space.

Value

The following are returned as an object to be saved for further use:

<code>main</code>	the plot title.
<code>input</code>	a text string containing the name of the n by p matrix containing the data, and a list of the row numbers of any individuals trimmed, if none are trimmed the entry is NULL.
<code>usage</code>	The projection option selected, and the values, TRUE or FALSE, for the ilr, log, robust normalization, normalization, and range transformation options.
<code>xlab</code>	the 2-d projection x-axis label.
<code>ylab</code>	the 2-d projection y-axis label.
<code>matnames</code>	the individual, sample, row identifiers and the names of the input variables. If there are no individual, sample, row identifiers then row numbers are used. If an ilr transform has been used the variable names will be the $(p-1)$ synthetic ilr variable names. If a trim has been executed only the row identifiers for the remaining data are stored.
<code>row.numbers</code>	the row numbers of the individuals, samples, remaining after a trim. If a trim has been executed only the row numbers for the remaining data are stored.
<code>x</code>	the n x-axis values for the 2-d projection.
<code>y</code>	the n y-axis values for the 2-d projection.
<code>stress</code>	the estimated stress of fitting 2-d projection to the p -space data.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with with NAs are removed prior to computing the 2-d projection. In the instance of an ilr transformation NAs have to be removed prior to undertaking the transformation, see [remove.na](#).

The results of repeated executions of the ‘fastICA’ implementation of Projection Pursuit lead to various mirror images of one another unless `set.seed` is used to ensure each execution commences with the same seed.

This function requires that packages MASS (Venables and Ripley) and fastICA (Marchini, Heaton and Ripley) both be available.

Author(s)

Robert G. Garrett

References

- Cox, T.F. and Cox, M.A.A., 2001. *Multidimensional Scaling*. Chapman and Hall, 308 p.
- Friedman, J.H. and Rafsky, L.C., 1981. Graphics for the multivariate two-sample problem. *Journal of the American Statistical Association*, 76(374):277-291.
- Hyvarinen, A. and Oja, E., 2000. Independent Component Analysis: Algorithms and Applications. *Neural Networks*, 13(4-5):411-430.
- Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. *Statistical Data Analysis Explained: Applied Environmental Statistics with R*. John Wiley & Sons, Ltd., 362 p.
- Venables, W.N. and Ripley, B.D., 2001. *Modern Applied Statistics with S-Plus*, 3rd Edition. Springer, 501 p.

See Also

[ltdl.fix.df](#), [remove.na](#), [gx.2dproj.plot](#), [sammon](#), [cmdscale](#), [isoMDS](#), [fastICA](#), [set.seed](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Display default, Sammon non-linear map, 2-d projection
sind.2dproj <- gx.2dproj(sind.mat2open, ifilr = TRUE)

## Display saved object identifying input matrix row numbers (cex = 0.7),
## and with an alternate main title (cex.main = 0.8)
gx.2dproj.plot(sind.2dproj, rowids = TRUE, cex = 0.7, cex.main = 0.8,
main = "Howarth & Sinding-Larsen\nStream Sediment ilr Transformed Data")

## Display Kruskal's non-metric multidimensional scaling 2-d projection
sind.2dproj <- gx.2dproj(sind.mat2open, proc = "iso", ifilr = TRUE)

## Display saved object identifying input matrix row numbers (cex = 0.7),
## and with an alternate main title (cex.main = 0.8)
gx.2dproj.plot(sind.2dproj, rowids = FALSE, cex = 0.7, cex.main = 0.8,
main = "Howarth & Sinding-Larsen\nStream Sediment ilr Transformed Data")

## Display default, Sammon non-linear map, 2-d projection, removing the three
## most extreme individuals
sind.2dproj.trim3 <- gx.2dproj(sind.mat2open, ifilr = TRUE, row.omits = c(13,15,16))

## Clean-up
rm(sind.2dproj)
rm(sind.2dproj.trim3)
```

 gx.2dproj.plot

 Function to Display a Saved 2-d Projection Object

Description

Displays the 2-d projection saved from [gx.2dproj](#), optionally the row numbers of the input matrix or the row identifiers, if available, may be displayed instead of the default plotting symbol.

Usage

```
gx.2dproj.plot(save, rowids = NULL, main = "", pch = 3, cex = 0.8, col = 1, ...)
```

Arguments

save	the saved object from gx.2dproj .
rowids	to display the input matrix row numbers set rowids = TRUE. Setting rowids = FALSE replaces the row numbers with the row identifiers from the input matrix. The default rowids = NULL causes the default symbol, a '+' to be plotted.
main	an alternative plot title to that in the saved object from gx.2dproj , see Details below.
pch	by default a '+', pch = 3 will be plotted. See Note below for alternate plotting symbols.
cex	by default symbols and characters are plotted at cex = 0.8, a 20% reduction in font size.
col	by default symbols and characters are plotted in black, col = 1, specify col = 2 to obtain red (see display.lty for the default colour palette). See Note below for alternate colours.
...	further arguments to be passed to methods concerning the plot. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Details

If main is undefined the plot title from the saved object from [gx.2dproj](#) is displayed. If no plot title is required set main = " ", or if a user defined plot title is required it should be defined in main, e.g., main = "Plot Title Text".

The x- and y-axis labels are those in the saved object from [gx.2dproj](#) and indicate the type of 2-d projection in the display.

Note

The available symbols are:

pch: 0 = square, 1 = circle, 2 = triangle, 3 = plus, 4 = X,
 5 = diamond, 6 = upside-down triangle, 7 = square with X,
 8 = asterisk, 9 = diamond with plus, 10 = circle with plus,

11 = double triangles, 12 = square with plus,
 13 = circle with X, 14 = square with upside-down triangle.
 Symbols 15 to 18 are solid in the colour specified:
 15 = square, 16 = circle, 17 = triangle, 18 = diamond.

The available colours from the default 'palette' are:
 Col: 0 = none, 1 = black, 2 = red, 3 = green, 4 = dark blue,
 5 = turquoise, 6 = pink, 7 = yellow, 8 = grey, 9 = black.

Author(s)

Robert G. Garrett

References

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p.

See Also

[gx.2dproj](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Display default 2-d projection
sind.save <- gx.2dproj(sind.mat2open, ifilr = TRUE)

## Display saved object with more informative title
gx.2dproj.plot(sind.save,
main = "Howarth & Sinding-Larsen\nStream Sediments, ilr Transformed Data")

## Display saved object with row identifiers in a smaller blue font
gx.2dproj.plot(sind.save,
main = "Howarth & Sinding-Larsen\nStream Sediments, ilr Transformed Data",
rowids = TRUE, cex = 0.7, col = 4)

## Display 2-d projection after trimming the 3 most extreme samples
sind.2dproj.trim3 <- gx.2dproj(sind.mat2open, ifilr = TRUE, row.omits = c(13,15,16))
gx.2dproj.plot(sind.save,
main = "Howarth & Sinding-Larsen\nStream Sediments, ilr Transformed Data",
rowids = FALSE, cex = 0.7, col = 4)

## Clean-up
rm(sind.save)
rm(sind.2dproj.trim3)
```

 gx.add.chisq

Function to Add Fences to Chi-square Plots

Description

This is an internal function used to plot fences at stated probability levels on a Chi-square plot to assist in the assessment of the plotted distribution. By default fences are plotted for the 90th, 95th and 98th percentiles of the Chi-square distribution. The function is called from `gx.md.plt0`, itself called from `gx.md.plot` that is used to display Chi-square plots generated by `gx.mva`, `gx.mva.closed`, `gx.robmva`, `gx.robmva.closed`, `gx.md.gait` and `gx.md.gait.closed`.

Usage

```
gx.add.chisq(p = c(0.98, 0.95, 0.9), df = NULL, ifflip = FALSE,
            cex = 0.6)
```

Arguments

<code>p</code>	the percentiles of the Chi-square distribution for the fences to be displayed, by default the 90th, 95th and 98th percentiles. If no fences are required set <code>p = NULL</code> .
<code>df</code>	the degrees of freedom for the Chi-square distribution, the number of variables in the multivariate distribution.
<code>ifflip</code>	by default fence labelling is placed to the left of the fences just above the x-axis. Setting <code>ifflip = TRUE</code> places the annotation to the right.
<code>cex</code>	the scale expansion factor for the fence labelling, by default <code>cex = 0.6</code> .

Author(s)

Robert G. Garrett

See Also

[gx.md.plot](#), [gx.md.gait](#), [gx.md.gait.closed](#)

Examples

```
## Synthesize test data
test <- mvrnorm(100, mu = c(40, 30), Sigma = matrix(c(6, 3, 3, 2), 2, 2))

## Display annotated Chi-square plot
gx.md.gait(test)
gx.md.gait(test, ifadd = c(0.9, 0.98))

## Clean-up
rm(test)
```

`gx.adj2`*Function to compute Adjusted r-squared values*

Description

Function to compute the Adjusted R-square value from the Multiple R-squared value displayed in the [summary](#) of a `lm` object. See Note below.

Usage

```
gx.adj2(mr2, n, p)
```

Arguments

<code>mr2</code>	the Multiple R-squared value.
<code>n</code>	the number of cases in the regression model.
<code>p</code>	the number of independent (explanatory or predictor) variables in the model.

Note

The Adjusted R-squared value is a long established criterion. It may be calculated casually by this function, or may be extracted from a `lm` object, using `summary(lm.object)[[9]]`. However, users are urged to investigate Akaike's Information Criterion, [AIC](#), as a procedure for comparing the fits of alternate models, and the use of the [step](#) function for automated model selection.

Author(s)

Robert G. Garrett

See Also

[summary](#), [AIC](#), [step](#)

Examples

```
gx.adj2(0.7394, 111, 11)
gx.adj2(0.713, 111, 6)
```

 gx.cnpplts

Multiple (max 9) Cumulative Normal Probability (CPP) plots

Description

Displays cumulative normal probability (CPP) plots for up to nine data subsets, using combinations of symbols and colours to identify each subset. Note CPP plots are equivalent to Q-Q plots and are more frequently used by physical scientists and engineers.

Usage

```
gx.cnpplts(xlab = " ", log = FALSE, xlim = NULL, main = "",
  iflgnd = FALSE, ...)
```

Arguments

xlab	a title for the x-axis must be provided, even if it is ‘no title’, i.e. xlab = "", or an informative title may be provided, see Examples.
log	to display the data with logarithmic (x-axis) scaling, set log = TRUE.
xlim	if the internally generated values for xlim are to be replaced see the Note below. If the internally generated x-axis limits are satisfactory omit any reference to xlim in the call to the function.
main	a title must be provided, even if it is ‘no title’, i.e. main = "". If main is specified a title will be added above the plot, e.g., main = "Kola Project, 1995".
iflgnd	iflgnd must be specified, TRUE or FALSE. If a R generated legend is to be placed on the plot, set iflgnd = TRUE. On completion of CPP plotting the cursor is activated, locate it at the top left of the space where the legend is to be added and ‘left button’ on the pointing device. The legend comprises the symbol/colour combination, the name of the subset plotted and the data subset size; this information is also displayed on the current device. If no legend is required, set iflgnd = FALSE.
...	the names of the data subsets (objects), separated by commas, to be plotted, up to a maximum of nine. See the example below for subset pre-processing steps that lead to a more presentable legend.

Details

Unlike most other functions in ‘rgr’ all the arguments must be specified explicitly, except xlim. This is the cost of being able to append up to nine subset names in the function call. The function needs to know where subset names start in the list passed to the function.

A default allocation of symbols and colours, and the size of the legend text, is provided in [gx.cnpplts.setup](#). These may be edited if required, they are imported into gx.cnpplts at function run time.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plot.

Although this function is most frequently used to compare the frequency distributions for the same element in multiple subsets of the data, it may also be used to display frequency distributions for multiple elements.

If it is required to set the x-axis limits to specific values they can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range a truncated plot will be displayed. Setting the limits wider than the default limits can provide additional space for annotation of the display.

By setting `iflgn = FALSE` no internally generated legend will be added. Alternately, a legend can be constructed with the `text` function and placed with the `locator` at execution of the `text` function.

Author(s)

Robert G. Garrett

See Also

[gx.cnpplots.setup](#), [display.marks](#), [display.lty](#), [ltdl.fix.df](#), [text](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## An example
gx.cnpplots(xlab = "Cu (mg/kg) in <2 mm Kola C-horizon soil", log = TRUE,
            xlim = NULL, main = "", iflgn = FALSE, Cu[COUNTRY == "RUS"],
            Cu[COUNTRY == "FIN"], Cu[COUNTRY == "NOR"])

## An example that leads to a cleaner legend
## First select data for the variable to be plotted for the subsets, from
## dimnames(kola.c) we know that Be is the 19th column in the data frame
Norway <- gx.subset(kola.c,COUNTRY=="NOR")[,19]
Russia <- gx.subset(kola.c,COUNTRY=="RUS")[,19]
Finland <- gx.subset(kola.c,COUNTRY=="FIN")[,19]
gx.cnpplots(xlab = "Be (mg/kg) in <2 mm Kola C-horizon soils", log = TRUE,
            xlim = NULL, main = "", iflgn = FALSE, Finland, Russia, Norway)

## An example where the limits of the x-axis are provided
gx.cnpplots(xlab = "Be (mg/kg) in <2 mm Kola C-horizon soils", log = TRUE,
            xlim = c(0.02, 20), main = "", iflgn = FALSE, Finland, Russia, Norway)

## An example of a multi-element display
gx.cnpplots(xlab = "Concentrations (mg/kg) in <2 mm Kola C-horizon soils",
```

```

log = TRUE, xlim = NULL, main = "Kola Project, 1995",
iflgnd = FALSE ,Cu, Ni, Co)

## Clean-up and detach test data
rm(Norway)
rm(Russia)
rm(Finland)
detach(kola.c)

```

gx.cnpplts.setup

Set Up and Display Symbolgy for function gx.cnpplts

Description

Permits a user to display the symbol mark and colour combinations to be used in function `gx.cnpplts`, and change them and the legend font size, if required. Any changes require editing the function and some elementary R-scripting skills, see Note below.

Usage

```
gx.cnpplts.setup(display = FALSE)
```

Arguments

`display` if `display = TRUE` the symbol mark and colour combinations are displayed on the current device. If `display = FALSE` output is suppressed.

Details

The available symbols are:

pch: 0 = square, 1 = circle, 2 = triangle, 3 = plus, 4 = X,
5 = diamond, 6 = upside-down triangle, 7 = square with X,
8 = asterisk, 9 = diamond with plus, 10 = circle with plus,
11 = double triangles, 12 = square with plus,
13 = circle with X, 14 = square with upside-down triangle.

Symbols 15 to 18 are solid in the colour specified:

15 = square, 16 = circle, 17 = triangle, 18 = diamond.

The available colours from the default 'palette' are:

Col: 0 = none, 1 = black, 2 = red, 3 = green, 4 = dark blue,
5 = turquoise, 6 = pink, 7 = yellow, 8 = grey, 9 = black.

Value

`pchs` a vector of 9 elements defining the symbols, marks, to use for plotting the 1 to 9 permissible subsets.

`symcols` a vector of 9 elements defining the colours from the 'default' palette to use for the colours of the 1 to 9 permissible subset symbols.

cex	the text scale expansion factor to use in the optional legend for function <code>gx.cnplots</code> , the default is 0.8.
cexp	the scale expansion factor for the plotting symbols in function <code>gx.cnplots</code> , the default is 0.9.

Note

To edit the function use `fix(gx.cnplots.setup)` to extract a copy of the function from the 'rgr' library for editing. It will help to have a colour printed copy of the display, `display = TRUE`, from this function at hand. Note that after editing and saving the function will remain in the workspace and you may get warning messages that can be ignored.

To return to the defaults delete the edited function from the workspace, i.e. `rm(gx.cnplots.setup)`.

Author(s)

Robert G. Garrett

See Also

[display.marks](#), [points](#), [display.lty](#)

 gx.ecdf

Empirical Cumulative Distribution Function (ECDF)

Description

Displays an empirical cumulative distribution function (ECDF) plot with a zero-to-one linear y-scale as part of the multi-panel display provided by [shape](#). The function may also be used stand-alone.

Usage

```
gx.ecdf(xx, xlab = deparse(substitute(xx)),
        ylab = "Empirical Cumulative Distribution Function", log = FALSE,
        xlim = NULL, main = "", pch = 3, ifqs = FALSE, cex = 0.8, ...)
```

Arguments

xx	name of the variable to be plotted.
xlab	by default the character string for xx is used for the x-axis title. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
ylab	a title for the y-axis, defaults to "Empirical Cumulative Distribution Function".
log	to display the data with logarithmic (x-axis) scaling, set <code>log = TRUE</code> .
xlim	when used in the shape function, <code>xlim</code> is determined by gx.hist and used to ensure all four panels in shape have the same x-axis scaling. However, when used stand-alone the limits may be user-defined by setting <code>xlim</code> , see Note below.

main	when used stand-alone a title may be added optionally above the plot by setting main, e.g., main = "Kola Project, 1995".
pch	by default the plotting symbol is set to a plus, pch = 3, alternate plotting symbols may be chosen from those displayed by <code>display.marks</code> , see also Notes below.
ifqs	setting ifqs = TRUE results in horizontal and vertical dotted lines being plotted at the three central quartiles and their values, respectively.
cex	by default the size of the text for data set size, N, is set to 80%, i.e. cex = 0.8, and may be changed if required.
...	further arguments to be passed to methods. The colour of the plotting symbols may be changed from default black, e.g., col = 2 for red. The size of the axis scale annotation can be change by setting cex.axis, the size of the axis titles by setting cex.lab, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Any NAs in the data vector are removed prior to displaying the plot.

Although the cumulative normal percentage probability (CPP) plot is often the preferred method for displaying the cumulative data distribution as it provides greater detail for inspection in the tails of the data, the ECDF is particularly useful for studying the central parts of data distributions as it has not been compressed to make room for the scale expansion in the tails of a cumulative normal percentage probability (CPP) plot.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scale plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range a truncated plot will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations.

The available symbols are:

pch: 0 = square, 1 = circle, 2 = triangle, 3 = plus, 4 = X,
 5 = diamond, 6 = upside-down triangle, 7 = square with X,
 8 = asterisk, 9 = diamond with plus, 10 = circle with plus,
 11 = double triangles, 12 = square with plus,
 13 = circle with X, 14 = square with upside-down triangle.

Symbols 15 to 18 are solid in the colour specified:

15 = square, 16 = circle, 17 = triangle, 18 = diamond.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or `xx` may be defined in the function call as, for example, `Cu[Cu < some.value]` which would remove the influence of one or more outliers having values greater than `some.value`. In this case the number of data values displayed will be the number that are `<some.value`.

Author(s)

Robert G. Garrett

See Also[display.marks](#), [ltdl.fix.df](#), [remove.na](#)**Examples**

```
## Make test data available
data(kola.o)
attach(kola.o)

## Plot a simple ECDF
gx.ecdf(Cu)

## Plot an ECDF with more appropriate labelling and with the quartiles
## indicated
gx.ecdf(Cu , xlab = "Cu (mg/kg) in <2 mm Kola O-horizon soil", log = TRUE,
ifqs = TRUE)

## Detach test data
detach(kola.o)
```

`gx.fractile`*Estimate the Fractile for a Specified Quantile*

Description

Estimates the fractile for a specified quantile of a data set by linear interpolation from the ranked data. If the function is run as `temp <- gx.fractile(xx, q)` the fractile is not displayed, but retained in `temp` for subsequent use or display.

Usage

```
gx.fractile(xx, q, display = TRUE)
```

Arguments

<code>xx</code>	the data set for which the quantile is to be estimated.
<code>q</code>	the fractile (value) for which the quantile is required, must be within the range of <code>xx</code> .
<code>display</code>	the default is to display the quantile and estimated fractile on the current device. If no display is required, set <code>display = FALSE</code> .

Value

<code>f</code>	the estimated fractile.
----------------	-------------------------

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to preparing the boxplots.

Author(s)

Based on a script shared on S-News by Nick Ellis, April 2002

See Also

[ltdl.fix.df](#), [remove.na](#), [gx.quantile](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Estimate the fractile for 20 mg/kg As
gx.fractile(As, 20)
temp <- gx.fractile(As, 20)
temp

## Clean-up and detach test data
rm(temp)
detach(kola.o)
```

gx.hist

Plot a Histogram

Description

Plots a histogram for a data set, the user has options for defining the axis and main titles, the x-axis limits, arithmetic or logarithmic x-axis scaling, the method for calculating the number of bins the data are displayed in, and the colour of the infill.

Usage

```
gx.hist(xx, xlab = deparse(substitute(xx)),
ylab = "Number of Observations", log = FALSE, xlim = NULL,
main = "", nclass = NULL, colr = NULL, ifnright = TRUE,
cex = 0.8, ...)
```

Arguments

xx	name of the variable to be plotted
xlab	by default the character string for xx is used for the x-axis title. An alternate title can be displayed with xlab = "text string", see Examples.
ylab	a default y-axis title of "Number of Observations" is provided, this may be changed, e.g., ylab = "Counts".
log	to display the data with logarithmic (x-axis) scaling, set log = TRUE.
xlim	default limits of the x-axis are determined in the function for use in other panel plots of function shape. However, when used stand-alone the limits may be user-defined by setting xlim, see Note below.
main	when used stand-alone a title may be added optionally above the plot by setting main, e.g., main = "Kola Project, 1995".
nclass	the default procedure for preparing the histogram depends on sample size. Where $N \leq 500$ the Scott (1979) rule is used, and when $N > 500$ the Freedman-Diaconis (1981) rule; both these rules are resistant to the presence of outliers, and usually provide informative histograms. Alternately, the user may define the histogram binning by setting nclass, i.e. nclass = "scott", nclass = "fd" or nclass = "sturges"; the latter being designed for normal distributions (Scott, 1992). See Venables and Ripley (2001) for details.
colr	by default the histogram is infilled in grey, colr = 8. If no infill is required, set colr = 0. See function display.lty for the range of available colours.
ifnright	controls where the sample size is plotted in the histogram display, by default this in the upper right corner of the plot. If the data distribution is such that the upper left corner would be preferable, set ifnright = FALSE. If neither option generates an acceptable plot, setting ifnright = NULL suppresses the display of the data set size.
cex	by default the size of the text for data set size, N, is set to 80%, i.e. cex = 0.8, and may be changed if required.
...	further arguments to be passed to methods. For example, the size of the axis titles may be changed by setting cex.lab, the size of the axis labels by setting cex.axis, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Value

xlim	A two element vector containing the actual minimum [1] and maximum [2] x-axis limits used in the histogram display are returned. These are used in function shape to ensure all panels have the same x-axis limits.
------	---

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plots.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range a truncated plot will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or `xx` may be defined in the function call as, for example, `Cu[Cu < some.value]` which would remove the influence of one or more outliers having values greater than `some.value`. In this case the number of data values displayed will be the number that are `<some.value`.

Author(s)

Robert G. Garrett

References

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p. See pp. 119 for a description of histogram bin selection computations.

See Also

[display.lty](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display to have a first look at the data and
## decide how best to proceed
gx.hist(Cu)

## Provides a more appropriate initial display
gx.hist(Cu, xlab = "Cu (mg/kg) in <2 mm Kola 0-horizon soil", log = TRUE)

## Causes the Sturges rule to be used to select the number
## of histogram bins
gx.hist(Cu, xlab = "Cu (mg/kg) in <2 mm Kola 0-horizon soil", log = TRUE,
nclass = "sturges")

## Detach test data
detach(kola.o)
```

`gx.hypergeom`*Compute Probabilities for Target Recognition*

Description

The hypergeometric distribution is used to infer if the number of anomalous sites along a traverse reliably reflect the presence of the dispersion pattern from a known mineral occurrence. The function displays the probability of the observed outcome could be due to chance alone.

Usage

```
gx.hypergeom(tt, aa, kk, xx)
```

Arguments

tt	total number of sites along a traverse.
aa	number of sites that a priori should be anomalous.
kk	total number of > threshold sites.
xx	number of the aa that are > threshold.

Details

See Stanley (2003) for details, the examples below reproduce the results in Table 1 and Table 2.

Note

Effectively, the hypothesis being tested is that the pattern of above threshold (see [fences](#)), sites coincides the the expected dispersion pattern from a known mineral occurrence. This requires that the geochemist uses knowledge of the dispersion processes active along the traverse, both chemical and mechanical, to predict an expected dispersion pattern.

Author(s)

Robert G. Garrett

References

Stanley, C.R., 2003. Statistical evaluation of anomaly recognition performance. *Geochemistry: Exploration, Environment, Analysis*, 3(1):3-12.

See Also

[gx.runs](#)

Examples

```
## From Stanley (2003) Tables 1 and 2

gx.hypergeom(31, 10, 5, 3)
gx.hypergeom(31, 10, 3, 2)
gx.hypergeom(31, 10, 4, 3)

gx.hypergeom(31, 10, 4, 4)
gx.hypergeom(31, 10, 6, 5)
gx.hypergeom(31, 10, 3, 3)
```

 gx.ks.test

Kolmogorov-Smirnov test with ECDF Plot

Description

Function to plot the Empirical Cumulative Distribution Functions (ECDFs) of two distributions and undertake a Kolmogorov-Smirnov test for the Hypothesis that both distributions were drawn from the same underlying distribution.

Usage

```
gx.ks.test(xx1, xx2, xlab = " ", x1lab = deparse(substitute(xx1)),
           x2lab = deparse(substitute(xx2)),
           ylab = "Empirical Cumulative Distribution Function", log = FALSE,
           main = "", pch1 = 3, col1 = 2, pch2 = 4, col2 = 4,
           ifresult = TRUE, cex = 0.8, cexp = 0.9, ...)
```

Arguments

xx1	name of the first variable to be plotted - distribution to be tested.
xx2	name of the second variable to be plotted - distribution to be tested.
xlab	a title for the x-axis, by default none is provided. For example, xlab = "Cu (mg/kg) in <2 mm C-horizon soil".
x1lab	the name for the first distribution to be plotted, defaults to x1lab = deparse(substitute(xx1)).
x2lab	the name for the second distribution to be plotted, defaults to x2lab = deparse(substitute(xx2)).
ylab	defaults to ylab = "Empirical Cumulative Distribution Function" and may be changed if required.
log	if it is required to display the data with logarithmic (x-axis) scaling, set log = TRUE. The Kolmogorov-Smirnov test is undertaken on untransformed data. If it is to be undertaken on transformed data, the transformation should be applied previously or in the call, e.g., log10(xx1), sqrt(xx1), etc.

main	a plot title if one is required, e.g., main = "Kola Ecogeochemistry Project, 1995".
pch1	the plotting symbol for the first distribution, defaults to a '+' sign, pch = 3, and may be changed if required, see display.marks .
col1	the colour of the plotting symbol for the first distribution, defaults to red, col1 = 2, and may be changed if required, see display.lty .
pch2	the plotting symbol for the second distribution, defaults to a 'x' sign, pch = 4, and may be changed if required, see display.marks .
col2	the colour of the plotting symbol for the second distribution, defaults to blue, col2 = 4, and may be changed if required, see display.lty .
ifresult	setting ifresult = FALSE suppresses the ability to add the results of the Kolmogorov-Smirnov test to the plot, the default is ifresult = TRUE.
cex	the scaling factor for the test results and legend identifying the symbology for each distribution and its population size is set to cex = 0.8 by default, it may be changed if required.
cexp	the scaling factor for the plotting symbol size is set to cexp = 0.9 by default, if may be changed if required.
...	further arguments to be passed to methods. For example, the size of the axis scale annotation can be change by setting cex.axis, the size of the axis titles by seetting cex.lab, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Details

By default the results of the Kolmogorov-Smirnov test are added to the plot. On completion of the ECDF plotting the cursor is activated, locate it at the centre of the area where the results are to be added and 'left button' on the pointing device. When ifresult = FALSE the cursor is not activated for this annotation; this is sometimes convenient if there is insufficient space for the results without overprinting on the ECDFs and report quality plots are required. Also by default a legend is added to the plot, the cursor is activated and should be placed at the top left corner of area where the legend is to be added and 'left button' on the pointing device. The legend consists of two lines indicating the symbology (symbol and colour), name and size of each distribution.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vectors are removed prior to displaying the plot and undertaking the Kolmogorov-Smirnov test.

Author(s)

Robert G. Garrett

See Also

[gx.cnpplts](#), [display.marks](#), [display.lty](#), [ltdl.fix.df](#), [text](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)

## First select data for the variable to be plotted for the subsets, from
## dimnames(kola.c) we know that Be is the 19th column in the data frame
Norway <- gx.subset(kola.c,COUNTRY=="NOR")[,19]
Russia <- gx.subset(kola.c,COUNTRY=="RUS")[,19]
Finland <- gx.subset(kola.c,COUNTRY=="FIN")[,19]

## NOTE: the examples below are commented out as gx.ks.test makes a
## call to the locator function that fails when the examples are run
## during package checking and building
## Initial plot
## gx.ks.test(Finland, Russia, xlab = "Be (mg/kg) in <2 mm Kola C-horizon soils",
## log = TRUE, main = "Kola Ecogeochemistry Project, 1995")

## The same plot as above, but with the results suppressed and the
## annotation better scaled, the legend and plot symbols at 75%, the
## plot title at 90% and the axis labelling at 80%
## gx.ks.test(Finland, Russia, xlab = "Be (mg/kg) in <2 mm Kola C-horizon soils",
## log = TRUE, main = "Kola Ecogeochemistry Project, 1995",
## ifresult = F, cex = 0.75, cexp = 0.75, cex.main = 0.9, cex.lab = 0.8,
## cex.axis = 0.8)

## Clean-up and detach test data
rm(Norway)
rm(Russia)
rm(Finland)
detach(kola.c)
```

gx.lm.vif

Estimate Variance Inflation Factors (VIFs)

Description

Function estimates Variance Inflation Factors (VIFs), measures of collinearity in a linear model. The VIF provides a measure of how much the variance of an estimated regression coefficient is increased because of collinearity. Collinearity is present when there is a high correlation between the independent, predictor, variables in a model, i.e. they tell the same ‘story’. Where collinearity exists it is often best to remove predictor variables with high VIFs from the model.

Usage

```
gx.lm.vif(object, ...)
```

Arguments

object a `lm` object.
... any additional parameters.

Value

A (structure) table of Variable Inflation Factors for the predictor variables.

Note

VIFs >5 are indicative of collinearity, and the information conveyed in that variable is also in the subset of the remaining variables.

Author(s)

W.N. Venables, function shared on S-News, October 21, 2002

References

<http://www.biostat.wustl.edu/archives/html/s-news/2001-10/msg00164.html>

Examples

```
## Make test data available
data(sind)
attach(sind)

## Model 1
sind.1 <- lm(log(Zn) ~ Fe + log(Mn) + log(Cu) + log(Cd))
summary(sind.1)
gx.lm.vif(sind.1)

## Model 2
sind.2 <- lm(log(Zn) ~ Fe + log(Mn))
summary(sind.2)
gx.lm.vif(sind.2)
AIC(sind.1, sind.2)

## Model 3
sind.3 <- lm(log(Zn) ~ log(Mn) + log(Cu))
summary(sind.3)
gx.lm.vif(sind.3)
AIC(sind.1, sind.2, sind.3)

## Clean-up and detach test data
rm(sind.1)
rm(sind.2)
rm(sind.3)
detach(sind)
```

gx.md.display	<i>Function to Display Membership Probabilities and Other Relevant Data</i>
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Description

Function to display the Mahalanobis distances (MDs) and predicted probabilities of membership (ppm or p_{gm}), together with other relevant data, following computations by functions `gx.md.gait`, `gx.md.gait.closed`, `gx.mva`, `gx.mva.closed`, `gx.robmva` or `gx.robmva.closed`. The user may select the predicted probability of membership below which the results are displayed. A simpler presentation is available with `gx.md.print`. Optionally the entire generated table may be saved as a '.csv' file for future use.

Usage

```
gx.md.display(xx, pcut = 0.1, ifprint = TRUE, file = NULL)
```

Arguments

xx	the data matrix to be displayed, see Details below.
pcut	the probability of group membership below which records will be displayed on the current device in ascending order of membership probability, i.e. the most outlying individuals first.
ifprint	by default, <code>ifprint = TRUE</code> , the most outlying individuals are displayed displayed on the current device, see <code>pcut</code> above. To suppress the display set <code>pcut = FALSE</code> .
file	the name of the folder where the file is to be saved, a '/' is appended before the synthesized file name, details of which are displayed on the current device. If no folder is specified, the files are saved to the Working Directory, see Details below. The default, <code>file = NULL</code> , results in no file being saved.

Details

The data frame from which the matrices were derived from for use by the above listed functions must be attached if such items as (numeric) sample IDs, coordinates and data values are to be displayed. Those numeric variables to be displayed must be appended in a `cbind` construct following the 'MDs' and 'ppms' extracted from the saved objects from the above listed functions. For example, `xx <- cbind(save.sind$md, save.sind$ppm, E, N, Zn, Cu)`. If row names were present in the original data frame they will have been carried forward in any saved object from an 'rgr' function and will be displayed in the first column of the table.

The table generated by the function may be saved as a '.csv' file in a folder of the user's choice or in the working directory, with the '.csv' being appended in the function, see Example below. If `file = ""` or `file = " "` the file is saved to the Working Directory. The file name is synthesized as 'MD_display.csv' and appended to the folder name.

Value

table.rows The displayed table is returned and may be saved as an object if required. It will contain the information passed to the function as xx sorted by MD and with appropriate column headings.

Note

This function is similar in purpose to [gx.mvalloc.print](#) for displaying multivariate outliers, however, it operates on a single population.

Author(s)

Robert G. Garrett

See Also

[gx.md.gait](#), [gx.md.gait.closed](#), [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.mvalloc](#), [gx.mvalloc.print](#), [gx.md.print](#).

Examples

```
## Make test data available
data(sind.mat2open)
data(sind)
attach(sind)
## data frame sind attached to provide access to row IDs

## Estimate and default display robust Mahalanobis distances
sind.save <- gx.robmva.closed(sind.mat2open)
xx <- cbind(sind.save$md, sind.save$ppm, E, N, Cu, Zn)
gx.md.display(xx)

## Save display for future use in the Working Directory
gx.md.display(xx, file = " ")

## Clean-up
rm(sind.save)
rm(xx)
detach(sind)
```

Description

Function to undertake the GAIT (Graphical Adaptive Interactive Trimming) procedure for multivariate distributions through Chi-square plots of Mahalanobis distances (MDs) as described in Garrett (1988, 1989). For closed compositional, geochemical, data sets use [gx.md.gait.closed](#). To carry out GAIT the function is called repeatedly with the weights from the previous iteration being used as a starting point. Either a percentage based MVT or a MCD robust start may be used as the first iteration.

Usage

```
gx.md.gait(xx, wts = NULL, trim = -1, mvtstart = FALSE,
mcdstart = FALSE, main = deparse(substitute(xx)),
ifadd = c(0.98, 0.95, 0.9), cexf = 0.6, cex = 0.8, ...)
```

Arguments

xx	the n by p matrix for which the Mahalanobis distances are required.
wts	the vector of weights for the n individuals, either zero or 1.
trim	the desired trim: trim < 0 - no trim, the default; trim >0 & <1 - the fraction, 0 to 1 proportion, of individuals to be trimmed; trim >= 1 - the number of individuals with the highest MDs from the previous iteration to trim.
mvtstart	set mvtstart = TRUE for a percentage based MVT (multivariate trim) start.
mcdstart	set mcdstart = TRUE for a minimum covariance determinant (mcd) robust start.
main	an alternative plot title to the default input data matrix name, see Details below.
ifadd	if probability based fences are to be displayed on the Chi-square plots enter the probabilities here, see Details below. For no fences set ifadd = NULL.
cexf	the scale expansion factor for the Chi-square fence annotation, by default cexf = 0.6.
cex	the scale expansion factor for the symbols and text annotation within the 'frame' of the Chi-square plot, by default cex = 0.8.
...	further arguments to be passed to methods concerning the generated plots. For example, if some colour other than black is required for the plotting characters, specify col = 2 to obtain red (see display.lty for the default colour palette). If it is required to make the plot title or axis labelling smaller, add cex.main = 0.9 or cex.lab = 0.9, respectively, to reduce the font size by 10%.

Details

If main is undefined the name of the matrix object passed to the function is used as the plot title. This is the recommended procedure as it helps to track the progression of the GAIT. Alternate plot titles can be defined if the final saved object is passed to [gx.md.plot](#). If no plot title is required set main = " ", or if a user defined plot title is required it may be defined, e.g., main = "Plot Title Text".

By default three fences are placed on the Chi-square plots at probabilities of membership of the current 'core' data subset, or total data if appropriate, with ifadd = c(0.98, 0.95, 0.9). Alternate probabilities may be defined as best for the display. If no fences are required set ifadd = NULL.

The Mahalanobis distance, Chi-square, plot x-axis label is set appropriately to indicated the type of robust start or trim using the value of `proc`.

Value

The following are returned as an object to be saved for the next iteration or final use:

<code>main</code>	by default (recommended) the input data matrix name.
<code>input</code>	the data matrix name, <code>input = deparse(substitute(xx))</code> , retained to be used by post-processing display functions.
<code>matnames</code>	the row numbers and column headings of the input matrix.
<code>proc</code>	the procedure followed for this iteration, used for subsequent Chi-square plot x-axis labelling.
<code>wts</code>	the vector of weights for the n individuals, either zero or 1.
<code>n</code>	the total number of individuals (observations, cases or samples) in the input data matrix.
<code>ptrim</code>	the percentage, as a fraction, of samples called to be trimmed in this iteration, otherwise <code>ptrim = -1</code> .
<code>mean</code>	the length p vector of means for the 'core' data following the current GAIT step.
<code>cov</code>	the $p \times p$ covariance matrix for the 'core' data following the current GAIT step.
<code>sd</code>	the length p vector of standard deviations for the 'core' data following the current GAIT step.
<code>md</code>	the vector of Mahalanobis distances for all the n individuals following the current GAIT step.
<code>ppm</code>	the vector of predicted probabilities of membership for all the n individuals following the current GAIT step.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Any rows in the data matrix with NAs are removed prior to computations. In the instance of a log-ratio, e.g., `ilr`, transformation NAs are removed.

Warnings are generated when the number of individuals (observations, cases or samples) falls below $5p$, and additional warnings when the number of individuals falls below $3p$. At these low ratios of individuals to variables the shape of the p -space hyperellipsoid is difficult to reliably define, and therefore the results may lack stability. These limits $5p$ and $3p$ are generous, the latter especially so; many statisticians would argue that the number of individuals should not fall below $9p$, see Garrett (1993).

Author(s)

Robert G. Garrett

References

- Garrett, R.G., 1988. IDEAS - An interactive computer graphics tool to assist the exploration geochemist. In Current Research Part F, Geological Survey of Canada Paper 88-1F, pp. 1-13.
- Garrett, R.G., 1993. Another cry from the heart. Explore - Assoc. Exploration Geochemists Newsletter, 81:9-14.
- Garrett, R.G., 1989. The Chi-square plot - a tool for multivariate outlier recognition. In Proc. 12th International Geochemical Exploration Symposium, Geochemical Exploration 1987 (Ed. S. Jenness). Journal of Geochemical Exploration, 32(1/3):319-341.

See Also

[ltdl.fix.df](#), [remove.na](#), [gx.md.plot](#), [gx.md.print](#)

Examples

```
## Note, the example below is presented for historical continuity. It is
## not recommended that this procedure be used for geochemical data. For
## such data function gx.md.gait.closed should be employed. However, the
## example below is consistent with Garrett (1989).
## Make test data available
data(sind)
attach(sind)
sind.mat <- as.matrix(sind[, -c(1:3)])

## Undertake original published GAIT procedure
gx.md.gait(sind.mat)
sind.gait.1 <- gx.md.gait(sind.mat, trim = 0.24, ifadd = 0.98)
sind.gait.2 <- gx.md.gait(sind.mat, wts = sind.gait.1$wts, mvtstart = TRUE,
trim = 4, ifadd = 0.98)
sind.gait.3 <- gx.md.gait(sind.mat, wts = sind.gait.2$wts, trim = 1,
ifadd = 0.9)
sind.gait.4 <- gx.md.gait(sind.mat, wts = sind.gait.3$wts, trim = 2,
ifadd = 0.9)

## Display saved object with alternate main titles and list outliers
## IDEAS procedure
gx.md.plot(sind.gait.4,
main = "Howarth & Sinding-Larsen\nStream Sediments, IDEAS procedure",
cex.main = 0.8, ifadd = 0.9)
gx.md.print(sind.gait.4, pcut = 0.2)

## Clean-up and detach test data
rm(sind.mat)
rm(sind.gait.1)
rm(sind.gait.2)
rm(sind.gait.3)
rm(sind.gait.4)
detach(sind)
```

gx.md.gait.closed	<i>Function for Multivariate Graphical Adaptive Interactive Trimming with Compositional Data</i>
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Description

Function to undertake the GAIT (Graphical Adaptive Interactive Trimming) procedure for multivariate distributions through Chi-square plots of Mahalanobis distances (MDs) as described in Garrett (1988), but for closed compositional, geochemical, data. To carry out GAIT the function is called repeatedly with the weights from the previous iteration being used as a starting point. Either a percentage based MVT or a MCD robust start may be used as the first iteration.

Usage

```
gx.md.gait.closed(xx, wts = NULL, trim = -1, mvtstart = FALSE,
  mcdstart = FALSE, main = deparse(substitute(xx)),
  ifadd = c(0.98, 0.95, 0.9), cexf = 0.6, cex = 0.8, ...)
```

Arguments

xx	the n by p matrix for which the Mahalanobis distances are required.
wts	the vector of weights for the n individuals, either zero or 1.
trim	the desired trim: trim < 0 - no trim, the default; trim >0 & <1 - the fraction, 0 to 1 proportion, of individuals to be trimmed; trim >= 1 - the number of individuals with the highest MDs from the previous iteration to trim.
mvtstart	set mvtstart = TRUE for a percentage based MVT (multivariate trim) start.
mcdstart	set mcdstart = TRUE for a minimum covariance determinant (mcd) robust start.
main	an alternative plot title to the default input data matrix name, see Details below.
ifadd	if probability based fences are to be displayed on the Chi-square plots enter the probabilities here, see Details below. For no fences set ifadd = NULL.
cexf	the scale expansion factor for the Chi-square fence annotation, by default cexf = 0.6.
cex	the scale expansion factor for the symbols and text annotation within the 'frame' of the Chi-square plot, by default cex = 0.8.
...	further arguments to be passed to methods concerning the generated plots. For example, if some colour other than black is required for the plotting characters, specify col = 2 to obtain red (see display.lty for the default colour palette). If it is required to make the plot title or axis labelling smaller, add cex.main = 0.9 or cex.lab = 0.9, respectively, to reduce the font size by 10%.

Details

The variables of the input data matrix must all be expressed in the same units. An isometric log-ratio (ilr) is undertaken and the transformed data used for the GAIT process. At the completion of the process the final ilr estimates, including the inverse of the covariance matrix, are transformed to the centred log-ratio (clr) basis. The vector of means and the inverse of the covariance matrix on a clr basis are required by function `gx.mvalloc.closed`, that is undertaken on a clr basis.

If `main` is undefined the name of the matrix object passed to the function is used as the plot title. This is the recommended procedure as it helps to track the progression of the GAIT. Alternate plot titles can be defined if the final saved object is passed to `gx.md.plot`. If no plot title is required set `main = ""`, or if a user defined plot title is required it may be defined, e.g., `main = "Plot Title Text"`.

By default three fences are placed on the Chi-square plots at probabilities of membership of the current 'core' data subset, or total data if appropriate, with `ifadd = c(0.98, 0.95, 0.9)`. Alternate probabilities may be defined as best for the display. If no fences are required set `ifadd = NULL`.

The Mahalanobis distance, Chi-square, plot x-axis label is set appropriately to indicated the type of robust start or trim using the value of `proc`.

Value

The following are returned as an object to be saved for the next iteration or final use:

<code>main</code>	by default (recommended) the input data matrix name.
<code>input</code>	the data matrix name, <code>input = deparse(substitute(xx))</code> , retained to be used by post-processing display functions.
<code>matnames</code>	the row numbers and column headings of the input matrix.
<code>proc</code>	the procedure followed for this iteration, used for subsequent Chi-square plot x-axis labelling.
<code>wts</code>	the vector of weights for the <code>n</code> individuals, either zero or 1.
<code>n</code>	the total number of individuals (observations, cases or samples) in the input data matrix.
<code>ptrim</code>	the percentage, as a fraction, of samples called to be trimmed in this iteration, otherwise <code>ptrim = -1</code> .
<code>mean</code>	the <code>p</code> length vector of clr basis means for the 'core' data following the current GAIT step.
<code>cov</code>	the <code>p x p</code> clr basis covariance matrix for the 'core' data following the current GAIT step.
<code>cov.inv</code>	the <code>p x p</code> inverse of the covariance matrix following its transformation to the clr basis from the ilr basis. For use by function <code>gx.mvalloc.closed</code> .
<code>sd</code>	the <code>p</code> length vector of clr basis standard deviations for the 'core' data following the current GAIT step.
<code>md</code>	the vector of Mahalanobis distances for all the <code>n</code> individuals following the current GAIT step.
<code>ppm</code>	the vector of predicted probabilities of membership for all the <code>n</code> individuals following the current GAIT step.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data matrix, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with NAs are removed in the ilr transformation.

Warnings are generated when the number of individuals (observations, cases or samples) falls below 5p, and additional warnings when the number of individuals falls below 3p. At these low ratios of individuals to variables the shape of the p-space hyperellipsoid is difficult to reliably define, and therefore the results may lack stability. These limits 5p and 3p are generous, the latter especially so; many statisticians would argue that the number of individuals should not fall below 9p, see Garrett (1993).

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1988. IDEAS - An interactive computer graphics tool to assist the exploration geochemist. In Current Research Part F, Geological Survey of Canada Paper 88-1F, pp. 1-13.

Garrett, R.G., 1993. Another cry from the heart. Explore - Assoc. Exploration Geochemists Newsletter, 81:9-14.

Garrett, R.G., 1989. The Chi-square plot - a tool for multivariate outlier recognition. In Proc. 12th International Geochemical Exploration Symposium, Geochemical Exploration 1987 (Ed. S. Jenness). Journal of Geochemical Exploration, 32(1/3):319-341.

See Also

[ltdl.fix.df](#), [remove.na](#), [gx.md.plot](#), [gx.md.print](#), [ilr](#)

Examples

```
## Make test data available
data(sind.mat2open)

## To multivariate trim as in IDEAS, see JGE (1989) 32(1-3):319-341,
## but recognizing that the data are of a closed compositional form
## and using a mcd start, execute:
gx.md.gait.closed(sind.mat2open,ifadd = 0.95)
sind.gait.1 <- gx.md.gait.closed(sind.mat2open, mcdstart = TRUE,
ifadd = NULL)
sind.gait.2 <- gx.md.gait.closed(sind.mat2open, wts = sind.gait.1$wts,
mvtstart = TRUE, trim = 3, ifadd = 0.9)
sind.gait.3 <- gx.md.gait.closed(sind.mat2open, wts = sind.gait.2$wts,
trim = 1, ifadd = 0.9)

## Display saved object with alternate main titles and list outliers
gx.md.plot(sind.gait.3, cex.main = 0.8, ifadd = 0.9,
main = "Howarth & Sinding-Larsen\nStream Sediments")
```

```

gx.md.print(sind.gait.3, pcut = 0.2)

## Clean-up
rm(sind.gait.1)
rm(sind.gait.2)
rm(sind.gait.3)

```

gx.md.plot

Function to Display Chi-square Plots of Mahalanobis Distances

Description

Function to display Chi-square plots of Mahalanobis distances from objects saved from [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.md.gait](#) and [gx.md.gait.closed](#). The actual plotting of the displays is undertaken by function [gx.md.plt0](#). The function facilitates making ‘cosmetic’ changes to the Chi-square plots not so easily achieved in function [gx.md.gait](#) and [gx.md.gait.closed](#), and not possible in functions [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#) or [gx.robmva.closed](#).

Usage

```

gx.md.plot(save, main = "", ifadd = c(0.98, 0.95, 0.9), cexf = 0.6,
cex = 0.8, ...)

```

Arguments

save	a saved object from the execution of function gx.mva , gx.mva.closed , gx.robmva , gx.robmva.closed , gx.md.gait or gx.md.gait.closed .
main	an alternate Chi-square plot title to that in the saved object, see Details below.
ifadd	if probability based fences are to be displayed on the Chi-square plots enter the probabilities here, see Details below. For no fences set ifadd = NULL.
cexf	the text scale expansion factor for the annotation of the probability based fences, by default cexf = 0.6.
cex	the text scale expansion factor for the other annotation within the ‘frame’ of the Chi-square plot, by default cex = 0.8.
...	further arguments to be passed to methods concerning the generated plots. For example, if some colour other than black is required for the plotting characters, specify col = 2 to obtain red (see display.lty for the default colour palette). If it is required to make the plot title or axis labelling smaller, add cex.main = 0.9 or cex.lab = 0.9, respectively, to reduce the font size by 10%.

Details

If `main` is undefined the name of the matrix object from which the Mahalanobis distances were derived is passed to the function via the saved object. Using the matrix name is the recommended procedure as it helps to track the progression during a GAIT exercise, and acts as a record of the data source. However, at a presentation stage an alternate plot title may preferred and can be defined in this function, e.g., `main = "Plot Title Text"`. If no plot title is required set `main = " "`.

By default three fences are placed on the Chi-square plots at probabilities of membership of the current 'core' data subset, or total data if appropriate, with `ifadd = c(0.98, 0.95, 0.9)`. Alternate probabilities may be defined as best for the display. If no fences are required set `ifadd = NULL`.

The Mahalanobis distance, Chi-square, plot x-axis label is set appropriately to indicated the type of robust start or trim using the value of `proc` from the saved object passed to the function.

Author(s)

Robert G. Garrett

See Also

[gx.md.gait](#), [gx.md.gait.closed](#), [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.add.chisq](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Save and display Chi-square plot
sind.save <- gx.mva(ilr(sind.mat2open))
gx.md.plot(sind.save)
gx.md.plot(sind.save,
  main = "Howarth & Sinding Larsen Stream Sediments\nilr transform",
  cexf = 0.8, cex = 1, col = 2)

## Save and display Chi-square plot with a
## mcd robust start and ilr transformation
sind.save <- gx.md.gait(ilr(sind.mat2open), mcdstart = TRUE, mvtstart = TRUE,
  trim = 3, ifadd = NULL)
gx.md.plot(sind.save)
gx.md.plot(sind.save,
  main = paste("Howarth & Sinding-Larsen\nStream Sediments, ilr Transformed Data",
    "\nMCD robust start"), ifadd = 0.9, cex.main = 0.8)

## Clean-up
rm(sind.save)
```

 gx.md.plt0

Function to Display Chi-square plots of Mahalanobis Distances

Description

This function is not called directly by the user but from functions that plot Mahalanobis distances, i.e. [gx.md.gait](#) and [gx.md.plot](#).

Usage

```
gx.md.plt0(md, n, p, trim = trim, ptrim = -1, proc = proc,
main = main, ifadd = ifadd, cexf = cexf, cex = cex, ...)
```

Arguments

md	a vector of Mahalanobis distances of length n.
n	the length of the vector of Mahalanobis distances.
p	the number of variables upon which the Mahalanobis distances are based.
trim	the number of individuals (observations or samples) that have been trimmed, and did not contribute to the estimation of covariance and means.
ptrim	the percentage trim requested, if a percentage (MVT) trim was requested.
proc	the procedure by which the Mahalanobis distances were estimated, used to ensure appropriate labelling of the Chi-square plot x-axis.
main	the title for the Chi-square plot.
ifadd	the probability based fences to be displayed on the Chi-square plots, set by the calling function and the user.
cexf	the text scale expansion factor for the annotation of the probability based fences, set by the calling function and the user.
cex	the text scale expansion factor for the other annotation within the ‘frame’ of the Chi-square plot, set by the calling function and the user.
...	further arguments to be passed to methods concerning the generated plots. For example, if some colour other than black is required for the plotting characters, specify <code>col = 2</code> to obtain red (see display.lty for the default colour palette). If it is required to make the plot title or axis labelling smaller, add <code>cex.main = 0.9</code> or <code>cex.lab = 0.9</code> , respectively, to reduce the font size by 10%.

Author(s)

Robert G. Garrett

See Also

[gx.md.gait](#), [gx.md.plot](#), [gx.add.chisq](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Generate and display sets of Mahalanobis distances
gx.md.gait(ilr(sind.mat2open))
gx.md.gait(ilr(sind.mat2open), mcdstart = TRUE, ifadd = NULL)
gx.md.gait(ilr(sind.mat2open), mcdstart = TRUE, mvtstart = TRUE, trim = 3,
ifadd = 0.9)
```

gx.md.print

*Function to Display Membership Probabilities***Description**

Function to display the Mahalanobis distances (MDs) and predicted probabilities of membership (ppm) following computations by functions [gx.md.gait](#), [gx.md.gait.closed](#), [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#) or [gx.robmva.closed](#). The user may select the predicted probability of membership below which the results are displayed. Alternately the Mahalanobis distances and group membership probabilities may be saved as a '.csv' file for future use.

Usage

```
gx.md.print(save, pcut = 0.1, ifprint = TRUE, file = NULL)
```

Arguments

save	a saved object from any of functions gx.md.gait , gx.md.gait.closed , gx.mva , gx.mva.closed , gx.robmva , or gx.robmva.closed .
pcut	the probability of group membership below which records will be displayed on the current device in ascending order of membership probability, i.e. the most outlying individuals first.
ifprint	by default, <code>ifprint = TRUE</code> , the most outlying individuals are displayed displayed on the current device, see <code>pcut</code> above. To suppress the display set <code>pcut = FALSE</code> .
file	the name of the folder where the file is to be saved, a '/' is appended before the synthesized file name, details of which are displayed on the current device. If no folder is specified, the files are saved to the Working Directory, see Details below. The default, <code>file = NULL</code> , results in no file being saved.

Details

The Mahalanobis distances, the membership probabilities and input matrix row numbers or IDs are extracted from the saved object and sorted in increasing order of probabilities of group membership for display on the current device. The full table of Mahalanobis distances and group membership probabilities may be saved as a '.csv' file. If a folder name has been provided in `file`, a '\ ' is appended prior to a file name synthesized from the name of the saved object concatenated with

'_MDs.csv'. If `file = ""` or `file = " "` the file will be saved to the Working Directory. Details of the file name are displayed on the current device, see Example below. For an alternative display see [gx.md.display](#).

Value

`table.rows` The last table processed is returned and may be saved as an object if required.

Note

This function is similar in purpose to [gx.mvalloc.print](#) for displaying multivariate outliers, however, it operates on a single population.

Author(s)

Robert G. Garrett

See Also

[gx.md.gait](#), [gx.md.gait.closed](#), [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.md.display](#), [gx.mvalloc](#), [gx.mvalloc.print](#).

Examples

```
## Make test data available
data(sind.mat2open)

## Estimate and display robust Mahalanobis distances
sind.save <- gx.robmva.closed(sind.mat2open)
gx.md.print(sind.save, pcut = 0.3)

## Save display for future use, not executed
## gx.md.print(sind.save, ifprint = FALSE, file = "D://R_work//Project3")

## Clean-up
rm(sind.save)
```

gx.mf

Prepare a Concentration-Number (C-N) Plot

Description

Display a log-log Concentration-Number (C-N) Plot to inspect a data set for multifractality.

Usage

```
gx.mf(xx, xlab = deparse(substitute(xx)),
      ylab = "Cumulative Percentage of Data", main = "Multifractality Plot",
      ifrev = FALSE, xlim = range(xx, na.rm = TRUE), ...)
```

Arguments

<code>xx</code>	name of the variable to be plotted.
<code>xlab</code>	by default the character string for <code>xx</code> is used for the x-axis plot titles. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
<code>ylab</code>	by default the y-axis is labelled 'Cumulative Percentage of Data'. This replaced with no title, <code>main = " "</code> , or text of the user's choice, <code>main = "text string"</code> .
<code>main</code>	by default a plot title of 'Multifractality Plot' is provided. This replaced with no title, <code>main = " "</code> , or text of the user's choice, <code>main = "text string"</code> .
<code>ifrev</code>	by default the empirical C-N function is plotted from highest value to lowest, <code>ifrev = FALSE</code> . As the C-N plot is a log-log display this provides greater detail for the highest values. The direction of accumulation can be key in detecting multifractal patterns, it is usually informative to also prepare a plot with <code>ifrev = TRUE</code> , i.e. accumulation from lowest to highest values. To see a dramatic example of this, run the Examples below.
<code>xlim</code>	by default this is the range of the data with any NAs removed, <code>xlim = range (xx, na.rm = TRUE)</code> . When called from <code>caplot</code> or <code>shape.alt</code> an appropriate value for <code>xlim</code> is determined by the calling functions to ensure all four panels have the same x-axis scaling. <code>xlim</code> may be defined, see Note below.
<code>...</code>	further arguments to be passed to methods. For example, the size of the axis scale annotation can be changed by setting <code>cex.axis</code> , the size of the axis titles by setting <code>cex.lab</code> , and the size of the plot title by setting <code>cex.main</code> . For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%. By default individual data points are marked by a plus sign, <code>pch = 3</code> , if a cross or open circle is desired, then set <code>pch = 4</code> or <code>pch = 1</code> , respectively. See display.marks for all available symbols.

Details

Any multi-fractality in the data, will be revealed as 'lines' of data points with different slopes.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plot.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range truncated plots will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations in the various panels.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or `xx` may be defined in the function call as, for example, `Cu[Cu < some.value]` which would remove the influence of one or more outliers having values greater than `some.value`. In this case the number of data values displayed will be the number that are `<some.value`.

In some R installations the generation of multi-panel displays and the use of function `eqsplot` from package MASS causes warning messages related to graphics parameters to be displayed on the

current device. These may be suppressed by entering options(warn = -1) on the R command line, or that line may be included in a ‘first’ function prepared by the user that loads the ‘rgr’ package, etc.

For summary statistics displays to complement the graphics see, [gx.summary1](#), [gx.summary2](#) and [inset](#).

Author(s)

Robert G. Garrett

References

Agterberg, F.P., 1995. Multifractal Modeling of the Sizes and Grades of Giant and Supergiant Deposits. *International Geology Review*, Vol. 37, No. 1, pp. 1-8.

See Also

[gx.hist](#), [cnpplt](#), [gx.ecdf](#), [gx.mf](#), [remove.na](#), [display.lty](#), [display.marks](#), [ltdl.fix.df](#), [inset](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display to have a first look at the data and
## decide how best to proceed
gx.mf(Cu)

## Provides a more appropriate initial display and indicates the
## quartiles
gx.mf(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", ifqs = TRUE)

## Causes the C-N plot to be cumulated in reverse order. This will reveal
## any multi-fractal properties of the data at lower concentrations
gx.mf(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", ifrev = TRUE)

## Detach test data
detach(kola.o)
```

Description

The function carries out a Principal Components Analysis (PCA) and estimates the Mahalanobis distances for a dataset and places them in an object to be saved and post-processed for display and further manipulation. Classical procedures are used, for robust procedures see [gx.robmva](#). For results display see [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#) and [gx.md.print](#). For Kaiser varimax rotation see [gx.rotate](#). For closed, compositional, data use [gx.mva.closed](#).

Usage

```
gx.mva(xx, main = deparse(substitute(xx)))
```

Arguments

xx	a n by p data matrix to be processed.
main	by default the name of the object xx, <code>main = deparse(substitute(xx))</code> , it may be replaced by the user, but this is not recommended, see Details below.

Details

If `main` is undefined the name of the matrix object passed to the function is used to identify the object. This is the recommended procedure as it helps to track the progression of a data analysis. Alternate plot titles are best defined when the saved object is passed to [gx.rqpca.plot](#), [gx.rqpca.screepplot](#) or [gx.md.plot](#) for display. If no plot title is required set `main = ""`, or if a user defined plot title is required it may be defined, e.g., `main = "Plot Title Text"`.

Value

The following are returned as an object to be saved for subsequent display, etc.:

main	by default (recommended) the input data matrix name.
input	the data matrix name, <code>input = deparse(substitute(xx))</code> , retained to be used by post-processing display functions.
proc	the procedure used, by default <code>proc = "cov"</code> to indicate a classical covariance matrix.
n	the total number of individuals (observations, cases or samples) in the input data matrix.
nc	the number of individuals remaining in the 'core' data subset after trimming. At this stage of a data analysis <code>nc = n</code> .
p	the number of variables on which the multivariate operations were based.
ifilr	flag for gx.md.plot , set to FALSE.
matnames	the row numbers or identifiers and column headings of the input matrix.
wts	the vector of weights for the n individuals used to compute the covariance matrix and means. At this stage of the data analysis all weights are set to '1'.
mean	the vector the weighted means for the p variables.
cov	the p by p weighted covariance matrix for the n by p data matrix.

sd	the vector of weighted standard deviations for the p variables.
snd	the n by p matrix of weighted standard normal deviates.
r	the p by p matrix of weighted Pearson product moment correlation coefficients.
eigenvalues	the vector of p eigenvalues of the scaled Pearson correlation matrix for RQ analysis, see Grunsky (2001).
econtrib	the vector of p eigenvalues each expressed as a percentage of the sum of the eigenvalues.
eigenvectors	the n by p matrix of eigenvectors.
rload	the p by p matrix of Principal Component (PC) loadings.
rcr	the p by p matrix containing the percentages of the variability of each variable (rows) expressed in each PC (columns).
rqscore	the n by p matrix of the n individuals scores on the p PCs.
vcontrib	a vector of p variances of the columns of rqscore.
pvcontrib	the vector of p variances of the columns of rqscore expressed as percentages. This is a check on vector econtrib, the values should be identical.
cpvcontrib	the vector of p cumulative sums of pvcontrib, see above.
md	the vector of n Mahalanobis distances (MDs) for the n by p input matrix.
ppm	the vector of n predicted probabilities of population membership, see Garrett (1990).
epm	the vector of n empirical Chi-square probabilities for the MDs.
nr	the number of PCs that have been rotated. At this stage of a data analysis nr = NULL in order to control PC plot axis labelling.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with NAs are removed prior to computations. In the instance of a compositional data opening transformation NAs have to be removed prior to undertaking the transformation, see [na.omit](#), [where.na](#) and [remove.na](#). When that procedure is followed the opening transformations may be executed on calling the function, see Examples below.

Note that, executing a [clr](#) transformation leads to a singular matrix that can not be inverted for the estimation of Mahalanobis distances. In that case the values of md, ppm and epm are all set to NULL.

Note that, executing a [ilr](#) transformation permits the estimation of Mahalanobis distances and associated probabilities through the use of p-1 synthetic variables. However, in that instance the loadings of the p-1 synthetic variables will be plotted by [gx.rqpca.plot](#) rather than the loadings for the elements.

Therefore, use function [gx.mva.closed](#) for compositional, geochemical, data.

Warnings are generated when the number of individuals (observations, cases or samples) falls below 5p, and additional warnings when the number of individuals falls below 3p. At these low ratios of individuals to variables the shape of the p-space hyperellipsoid is difficult to reliably define, and therefor the results may lack stability. These limits 5p and 3p are generous, the latter especially so; many statisticians would argue that the number of individuals should not fall below 9p, see Garrett (1993).

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1990. A robust multivariate allocation procedure with applications to geochemical data. In Proc. Colloquium on Statistical Applications in the Earth Sciences (Eds F.P. Agterberg & G.F. Bonham-Carter). Geological Survey of Canada Paper 89-9, pp. 309-318.

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Grunsky, E.C., 2001. A program for computing RQ-mode principal components analysis for S-Plus and R. Computers & Geosciences, 27(2):229-235.

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

See Also

[ltdl.fix.df](#), [remove.na](#), [na.omit](#), [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#), [gx.md.print](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.rotate](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Generate gx.mva object, for demonstration purposes only
## These are compositional data - gx.mva.closed should be used
sind.save <- gx.mva(sind.mat2open)
gx.rqpca.screepplot(sind.save)
gx.rqpca.loadplot(sind.save)
gx.rqpca.plot(sind.save)
## Display saved object with alternate main titles
gx.rqpca.loadplot(sind.save,
  main = "Howarth & Sinding-Larsen\nStream Sediments, clr Transformed Data",
  cex.main = 0.8)
gx.rqpca.plot(sind.save,
  main = "Howarth & Sinding-Larsen\nStream Sediments, clr Transformed Data",
  cex.main = 0.8)

## Display Mahalanobis distances in a Chi-square plot
gx.md.plot(sind.save)
## Display saved object with alternate main titles
gx.md.plot(sind.save,
  main = "Howarth & Sinding-Larsen\nStream Sediments, ilr Transformed Data",
  cex.main = 0.8)

## Clean-up
rm(sind.save)
```

gx.mva.closed	<i>Function to undertake an Exploratory Multivariate Data Analysis on Compositional, geochemical data</i>
---------------	---

Description

The function carries out a Principal Components Analysis (PCA) and estimates the Mahalanobis distances for a compositional dataset and places them in an object to be saved and post-processed for display and further manipulation. Classical procedures are used, for robust procedures see [gx.robmva.closed](#). For results display see [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#) and [gx.md.print](#). For Kaiser varimax rotation see [gx.rotate](#).

Usage

```
gx.mva.closed(xx, main = deparse(substitute(xx)))
```

Arguments

xx	a n by p data matrix to be processed.
main	by default the name of the object xx, <code>main = deparse(substitute(xx))</code> , it may be replaced by the user, but this is not recommended, see Details below.

Details

If `main` is undefined the name of the matrix object passed to the function is used to identify the object. This is the recommended procedure as it helps to track the progression of a data analysis. Alternate plot titles are best defined when the saved object is passed to [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.screepplot](#) or [gx.md.plot](#) for display. If no plot title is required set `main = ""`, or if a user defined plot title is required it may be defined, e.g., `main = "Plot Title Text"`.

The data are centre log-ratio transformed prior to undertaking the PCA. For the computation of Mahalanobis distances the data are isometrically log-ratio transformed, this results in the loss of one degree of freedom.

Value

The following are returned as an object to be saved for subsequent display, etc.:

main	by default (recommended) the input data matrix name.
input	the data matrix name, <code>input = deparse(substitute(xx))</code> , retained to be used by post-processing display functions.
proc	the procedure used, by default <code>proc = "cov"</code> to indicate a classical covariance matrix.
n	the total number of individuals (observations, cases or samples) in the input data matrix.
nc	the number of individuals remaining in the 'core' data subset after trimming. At this stage of a data analysis <code>nc = n</code> .

p	the number of variables on which the multivariate operations were based.
ifilr	flag for <code>gx.md.plot</code> , set to TRUE.
matnames	the row numbers or identifiers and column headings of the input matrix.
wts	the vector of weights for the n individuals used to compute the covariance matrix and means. For a classical, non-robust, estimation all weights are set to '1'.
mean	the vector the clr means for the p variables.
cov	the p by p clr covariance matrix for the n by p data matrix.
sd	the vector of clr standard deviations for the p variables.
snd	the n by p matrix of clr standard normal deviates.
r	the p by p matrix of clr Pearson product moment correlation coefficients.
eigenvalues	the vector of p eigenvalues of the scaled Pearson correlation matrix for RQ analysis, see Grunsky (2001).
econtrib	the vector of p eigenvalues each expressed as a percentage of the sum of the eigenvalues.
eigenvectors	the n by p matrix of eigenvectors.
rload	the p by p matrix of Principal Component (PC) loadings.
rcr	the p by p matrix containing the percentages of the variability of each variable (rows) expressed in each PC (columns).
rqscore	the n by p matrix of the n individuals scores on the p PCs.
vcontrib	a vector of p variances of the columns of <code>rqscore</code> .
pvcontrib	the vector of p variances of the columns of <code>rqscore</code> expressed as percentages. This is a check on vector <code>econtrib</code> , the values should be identical.
cpvcontrib	the vector of p cumulative sums of <code>pvcontrib</code> , see above.
md	the vector of n Mahalanobis distances (MDs) for the n by p, now (p-1), input matrix.
ppm	the vector of n predicted probabilities of population membership, see Garrett (1990).
epm	the vector of n empirical Chi-square probabilities for the MDs.
nr	the number of PCs that have been rotated. At this stage of a data analysis <code>nr = NULL</code> in order to control PC plot axis labelling.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with NAs are removed prior to computations. In the instance of a compositional data opening transformation NAs have to be removed prior to undertaking the transformation, see [na.omit](#), [where.na](#) and [remove.na](#). When that procedure is followed the opening transformations may be executed on calling the function, see Examples below.

Warnings are generated when the number of individuals (observations, cases or samples) falls below 5p, and additional warnings when the number of individuals falls below 3p. At these low ratios of individuals to variables the shape of the p-space hyperellipsoid is difficult to reliably define, and

therefor the results may lack stability. These limits 5p and 3p are generous, the latter especially so; many statisticians would argue that the number of individuals should not fall below 9p, see Garrett (1993).

Author(s)

Robert G. Garrett

References

- Garrett, R.G., 1990. A robust multivariate allocation procedure with applications to geochemical data. In Proc. Colloquium on Statistical Applications in the Earth Sciences (Eds F.P. Agterberg & G.F. Bonham-Carter). Geological Survey of Canada Paper 89-9, pp. 309-318.
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See Also

[ltdl.fix.df](#), [remove.na](#), [na.omit](#), [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#), [gx.md.print](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.rotate](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Generate gx.mva object after an clr transform for a PCA
sind.closed <- gx.mva.closed(sind.mat2open)
gx.rqpca.screepplot(sind.closed)
gx.rqpca.plot(sind.closed)
gx.rqpca.loadplot(sind.closed)
## Display saved object with alternate main titles
gx.rqpca.loadplot(sind.closed,
  main = "Howarth & Sinding-Larsen\nStream Sediments, clr Transformed Data",
  cex.main = 0.8)
gx.rqpca.plot(sind.closed,
  main = "Howarth & Sinding-Larsen\nStream Sediments, clr Transformed Data",
  cex.main = 0.8)

## Display Mahalanobis distances with alternate main title
gx.md.plot(sind.closed,
  main = "Howarth & Sinding-Larsen\nStream Sediments, ilr Transformed Data",
  cex.main = 0.8)

## Clean-up
rm(sind.closed)
```

gx.mvalloc

*Function for Allocation on the basis of Multivariate Data***Description**

Function to allocate individuals (observations, cases or samples) into one of multiple reference groups (populations) on the basis of their Mahalanobis distances. If an individual's predicted probability of group membership (typicality) falls below a user defined 'cut-off', `pcrit`, the individual is allocated to an 'outlier bin'.

Usage

```
gx.mvalloc(pcrit = 0.05, x, ...)
```

Arguments

<code>pcrit</code>	the critical cut-off probability for group membership below which an individual will be classified as an 'outlier'. By default the critical probability of group membership is set to <code>pcrit = 0.05</code> .
<code>x</code>	a n by p matrix containing the n individuals, with p variables determined on each, to be allocated, see Details below.
<code>...</code>	a list of objects saved from any of functions gx.md.gait , gx.md.gait.closed , gx.mva , gx.robmva or gx.robmva.closed , containing the vectors of means and covariance matrices for the 'reference' groups into which the individuals are to be classified.

Details

It is imperative that the data matrix `x` contains no special codes and all records (individuals) with NAs have been removed, see Notes below. It is also imperative that the variables in the reference groups and in the matrix `x` of individuals to be classified are identical and in the same order.

The allocations are made on the assumption that the covariance structures are inhomogeneous, i.e. that the population hyperellipsoids are of different size, shape and orientation in p -space.

Value

The following are returned as an object to be saved for display with [gx.mvalloc.print](#):

<code>groups</code>	a list of the names of the <code>kk</code> reference groups.
<code>kk</code>	the number of reference groups passed to the function.
<code>n</code>	the number of individuals (observations, cases or samples) allocated.
<code>p</code>	the number of variables in the reference and allocated data.
<code>pcrit</code>	the critical cut-off probability for reference group membership.
<code>pgm</code>	a vector of <code>kk</code> predicted probabilities of reference group memberships.
<code>xalloc</code>	the reference group, <code>1:kk</code> , that the individual was allocated into. All outliers, i.e. all <code>pgm(1:kk) < crit</code> are allocated to group zero, \emptyset . Therefore <code>xalloc</code> will be in the range of $\emptyset:kk$.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed from the matrix *x* prior to executing this function, see [ltdl.fix.df](#). Additionally, any rows in the data matrix with NAs also must have been removed prior to computations, see [na.omit](#) and [remove.na](#).

It is recommended that when applying this procedure to compositional data an ilr transformation be undertaken, this can be done at execution time. This implies that the reference group means and covariance matrices must have also been estimated following an ilr transformation.

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1990. A robust multivariate allocation procedure with applications to geochemical data. In Proc. Colloquium on Statistical Applications in the Earth Sciences (Eds F.P. Agterberg & G.F. Bonham-Carter). Geological Survey of Canada Paper 89-9, pp. 309-318.

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

See Also

[gx.md.gait](#), [gx.md.gait.closed](#), [gx.mva](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.mvalloc.print](#), [ltdl.fix.df](#), [remove.na](#), [na.omit](#)

Examples

```
## Generate three groups of synthetic bivariate normal data
grp1 <- mvrnorm(100, mu = c(40, 30), Sigma = matrix(c(6, 3, 3, 2), 2, 2))
grp1 <- cbind(grp1, rep(1, 100))
grp2 <- mvrnorm(100, mu = c(50, 40), Sigma = matrix(c(4, -3, -3, 5), 2, 2))
grp2 <- cbind(grp2, rep(2, 100))
grp3 <- mvrnorm(100, mu = c(30, 45), Sigma = matrix(c(6, 4, 4, 5), 2, 2))
grp3 <- cbind(grp3, rep(3, 100))
## Generate a set of six (6) outliers
anom <- matrix(c(35, 40, 25, 60, 25, 60, 35, 40, 25, 60, 60, 25), 6, 2)
anom <- cbind(anom, rep(4, 6))
## Bind the test data sets together and display the test data
test.mvalloc.mat <- rbind(grp1, grp2, grp3, anom)
test.mvalloc <- as.data.frame(test.mvalloc.mat)
dimnames(test.mvalloc)[[2]] <- c("x", "y", "grp")
attach(test.mvalloc)
xyplot.tags(x, y, grp, cex = 0.75)

## Generate robust reference groups
test.save.grp1 <- gx.md.gait(grp1[, -3], mcdstart = TRUE)
test.save.grp2 <- gx.md.gait(grp2[, -3], mcdstart = TRUE)
test.save.grp3 <- gx.md.gait(grp3[, -3], mcdstart = TRUE)
```

```

## Allocate the synthetic data into the three reference groups
test.save.mvalloc <- gx.mvalloc(pcrit = 0.05, test.mvalloc.mat[, -3],
test.save.grp1, test.save.grp2, test.save.grp3)
## Display the results of the allocation
xyplot.tags(x, y, test.save.mvalloc$xalloc, cex = 0.75)
gx.mvalloc.print(test.save.mvalloc)

## Save the allocation as a csv file
gx.mvalloc.print(test.save.mvalloc, ifprint = FALSE,
file = " ")

## Clean-up and detach synthetic test data
rm(grp1)
rm(grp2)
rm(grp3)
rm(anom)
rm(test.mvalloc)
rm(test.save.grp1)
rm(test.save.grp2)
rm(test.save.grp3)
rm(test.save.mvalloc)
detach(test.mvalloc)

```

gx.mvalloc.closed	<i>Function for Allocation on the basis of Multivariate Data for closed Compositional Data</i>
-------------------	--

Description

Function to allocate individuals (observations, cases or samples) from closed compositional, geochemical, data sets into one of multiple reference groups (populations) on the basis of their Mahalanobis distances. If an individual's predicted probability of group membership (typicality) falls below a user defined 'cut-off', `pcrit`, the individual is allocated to an 'outlier bin'.

Usage

```
gx.mvalloc.closed(pcrit = 0.05, xx, ...)
```

Arguments

<code>pcrit</code>	the critical cut-off probability for group membership below which an individual will be classified as an 'outlier'. By default the critical probability of group membership is set to <code>pcrit = 0.05</code> .
<code>xx</code>	a n by p matrix containing the n individuals, with p variables determined on each, to be allocated, see Details below.
<code>...</code>	a list of objects saved from either function gx.md.gait.closed or gx.robmva.closed , containing the vectors of means and inverse covariance matrices for the 'reference' groups into which the individuals are to be classified.

Details

It is imperative that the data matrix `xx` contains no special codes, see Note below. It is also imperative that the variables in the reference groups and in the matrix `x` of individuals to be classified are identical in number and in the same order.

The allocations are made on the assumption that the covariance structures are inhomogeneous, i.e. that the population hyperellipsoids are of different size, shape and orientation in p -space.

Value

The following are returned as an object to be saved for display with `gx.mvalloc.print`:

<code>groups</code>	a list of the names of the <code>kk</code> reference groups.
<code>kk</code>	the number of reference groups passed to the function.
<code>n</code>	the number of individuals (observations, cases or samples) allocated.
<code>p</code>	the number of variables in the reference and allocated data.
<code>pcrit</code>	the critical cut-off probability for reference group membership.
<code>p_{gm}</code>	a vector of <code>kk</code> predicted probabilities of reference group memberships.
<code>xalloc</code>	the reference group, <code>1:kk</code> , that the individual was allocated into. All outliers, i.e. all <code>p_{gm}(1:kk) < crit</code> are allocated to group zero, \emptyset . Therefore <code>xalloc</code> will be in the range of $\emptyset:kk$.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed from the matrix `xx` prior to executing this function, see `ltdl.fix.df`. Any rows in the input data matrix `xx` with NAs are removed prior to computations.

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1990. A robust multivariate allocation procedure with applications to geochemical data. In Proc. Colloquium on Statistical Applications in the Earth Sciences (Eds F.P. Agterberg & G.F. Bonham-Carter). Geological Survey of Canada Paper 89-9, pp. 309-318.

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

See Also

[gx.md.gait.closed](#), [gx.robmva.closed](#), [gx.mvalloc.print](#), [ltdl.fix.df](#), [remove.na](#), [na.omit](#)

Examples

```

## Make test data available
data(ogrady)
attach(ogrady)
ogrady.grdr <- gx.subset(ogrady, Lith == "GRDR")
ogrady.grnt <- gx.subset(ogrady, Lith == "GRNT")
## Ensure all data are in the same units (mg/kg)
ogrady.grdr.2open <- ogrady.grdr[, c(5:14)]
ogrady.grdr.2open[, 1:7] <- ogrady.grdr.2open[, 1:7] * 10000
ogrady.grnt.2open <- ogrady.grnt[, c(5:14)]
ogrady.grnt.2open[, 1:7] <- ogrady.grnt.2open[, 1:7] * 10000
ogrady.2open <- ogrady[, c(5:14)]
ogrady.2open[, 1:7] <- ogrady.2open[, 1:7] * 10000

## Create reference data sets
ogrady.grdr.save <- gx.md.gait.closed(as.matrix(ogrady.grdr.2open),
mcdstart = TRUE)
ogrady.grnt.save <- gx.md.gait.closed(as.matrix(ogrady.grnt.2open),
mcdstart = TRUE)

## Allocate all 0'Grady granitoids
ogrady.mvalloc <- gx.mvalloc.closed(pcrit = 0.02, as.matrix(ogrady.2open),
ogrady.grdr.save, ogrady.grnt.save)

## Display list of outliers
gx.mvalloc.print(ogrady.mvalloc)

## Display allocations
ogrady.mvalloc$xalloc

## Save allocations as a csv file
gx.mvalloc.print(ogrady.mvalloc, ifprint = FALSE, file = " ")

## Clean-up and detach test data
rm(ogrady.grdr)
rm(ogrady.grnt)
rm(ogrady.grdr.2open)
rm(ogrady.grnt.2open)
rm(ogrady.2open)
rm(ogrady.grdr.save)
rm(ogrady.grnt.save)
rm(ogrady.mvalloc)
detach(ogrady)

```

gx.mvalloc.print

Function to display the results of Multivariate Allocation

Description

Function to extract and display the results from the saved object from `gx.mvalloc` or `gx.mvalloc.closed`. The function displays on the current device only those individuals (obser-

variations, cases or samples) whose predicted probability of reference group membership (typicality) was less than the value provided, `pcrit`, for all reference groups, i.e. the outliers. Alternately, the results can be saved as a '.csv' file for viewing with a spreadsheet program and any subsequent post-processing.

Usage

```
gx.mvalloc.print(save, ifprint = TRUE, unalloc = TRUE, file = NULL)
```

Arguments

<code>save</code>	an object saved from <code>gx.mvalloc</code> .
<code>ifprint</code>	by default the 'outliers', i.e. individuals classified into group 'zero', are displayed on the current device. The display consists of the input matrix row numbers together with the predicted probabilities of reference group membership for the 1:kk reference groups. To suppress the display set <code>ifprint = FALSE</code> .
<code>unalloc</code>	by default, <code>unalloc = TRUE</code> , individuals that were not allocated to one of the reference groups are displayed. To suppress displaying these individuals, set <code>unalloc = FALSE</code> .
<code>file</code>	the name of the folder where the file is to be saved, a '/' is appended before the synthesized file name, details of which are displayed on the current device. If no file is specified, the files are saved to the Working Directory, see Details below.

Details

The full table of Mahalanobis distances and group membership probabilities may be saved as a '.csv' file. If a folder name has been provided in `file`, a '/' is appended prior a file name synthesized from the name of the saved object concatenated with '_mvalloc.csv', see Example below. If `file = ""` or `file = " "` the file will be saved to the Working Directory. The file name for the .csv file is displayed on the current device, and all other output is suppressed. The output file can be inspected with spread sheet software or a viewer of the user's choice.

Note

Included in the display on the current device are the names of the `kk` reference group objects supplied to `gx.mvalloc` together with the value of `pcrit`.

Author(s)

Robert G. Garrett

See Also

[gx.mvalloc](#), [gx.mvalloc.closed](#)

Examples

```

## Make test data available
data(ogrady)
attach(ogrady)
ogrady.grdr <- gx.subset(ogrady, Lith == "GRDR")
ogrady.grnt <- gx.subset(ogrady, Lith == "GRNT")
## Ensure all data are in the same units (mg/kg)
ogrady.grdr.2open <- ogrady.grdr[, c(5:14)]
ogrady.grdr.2open[, 1:7] <- ogrady.grdr.2open[, 1:7] * 10000
ogrady.grnt.2open <- ogrady.grnt[, c(5:14)]
ogrady.grnt.2open[, 1:7] <- ogrady.grnt.2open[, 1:7] * 10000
ogrady.2open <- ogrady[, c(5:14)]
ogrady.2open[, 1:7] <- ogrady.2open[, 1:7] * 10000

## Create reference data sets
ogrady.grdr.save <- gx.md.gait(ilr(as.matrix(ogrady.grdr.2open)),
mcdstart = TRUE)
ogrady.grnt.save <- gx.md.gait(ilr(as.matrix(ogrady.grnt.2open)),
mcdstart = TRUE)

## Allocate all 0'Grady granitoids
ogrady.mvalloc <- gx.mvalloc(pcrit = 0.02, ilr(as.matrix(ogrady.2open)),
ogrady.grdr.save, ogrady.grnt.save)

## Display list of outliers
gx.mvalloc.print(ogrady.mvalloc)

## Save allocations as a .csv file, not executed
## gx.mvalloc.print(ogrady.mvalloc, ifprint = FALSE, file = "D://R_work//Project3")

## Clean-up and detach test data
rm(ogrady.grdr)
rm(ogrady.grnt)
rm(ogrady.grdr.2open)
rm(ogrady.grnt.2open)
rm(ogrady.2open)
rm(ogrady.grdr.save)
rm(ogrady.grnt.save)
rm(ogrady.mvalloc)
detach(ogrady)

```

gx.ngr.ltdl

Displays a Count of Negative Values and NAs for a Matrix or Data Frame

Description

Displays a table of the counts of negative values, indicating less than the detection limit, and NAs to support the preparation of a NGR Report Table of summary statistics. The table for numerical columns in the matrix or data frame displays the variable name, the number of NAs and the number of negative values, and the number of times the value in coded occurs.

Usage

```
gx.ngr.ltdl(xmat, vars, coded = -9999)
```

Arguments

xmat	name of the matrix or data frame.
vars	the indices, or names, of the columns of the matrix or data frame for the variables for which counts are to be displayed. By default, i.e. vars is undefined, counts for all numeric columns are displayed.
coded	a coded value, default -9999 whose occurrence will be counted. In some data sets values greater than the upper limit of quantification are set to such a coded value.

Examples

```
## Make test data available
data(fix.test)

## Generate table of counts
gx.ngr.ltdl(fix.test)
```

 gx.ngr.skew

Estimate the Skewness of a Data Vector

Description

Estimates the skew of a data vector for [gx.ngr.stats](#) to be used by [gx.ngr.summary](#) to output a NGR Table of summary statistics as a '.csv' file.

Usage

```
gx.ngr.skew(xx)
```

Arguments

xx	vector for which the skewness is required.
----	--

Details

The formula used, $(m_3 / m_2^{1.5}) * ((n-1)/n)^{1.5}$, where m_3 and m_2 are the third and second moments about the mean, respectively, provides an estimate similar to that obtained by Minitab and BMDP.

Value

skew	the estimate of the skewness of the data vector passed to the function.
------	---

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation.

Author(s)

Robert G. Garrett

Examples

```
## Make test data available
data(sind)
attach(sind)

## Estimate and display the skewness of the Zn data
skew <- gx.ngr.skew(Zn)
skew

## Alternately
skew <- gx.ngr.skew(rnorm(0,1,1000))
skew

## Detach test data
detach(sind)
```

gx.ngr.stats

Computes Summary Statistics for a NGR Report Table

Description

Builds a vector of summary statistics for [gx.ngr.summary](#) to output a NGR Report table of summary statistics as a '.csv' file from estimates made by [gx.stats](#) and [gx.ngr.skew](#).

Usage

```
gx.ngr.stats(xx)
```

Arguments

xx vector for which NGR Report Table summary statistics are required.

Value

table	the computed summary statistics to be used in function gx.ngr.summary . The list returned, table, is a 29-element vector, see below:
[1]	the sample size, N.
[2]	the number of NAs in the input vector.
[3]	the data (sample) Mean.
[4]	the data (sample) Standard Deviation (SD).
[5]	the data (sample) Skew.
[6]	the Coefficient of Variation as a percentage (CV%).
[7]	the data (sample) Geometric Mean.
[8]	the data (sample) Median.
[9]	the data (sample) Median Absolute Deviation (MAD).
[10]	the Robust Coefficient of Variation as a percentage (RCV%).
[11:20]	the minimum value, and the 1st, 2nd, 5th, 10th, 20th, 25th (Q1), 30th, 40th and 50th (Q2) percentiles.
[21:29]	the 60th, 70th, 75th (Q3), 80th 90th, 95th, 98th and 99th percentiles and the maximum value.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation in function [gx.stats](#). Display of the number of NAs found by function `remove.na` is suppressed in `remove.na` as the information is included in the display from this function.

Author(s)

Robert G. Garrett

See Also

[gx.summary1](#), [gx.summary](#), [gx.stats](#), [ltdl.fix.df](#), [remove.na](#), [gx.summary2](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Generate and display the results for Zn
table <- gx.ngr.stats(Zn)
table

## Detach test data
detach(sind)
```

gx.ngr.summary	<i>Saves a NGR Report Summary Statistics Table to a .csv File</i>
----------------	---

Description

Prepares a .csv file with the standard set of NGR Report summary statistics for further processing with a spread sheet program. The table includes: N (data set size), number of NAs, mean, SD, skew, CV%, geometric mean, median, MAD, robust CV%, and the minimum and maximum values and 17 intermediate percentiles.

Usage

```
gx.ngr.summary(xmat, vars, banner = deparse(substitute(xmat)),
file = NULL)
```

Arguments

xmat	name of the matrix or data frame.
vars	the indices, or names (see Example), of the columns of the matrix or data frame for the variables whose summary statistics are to be displayed.
banner	by default the character string for xmat, the input matrix, is used in the file name. An alternate title can be displayed with banner = "text string", see Details below.
file	the name of the folder where the files are to be saved, a '/' is appended before the synthesized file name, details of which are displayed on the current device. If no file is specified, the files are saved to the Working Directory.

Details

The file name is synthesized by concatenating the input matrix name and '_NGR_Summary.csv', and prefixed by sQuote/. If a data frame is the data source, the synthesized file name will be '/dataframe_NGR_Summary.csv', as it is not possible to obtain the text string for the data frame name from xmat. In such cases the user should rename the .csv file more appropriately after it has been saved. If a folder name has been provided in file, for example, file = "D://R_work//Project3", a '/' is appended prior to the synthesized file name.

The file name for the .csv file is displayed on the current device, and all other output is suppressed. The output file can be inspected with spread sheet software or a viewer of the user's choice.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation in function [gx.stats](#). Display of the number of NAs found by function [remove.na](#) is suppressed in [remove.na](#) as the information is included in the display from this function that is prepared by [gx.ngr.stats](#).

For a more extensive summary statistics display a variable at a time, see [gx.summary2](#).

Author(s)

Robert G. Garrett

See Also[gx.ngr.stats](#), [gx.stats](#), [gx.ngr.skew](#), [ltdl.fix.df](#), [remove.na](#), [gx.summary1](#), [gx.summary2](#)**Examples**

```
## Make test data available
data(sind.mat2open)

## Prepare and export a .csv file
gx.ngr.summary(sind.mat2open)
```

gx.pairs4parts

*Display a Graphical Matrix for Parts of a Compositional Data Set***Description**

Displays a graphical matrix of log₁₀ scaled x-y plots in the upper triangle and boxplots of the ilr transforms in the lower triangle for the parts of a compositional matrix. The robust ilr stability (Filzmoser et al., 2010) for each x-y pair is displayed as the boxplot title.

Usage

```
gx.pairs4parts(xx, cex = 2, ifwarn = TRUE, ...)
```

Arguments

xx	a matrix, or sub-matrix, of parts from a compositional data set.
cex	by default the size of the text of the variable names in the diagonal of the graphical matrix. By default <code>cex = 2</code> , and may be changed if required.
ifwarn	by default <code>ifwarn = TRUE</code> which generates a reminder/warning that when carrying out analyses of compositional data all data must be in the same measurement units. The message can be suppressed by setting <code>ifwarn = FALSE</code> .
...	further arguments to be passed to plot or bplot. For example, the size of the axis scale annotation can be change by setting <code>cex.axis</code> and the size of the plot title by setting <code>cex.main</code> . For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors (rows) containing NAs are removed prior to computation.

Author(s)

Robert G. Garrett

References

Filzmoser, P, Hron, K. and Reimann, C., 2010. The bivariate statistical analysis of environmental (compositional) data. *Science of the Total Environment*, 408(19):4230-4238.

See Also[ltdl.fix.df](#), [remove.na](#), [bxplot](#), [ilr.stab](#)**Examples**

```
## Make test data available
data(sind.mat2open)

## Display 'pairs' plots for a set, or sub-set, of parts of a
## compositional data matrix
gx.pairs4parts(sind.mat2open)
```

`gx.pearson`*Display Pearson Correlation Coefficients and their Significances*

Description

The function computes Pearson product moment correlation coefficients and places them in the upper triangle of a printed matrix displayed on the current device, the probabilities that the coefficients are not due to chance (H_0 : Coefficient = 0) are printed in the lower triangle. The diagonal is filled with NAs to visually split the two triangles.

Usage

```
gx.pearson(xx, log = FALSE, ifclr = FALSE, ifwarn = TRUE)
```

Arguments

<code>xx</code>	a matrix of numeric data.
<code>log</code>	if <code>log = TRUE</code> the data are log10 transformed prior to computation of the Pearson coefficients. The default is no transformation.
<code>ifclr</code>	if <code>ifclr = TRUE</code> the data are Centred Log-Ratio transformed prior to the computation of the Pearson Coefficients. The default is no transformation.
<code>ifwarn</code>	by default <code>ifwarn = TRUE</code> which generates a reminder/warning that when carrying out a centred log-ratio transformation all the data must be in the same measurement units. The message can be suppressed by setting <code>ifwarn = FALSE</code> .

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors (rows) containing NAs are removed prior to computation.

This function is not recommended for use with closed compositional data sets, i.e. geochemical analyses, unless correlations are sought between a non-compositional variable and individual compositional variables. If it is used with compositional data, it is highly recommended that `ifclr` be set to TRUE to remove the effects of closure and display the ‘true’ inter-element variability. However, different groups of elements, subsets, of a data set will yield different inter-element correlations for the same pair of elements due to the nature of the `clr` transform. When carrying out a centred log-ratio transformation it is essential that the data are all in the same measurement units, and by default a reminder/warning is display if the data are centred log-ratio transformed, see `ifwarn` above.

For working with compositional data sets functions [gx.vm](#) and [gx.sm](#) are recommended. For visual displays see [gx.pairs4parts](#) and [gx.plot2parts](#).

When a centred log-ratio transformation is undertaken the `log` ‘switch’ is ignored.

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [remove.na](#), [clr](#), [sind.mat2open](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Compute Pearson correlation coefficients
gx.pearson(sind.mat2open)

## Compute Pearson correlation coefficients following
## a logarithmic transformation
gx.pearson(sind.mat2open, log = TRUE)

## Compute Pearson correlation coefficients following
## a centred log-ratio transformation
gx.pearson(sind.mat2open, ifclr = TRUE)
```


Description

Displays a panel of four plots for a pair of parts from a compositional data set. The displays consist of a log₁₀ scaled x-y plot, a boxplot of the corresponding values of $\text{ilr}(x,y)$ annotated with the robust ilr stability measure, and sequential index and ECDF plots of the ilr values. The display is based on those used in Filzmoser et al. (2010).

Usage

```
gx.plot2parts(xx1, xx2, x1lab = deparse(substitute(xx1)),
             x2lab = deparse(substitute(xx2)), cex = 0.8, ifwarn = TRUE, ...)
```

Arguments

xx1	a column vector from a matrix or data frame of compositional data, xx1[1], ..., xx1[n].
xx2	another column vector from the matrix or data frame of compositional data, xx2[1], ..., xx2[n]. xx1 and xx2 must be of identical length, n.
x1lab	the x-axis title, by default the variable name, deparse(substitute(xx1)). It is often desirable to replace this with a more informative title, e.g., x1lab = "Cu (mg/kg) in stream sediment".
x2lab	the y-axis title, by default the variable name, deparse(substitute(xx2)). It is often desirable to replace this with a more informative title, e.g., x2lab = "Zn (mg/kg) in stream sediment".
cex	by default the size of the text for data set size, N, and the robust ilr stability is set to 80%, i.e. $cex = 0.8$, and may be changed if required.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out analyses of compositional data all data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.
...	further arguments to be passed to methods. For example, the size of the axis scale annotation can be change by setting cex.axis, the size of the axis titles by setting cex.lab, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#). Any data vectors (rows) containing NAs are removed prior to computation.

Author(s)

Robert G. Garrett

References

Filzmoser, P, Hron, K. and Reimann, C., 2010. The bivariate statistical analysis of environmental (compositional) data. *Science of the Total Environment*, 408(19):4230-4238.

See Also

[ltdl.fix.df](#), [remove.na](#), [bxplot](#), [gx.ecdf](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Display
gx.plot2parts(Cu, Zn)

## Display with alternate xy-plot titling
gx.plot2parts(Cu, Zn, x1lab = "Cu (mg/kg) in stream sediment",
x2lab = "Zn (mg/kg) in stream sediment")

## Detach test data
detach(sind)
```

gx.quantile

Estimate the Quantile for a Specified Fractile

Description

Estimates and displays the quantile for a specified fractile of a data set by linear interpolation from the ranked data. If the function is run as `temp <- gx.quantile(xx, f)` the quantile is not displayed, but retained in `temp` for subsequent use or display.

Usage

```
gx.quantile(xx, f, display = TRUE)
```

Arguments

<code>xx</code>	the data set for which the quantile is to be estimated.
<code>f</code>	the fractile for which the quantile is required, must be in the range zero to 1.
<code>display</code>	the default is to display the fractile and estimated quantile on the current device. If no display is required, set <code>display = FALSE</code> .

Value

<code>q</code>	the estimated quantile.
----------------	-------------------------

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to preparing the boxplots.

The result is an approximation, and the result from the quantile function will likely differ by some small amount.

Author(s)

Based on a script shared on S-News by Nick Ellis, April 2002

See Also

[ltdl.fix.df](#), [remove.na](#), [gx.fractile](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Estimate the 80th percentile, f = 0.8
gx.quantile(As, 0.8)
temp <- gx.quantile(As, 0.8)
temp

## Clean-up and detach test data
rm(temp)
detach(kola.o)
```

gx.rma

Estimate the Coefficients of the Reduced Major Axis

Description

Function to estimate the coefficients and their standard errors of the Reduced Major Axis, the case of orthogonal regression, and also known as total least squares or errors in variables regression. The procedure is based on the methodology described in Miller and Kahn (1962). Function will also prepare a Youden plot.

Usage

```
gx.rma(xx1, xx2, x1lab = NULL, x2lab = NULL, log = FALSE, if.plot = FALSE,
if.rma = FALSE, if.coeffs = FALSE, ...)
```

Arguments

xx1	the name of the first independent variable, or a n by 2 matrix of independent variables, for instance from <code>alts2dups</code> .
xx2	the name of the second independent variable, omit if xx1 is a matrix.
x1lab	a title for the first independent variable, the default is the variable name, <code>deparse(substitute(xx1))</code> or the name from the first column of the n by 2 matrix. It is often desirable to replace the default title of the input variable name text string with a more informative title, e.g., <code>x1lab = "Magnetic Susceptibility - Measurement 1"</code> .
x2lab	a title for the second independent variable, the default is the variable name, <code>deparse(substitute(xx2))</code> or the name from the second column of the n by 2 matrix. It is often desirable to replace the default title of the input variable name text string with a more informative title, e.g., <code>x2lab = "Magnetic Susceptibility - Measurement 2"</code> .
log	if a logarithmic transformation (base 10) of the data is required to meet homogeneity of variance considerations (i.e. severe heteroscedasticity) set <code>log = TRUE</code> . This is also advisable if the range of the observations exceeds 1.5 orders of magnitude.
if.plot	if a x-y plot of the independent variables is required set <code>if.plot = TRUE</code> . The plot is equi-scaled and the 1:1 line is added. If <code>if.rma = TRUE</code> <code>if.plot</code> will be set to TRUE.
if.rma	if the Reduced Major Axis is to be added to the plot set <code>if.rma = TRUE</code> .
if.coeffs	if the coefficients of the Reduced Major Axis and the result of the test for the RMA being (0,1) are to be displayed on the plot, set <code>if.coeffs = TRUE</code> . The cursor will be activated, and the upper left corner of the text block may be placed where most appropriate, then 'left button' the pointing device.
...	further arguments to be passed to methods concerning the plot. For example, if it is required to add a title, add <code>main = "Youden Plot"</code> .

Value

A list comprising of:

alen	the data set size.
mean	a two-element vector with the means of x1 and x2.
sd	a two-element vector with the standard deviations of x1 and x2.
corr	the Pearson correlation coefficient for x1 and x2.
a0	the intercept of the reduced major axis.
a1	the slope of the reduced major axis.
sea0	the standard error of the intercept estimate.
aea1	the standard error of the slope estimate.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data pairs, xx1, xx2, containing any NAs are omitted from the calculations.

If a log transformation is undertaken and any less than or equal to zero values occur in the data the function will halt with a warning to that effect.

The coefficients may be used to plot the RMA on a x-y plot of the two measures, see example below.

Author(s)

Robert G. Garrett

References

Miller, R.L. and Kahn, J.S., 1962. Statistical Analysis in the Geological Sciences, John Wiley & Sons, New York, U.S.A., 483 p. Specifically pp. 204-209.

See Also

[ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test available
data(ms.data1)
attach(ms.data1)

## Estimate RMA coefficients for duplicate measurements on rock samples
gx.rma(MS.1, MS.2, x1lab = "MS - 1", x2lab = "MS - 2", log = TRUE)

## Display an x-y plot of the data and the RMA
gx.rma(MS.1, MS.2, x1lab = "MS - 1", x2lab = "MS - 2", log = TRUE,
if.plot = TRUE, if.rma = TRUE)

## Detach test data
detach(ms.data1)

## Using function alts2dups
data(ms.data3)
attach(ms.data3)
gx.rma(alts2dups(MS, ifalt = TRUE), log = TRUE, if.plot = TRUE,
if.rma = TRUE)

## Detach test data
detach(ms.data3)
```

gx.robmva	<i>Function to undertake a Robust Exploratory Multivariate Data Analysis</i>
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Description

The function carries out a robust Principal Components Analysis (PCA) and estimates the Mahalanobis distances for a non-compositional dataset and places them in an object to be saved and post-processed for display and further manipulation. For closed compositional, geochemical, data use function [gx.robmva.closed](#). Robust procedures are used, 'MCD', 'MVE' or user supplied weights, for classical procedures see [gx.mva](#). For results display see [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#) and [gx.md.print](#). For Kaiser varimax rotation see [gx.rotate](#).

Usage

```
gx.robmva(xx, proc = "mcd", wts = NULL, main = deparse(substitute(xx)))
```

Arguments

xx	a n by p data matrix to be processed.
proc	by default proc = "mcd" for the Minimum Covariance Determinant (MCD) robust procedure. Setting proc = "mve" results in the Minimum Volume Ellipsoid (MVE) procedure being used. If p > 50 the MVE procedure is used. See wts below.
wts	by default wts = NULL and the MCD or MVE estimation procedures will be used. If, however, a vector of n zero or 1 weights are supplied these will be used for robust estimation and the value of proc ignored.
main	by default the name of the object xx, main = deparse(substitute(xx)), it may be replaced by the user, but this is not recommended, see Details below.

Details

If main is undefined the name of the matrix object passed to the function is used to identify the object. This is the recommended procedure as it helps to track the progression of a data analysis. Alternate plot titles are best defined when the saved object is passed to [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#) or [gx.md.plot](#) for display. If no plot title is required set main = " ", or if a user defined plot title is required it may be defined, e.g., main = "Plot Title Text".

The variances of the robust Principal Component scores are displayed, in a non-robust PCA these decrease with increasing component rank. However, in a robust PCA this may not be the case, and lower-order scores with high variances are often worthy of further inspection.

Value

The following are returned as an object to be saved for subsequent display, etc.:

main	by default (recommended) the input data matrix name.
input	the data matrix name, <code>input = deparse(substitute(xx))</code> , retained to be used by post-processing display functions.
proc	the robust procedure used, the value of <code>proc</code> will be "mcd", "mve" or "wts".
n	the total number of individuals (observations, cases or samples) in the input data matrix.
nc	the number of individuals remaining in the 'core' data subset following the robust estimation, i.e. the sum of those individuals with <code>wts = 1</code> .
p	the number of variables on which the multivariate operations were based.
ifilr	flag for <code>gx.md.plot</code> , set to FALSE.
matnames	the row numbers or identifiers and column headings of the input matrix.
wts	the vector of weights for the <code>n</code> individuals arising from the robust estimation of the covariance matrix and means.
mean	the length <code>p</code> vector the weighted means for the variables.
cov	the <code>p</code> by <code>p</code> weighted covariance matrix for the <code>n</code> by <code>p</code> data matrix.
sd	the length <code>p</code> vector of weighted standard deviations for the variables.
snd	the <code>n</code> by <code>p</code> matrix of weighted standard normal deviates.
r	the <code>p</code> by <code>p</code> matrix of weighted Pearson product moment correlation coefficients.
eigenvalues	the vector of <code>p</code> eigenvalues of the scaled Pearson robust correlation matrix for RQ analysis, see Grunsky (2001).
econtrib	the vector of <code>p</code> robustly estimated eigenvalues each expressed as a percentage of the sum of the eigenvalues.
eigenvectors	the <code>n</code> by <code>p</code> matrix of robustly estimated eigenvectors.
rload	the <code>p</code> by <code>p</code> matrix of robust Principal Component (PC) loadings.
rcr	the <code>p</code> by <code>p</code> matrix containing the percentages of the variability of each variable (rows) expressed in each robust PC (columns).
rqscore	the <code>n</code> by <code>p</code> matrix of the <code>n</code> individuals scores on the <code>p</code> robust PCs.
vcontrib	a vector of <code>p</code> variances of the columns of <code>rqscore</code> .
pvcontrib	the vector of <code>p</code> variances of the columns of <code>rqscore</code> expressed as percentages. This is a check on vector <code>econtrib</code> , the values should be identical for a classical PCA. However, for robust PCAs this is not so as the trimmed individuals from the robust estimation have been re-introduced. As a consequence <code>pvcontrib</code> can be very different from <code>econtrib</code> . The plotting of PCs containing high proportions of the variance in robust PCAs can be useful for identifying outliers
cpvcontrib	the vector of <code>p</code> cumulative sums of <code>pvcontrib</code> , see above.
md	the vector of <code>n</code> robust Mahalanobis distances (MDs) for the <code>n</code> by <code>p</code> input matrix.
ppm	the vector of <code>n</code> robust predicted probabilities of population membership, see Garrett (1990).
epm	the vector of <code>n</code> robust empirical Chi-square probabilities for the MDs.
nr	the number of PCs that have been rotated. At this stage of a data analysis <code>nr = NULL</code> in order to control PC plot axis labelling.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with NAs are removed prior to computations. In the instance of a compositional data opening transformation NAs have to be removed prior to undertaking the transformation, see [na.omit](#), [where.na](#) and [remove.na](#). When that procedure is followed the opening transformations may be executed on calling the function, see Examples below.

Passing a set of weights from an investigation with [gx.md.gait](#) or on the basis of some prior knowledge permits the use of a [clr](#) transformation. In this instance a Moore-Penrose inverse is computed and used for the estimation of Mahalanobis distances. See example below. With reference to weights based on prior knowledge, the weights are not necessarily constrained to be '0' or '1', intermediate values may be employed.

Executing a [clr](#) transformation leads to both collinearity and singularity such that neither a PCA can be undertaken or Mahalanobis distances be estimated. The function fails - do not use with a [clr](#) transformation.

Executing a [ilr](#) transformation permits the estimation of both Principal Components and Mahalanobis distances and associated probabilities through the use of (p-1) synthetic variables. However, in that instance the loadings of the (p-1) synthetic variables will be plotted by [gx.rqpca.plot](#) rather than the loadings for the elements.

Warnings are generated when the number of individuals (observations, cases or samples) falls below 5*p, and additional warnings when the number of individuals falls below 3*p. At these low ratios of individuals to variables the shape of the p-space hyperellipsoid is difficult to reliably define, and therefor the results may lack stability. These limits 5*p and 3*p are generous, the latter especially so; many statisticians would argue that the number of individuals should not fall below 9*p, see Garrett (1993).

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1990. A robust multivariate allocation procedure with applications to geochemical data. In Proc. Colloquium on Statistical Applications in the Earth Sciences (Eds F.P. Agterberg & G.F. Bonham-Carter). Geological Survey of Canada Paper 89-9, pp. 309-318.

Garrett, R.G., 1993. Another cry from the heart. Explore - Assoc. Exploration Geochemists Newsletter, 81:9-14.

Grunsky, E.C., 2001. A program for computing RQ-mode principal components analysis for S-Plus and R. Computers & Geosciences, 27(2):229-235.

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

See Also

[ltdl.fix.df](#), [remove.na](#), [na.omit](#), [gx.rqpca.screplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#), [gx.md.print](#), [gx.robmva.closed](#), [gx.rotate](#)

Examples

```
## Generate a population of synthetic bivariate normal data
grp1 <- mvrnorm(100, mu = c(40, 30), Sigma = matrix(c(6, 3, 3, 2), 2, 2))
grp1 <- cbind(grp1, rep(1, 100))
## Generate a set of six (6) outliers
anom <- matrix(c(43, 34, 50, 37, 47, 30, 27, 29, 35, 33, 32, 25), 6, 2)
anom <- cbind(anom, rep(2, 6))
## Bind the test data together and display the test data
test.robmva.mat <- rbind(grp1, anom)
test.robmva <- as.data.frame(test.robmva.mat)
dimnames(test.robmva)[[2]] <- c("x", "y", "grp")
attach(test.robmva)
xyplot.tags(x, y, dimnames(test.robmva)[[1]], cex = 0.75)

## Generate gx.robmva saved object
save.rob <- gx.robmva(as.matrix(test.robmva[, c(1:2)]))
## Display saved object with alternate main titles
gx.rqpca.screepplot(save.rob, main = "Bivariate synthetic data")
gx.rqpca.plot(save.rob, cex.lab = 0.8, rowids = TRUE,
main = "Bivariate synthetic data")
gx.md.plot(save.rob, cex.main = 0.9, cex.lab = 0.8, cex.axis = 0.8,
main = "Bivariate synthetic data")
gx.md.print(save.rob, pcut = 0.05)

## Clean-up and detach test data
rm(grp1)
rm(anom)
rm(test.robmva.mat)
rm(test.robmva)
rm(save.rob)
detach(test.robmva)
```

gx.robmva.closed

Function to undertake a Robust Closed Data Multivariate EDA

Description

The function carries out a robust Principal Components Analysis (PCA) and estimates the Mahalanobis distances for a closed compositional, geochemical, dataset and places the results in an object to be saved and post-processed for display and further manipulation. Robust procedures are used, ‘MCD’, ‘MVE’ or user supplied weights, for classical procedures see [gx.mva.closed](#), or for non-compositional data and robust procedures see [gx.robmva](#). For results display see [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#) and [gx.md.print](#). For Kaiser varimax rotation see [gx.rotate](#).

Usage

```
gx.robmva.closed(xx, proc = "mcd", wts = NULL,
main = deparse(substitute(xx)))
```

Arguments

xx	a n by p data matrix to be processed.
proc	by default proc = "mcd" for the Minimum Covariance Determinant (MCD) robust procedure. Setting proc = "mve" results in the Minimum Volume Ellipsoid (MVE) procedure being used. If $p > 50$ the MVE procedure is used. See wts below.
wts	by default wts = NULL and the MCD or MVE estimation procedures will be used. If, however, a vector of n zero or 1 weights are supplied these will be used for robust estimation and the value of proc ignored.
main	by default the name of the object xx, main = deparse(substitute(xx)), it may be replaced by the user, but this is not recommended, see Details below.

Details

The data are initially isometrically log-ratio transformed and a robust covariance matrix and vector of means estimated, by either the Minimum Covariance Determinant (MCD) or Minimum Volume Ellipsoid (MVE) procedures, or on the basis of a vector of user supplied weights. The Mahalanobis distances are computed on the basis of the ilr transformed data. The ilr transformed data and robust estimates, including the inverse of the covariance matrix, are then back-transformed to the centred log-ratio basis and a Principal Components Analysis (PCA) undertaken (see Filzmoser, et al., 2009), permitting the results to be interpreted in the original variable space.

The variances of the robust Principal Component scores are displayed, in a non-robust PCA these decrease with increasing component rank. However, in a robust PCA this may not be the case, and lower-order scores with high variances are often worthy of further inspection.

If main is undefined the name of the matrix object passed to the function is used to identify the object. This is the recommended procedure as it helps to track the progression of a data analysis. Alternate plot titles are best defined when the saved object is passed to `gx.rqpca.plot`, `gx.rqpca.screeplot` or `gx.md.plot` for display. If no plot title is required set main = " ", or if a user defined plot title is required it may be defined, e.g., main = "Plot Title Text".

Value

The following are returned as an object to be saved for subsequent display, etc.:

main	by default (recommended) the input data matrix name.
input	the data matrix name, input = deparse(substitute(xx)), retained to be used by post-processing display functions.
proc	the robust procedure used, the value of proc will be "mcd", "mve" or "wts".
n	the total number of individuals (observations, cases or samples) in the input data matrix.
nc	the number of individuals remaining in the 'core' data subset following the robust estimation, i.e. the sum of those individuals with wts = 1.
p	the number of variables on which the multivariate operations were based.
ifilr	flag for gx.md.plot, set to TRUE.
matnames	the row numbers or identifiers and column headings of the input matrix.

wt	the vector of weights for the n individuals arising from the robust estimation of the covariance matrix and means.
mean	the length p vector of clr-based weighted means for the variables.
cov	the p by p weighted clr-based covariance matrix for the n by p data matrix.
cov.inv	the p by p weighted clr-based inverse of the covariance matrix, for use by function <code>gx.mvalloc.closed</code> .
sd	the length p vector of weighted clr-based standard deviations for the variables.
snd	the n by p matrix of clr-based weighted standard normal deviates.
r	the p by p matrix of weighted clr-based Pearson product moment correlation coefficients.
eigenvalues	the vector of p eigenvalues of the scaled clr-based Pearson robust correlation matrix for RQ analysis, see Grunsky (2001).
econtrib	the vector of p robustly estimated eigenvalues each expressed as a percentage of the sum of the eigenvalues.
eigenvectors	the n by p matrix of clr-based robustly estimated eigenvectors.
rload	the p by p matrix of robust clr-based Principal Component (PC) loadings.
rcr	the p by p matrix containing the percentages of the variability of each variable (rows) expressed in each robust clr-based PC (columns).
rqscore	the n by p matrix of the n individuals scores on the p robust clr-based PCs.
vcontrib	a vector of p variances of the columns of <code>rqscore</code> .
pvcontrib	the vector of p variances of the columns of <code>rqscore</code> expressed as percentages. This is a check on vector <code>econtrib</code> , the values should be identical for a classical PCA. However, for robust PCAs this is not so as the trimmed individuals from the robust estimation have been re-introduced. As a consequence <code>pvcontrib</code> can be very different from <code>econtrib</code> . The plotting of PCs containing high proportions of the variance in robust PCAs can be useful for identifying outliers.
cpvcontrib	the vector of p cumulative sums of <code>pvcontrib</code> , see above.
md	the vector of n robust ilr-based Mahalanobis distances (MDs) for the n by p input matrix.
ppm	the vector of /coden robust ilr-based predicted probabilities of population membership, see Garrett (1990).
epm	the vector of n robust ilr-based empirical Chi-square probabilities for the MDs.
nr	the number of PCs that have been rotated. At this stage of a data analysis <code>nr = NULL</code> in order to control PC plot axis labelling.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with NAs are removed prior to computations. In the instance of a compositional data opening transformation NAs have to be removed prior to undertaking the transformation, see [na.omit](#), [where.na](#) and [remove.na](#). When that procedure is followed the opening transformations may be executed on calling the function, see Examples below.

Warnings are generated when the number of individuals (observations, cases or samples) falls below $5 \cdot p$, and additional warnings when the number of individuals falls below $3 \cdot p$. At these low ratios of individuals to variables the shape of the p -space hyperellipsoid is difficult to reliably define, and therefore the results may lack stability. These limits $5 \cdot p$ and $3 \cdot p$ are generous, the latter especially so; many statisticians would argue that the number of individuals should not fall below $9 \cdot p$, see Garrett (1993).

Author(s)

Robert G. Garrett

References

Filzmoser, P., Hron, K., Reimann, C. and Garrett, R., 2009. Robust factor analysis for compositional data. *Computers & Geosciences*, 35(9):1854-1861.

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Garrett, R.G., 1993. Another cry from the heart. *Explore - Assoc. Exploration Geochemists Newsletter*, 81:9-14.

Grunsky, E.C., 2001. A program for computing RQ-mode principal components analysis for S-Plus and R. *Computers & Geosciences*, 27(2):229-235.

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. *Statistical Data Analysis Explained: Applied Environmental Statistics with R*. John Wiley & Sons, Ltd., 362 p.

See Also

[ltdl.fix.df](#), [remove.na](#), [na.omit](#), [orthonorm](#), [gx.rqpca.screepplot](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#), [gx.rqpca.print](#), [gx.md.plot](#), [gx.md.print](#), [gx.robmva](#), [gx.rotate](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Generate gx.robmva.closed object
sind.save <- gx.robmva.closed(sind.mat2open)

## Display Mahalanobis distances
gx.md.plot(sind.save)

## Display default PCA results
gx.rqpca.screepplot(sind.save)
gx.rqpca.loadplot(sind.save)

## Display appropriately annotated results
gx.md.plot(sind.save,
main = "Howarth & Sinding-Larsen\nStream Sediments, Opened Data",
cex.main=0.8)
gx.rqpca.screepplot(sind.save,
```

```

main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")
gx.rqpca.plot(sind.save,
main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")
gx.rqpca.plot(sind.save, rowids = TRUE,
main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")
sind.save$pvcontrib
gx.rqpca.plot(sind.save, v1 = 3, v2 =4, rowids = TRUE,
main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")

## Display Kaiser Varimax rotated (nrot = 4) results
sind.save.rot4 <- gx.rotate(sind.save, 4)
gx.rqpca.plot(sind.save.rot4,
main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")
gx.rqpca.plot(sind.save.rot4, rowids = TRUE,
main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")
gx.rqpca.plot(sind.save.rot4, v1 = 3, v2 =4, rowids = TRUE,
main = "Howarth & Sinding-Larsen Stream Sediments\nOpened Data")

## Clean-up
rm(sind.save)
rm(sind.save.rot4)

```

gx.rotate

Function to Perform a Kaiser Varimax Rotation

Description

Function to perform a Kaiser Varimax rotation on Principal Component (PCA) loadings and scores in an object saved from [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#) or [gx.robmva.closed](#).

Usage

```
gx.rotate(save, nrot = 2)
```

Arguments

save	a saved object from the execution of function gx.mva , gx.mva.closed , gx.robmva , or gx.robmva.closed .
nrot	the number of component loadings to be rotated, by default the first two components are rotated, nrot = 2.

Value

The value of nr is modified in, and the following are appended to, the object that was saved from [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), or [gx.robmva.closed](#):

nr	modified to equal the number of components rotated.
vload	the new loadings after Varimax rotation.
vscore	the new scores after Varimax rotation.

vvcontrib	the contribution of the rotated Varimax component to the total data variability.
pvvcontrib	the contribution of the rotated Varimax component to the total data variability as a percentage.
cpvvcontrib	the cumulative contribution of the rotated Varimax component to the total data variability as a percentage.

Author(s)

Robert G. Garrett

References

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p.

See Also

[gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [varimax](#), [gx.rqpca.loadplot](#), [gx.rqpca.plot](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Save PCA results and display biplots before and after Varimax rotation
sind.save <- gx.mva(clr(sind.mat2open))
gx.rqpca.plot(sind.save)
gx.rqpca.plot(sind.save,
  main = "Howarth & Sinding Larsen Stream Sediments\nclr transform",
  pch = 4, cex.main = 0.9)
sind.save.rot2 <- gx.rotate(sind.save)
gx.rqpca.plot(sind.save.rot2,
  main = "Howarth & Sinding Larsen Stream Sediments\nclr transform",
  pch = 4, cex.main = 0.9)

## Clean-up
rm(sind.save)
rm(sind.save.rot2)
```

gx.rqpca.loadplot *Function to Graphically Display PCA Loadings*

Description

Function to graphically display PCA loadings computed by functions `gx.mva`, `gx.mva.closed`, `gx.robmva`, `gx.robmva.closed` or `gx.rotate`. The user may define the minimum absolute loading below which variables will not be graphically displayed, and modify the display title and text size as required.

Usage

```
gx.rqpca.loadplot(save, main = "", crit = 0.3, cex = 0.8,  
cex.axis = 0.7, cex.main = 0.8)
```

Arguments

save	a saved object from any of functions <code>gx.mva</code> , <code>gx.mva.closed</code> , <code>gx.robmva</code> or <code>gx.robmva.closed</code> .
main	an alternate plot title from that generated automatically from information in the saved object, see Details below.
crit	the lower limit of the absolute value of a loading for a variable to be displayed, by default <code>crit = 0.3</code> .
cex	the text scale expansion factor for the variable names in the display, by default <code>cex = 0.8</code> , a 20% font size reduction.
cex.axis	the text scale expansion factor for the axis labels of the display, by default <code>cex.axis = 0.7</code> , a 30% font size reduction.
cex.main	the text scale expansion factor for the display title, by default <code>cex.axis = 0.8</code> , a 20% font size reduction.

Details

If `main` is undefined the name of the matrix object supplied to the function is displayed, together with the value of `crit`. On the line below the name of the data matrix from which the PCA was derived is displayed. However, if an alternate plot title is preferred it may be defined, e.g., `main = "Plot Title Text"`. If no plot title is required set `main = ""`.

If the variable names are longer than three characters the display can easily become cluttered. In which case the user should redefine the variable names in the input matrix from which the PCA was derived using the `dimnames(matrix.name)[[2]]` construct, and run the generating function again. Alternately, the variable names in the saved object may be changed directly via a redefinition of `save$matnames[[2]]`.

Author(s)

Robert G. Garrett, based on a script by Peter Filzmoser

References

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. Statistical Data Analysis Explained: Applied Environmental Statistics with R. John Wiley & Sons, Ltd., 362 p.

See Also

[gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.rotate](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Estimate and display robust PCA loadings
sind.save <- gx.robmva.closed(sind.mat2open)
gx.rqpca.loadplot(sind.save)

## Clean-up
rm(sind.save)
```

gx.rqpca.plot

Function to Plot Principal Component Analysis Loadings and Scores

Description

Function to display the results of a Principal Components Analysis (PCA) from the saved object from [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#) or [gx.rotate](#) as biplots. Various options for displaying loadings and scores are available, see Details below.

Usage

```
gx.rqpca.plot(save, v1 = 1, v2 = 2, rplot = TRUE, qplot = TRUE,
rowids = NULL, ifrot = TRUE, main = "", cex.lab = 0.9,
cex.main = 0.9, rcex = 1, qcex = 0.8, rcol = 1, qcol = 1,
ifray = TRUE, if34 = TRUE, ...)
```

Arguments

save	a saved object from the execution of function gx.mva , gx.mva.closed , gx.robmva , gx.robmva.closed or gx.rotate .
v1	the component to be plotted on the x-axis of the biplot, default is the first component, v1 = 1.
v2	the component to be plotted on the y-axis of the biplot, default is the second component, v1 = 2.
rplot	the default is to plot the variables. If the variables are not required set rplot = FALSE. Note, if an ilr transform has been undertaken the loadings of the (p-1) synthetic variables will be displayed.

qplot	the default is to plot the observation (individual, case or sample) scores. If scores are not required set <code>qplot = FALSE</code> .
rowids	'switch' to determine if the input matrix row numbers are to be displayed instead of default plotting symbols. The default is for default plotting symbols, i.e. <code>rowids = NULL</code> , set <code>rowids = TRUE</code> if the row numbers are to be displayed, or set <code>rowids = FALSE</code> for the sample IDs to be displayed if they are present in the input matrix, if they are not, row numbers will be displayed.
ifrot	by default the post-Varimax rotation scores are displayed if a rotation has been made, see gx.rotate . If rotated scores are available in the saved object but the unrotated biplot is to be displayed set <code>ifrot = FALSE</code> .
main	an alternate plot title from that generated automatically from information in the saved object, see Details below.
cex.lab	the text scale expansion factor for the axis labels of the display, by default <code>cex.axis = 0.9</code> , a 10% font size reduction.
cex.main	the text scale expansion factor for the display title, by default <code>cex.axis = 0.9</code> , a 10% font size reduction.
rcex	the text scale expansion factor for the variable names in the display, by default <code>cex = 1</code> , 100%, no font size reduction.
qcex	the text scale expansion factor for the observation symbols, row numbers or sample IDs in the display, by default <code>cex = 0.8</code> , a 20% font size reduction.
rcol	the colour of the variable names in the display, by default <code>rcol = 1</code> , black, setting <code>rcol = 2</code> will result in red variable names (see display.lty for the default colour palette).
qcol	the colour of the observation symbols, row numbers or sample IDs in the display, by default <code>qcol = 1</code> , black, setting <code>qcol = 4</code> will result in blue variable names (see display.lty for the default colour palette).
ifray	by default 'rays' are plotted from the R loadings origin to the variable name in the same colour as the variable names, to omit the 'rays' set <code>ifray = FALSE</code> .
if34	this applies to a robust biplot only, by default the top (3) and right side (4) of the plot frame are annotated in black with the R loadings scale and with axis titles in the same colour as the variable names. To suppress these annotations set <code>if34 = FALSE</code> . When this extra annotation is provided the top (3) scale and annotation overwrites the plot title, therefore, set <code>main = " "</code> to suppress adding the plot title.
...	further arguments to be passed to methods concerning the plot.

Details

If `main` is undefined the name of the matrix object supplied to the function is displayed in the plot title. On the line below the name of the data matrix from which the PCA was derived is displayed. However, if an alternate plot title is preferred it may be defined, e.g., `main = "Plot Title Text"`. If no plot title is required set `main = " "`.

If the variable names are longer than three characters the display can easily become cluttered. In which case the user should redefine the variable names in the input matrix from which the PCA was derived using the `dimnames(matrix.name)[[2]]` construct, and run the generating function again.

Alternately, the variable names in the saved object may be changed directly via a redefinition of `save$matnames[[2]]`.

Information on the percentage of the variability explained by each component, and whether or not rotation has been undertaken, is recovered from the saved object and used to appropriately label the plot axes. Note that for non-robust models the percentage variability explained will be the same as the percentage variability explained by the corresponding eigenvalues.

The following describes the available plot option combinations, the first being the default:

`rplot = TRUE & qplot = TRUE & rowids = NULL`, crosses (pch default) and variable names

`rplot = TRUE & qplot = FALSE & rowids = NULL`, variable names only

`rplot = FALSE & qplot = TRUE & rowids = NULL`, crosses (pch default) only

`rplot = FALSE & qplot = TRUE & rowids = TRUE`, input matrix row numbers only

`rplot = FALSE & qplot = TRUE & rowids = FALSE`, input matrix row identifiers

`rplot = TRUE & qplot = TRUE & rowids = TRUE`, input matrix row numbers and variable names

Because functions `gx.mva`, `gx.mva.closed`, `gx.robmva` or `gx.robmva.closed` require a matrix as input the sample IDs that may be in a data frame may be lost. They may be re-inserted by copying `dimnames(...)[[1]]` from a data frame into the matrix. Alternately, to plot in the component score space with Sample IDs, the scores can be recovered from the saved object, e.g., `save$rqscore[, 1]` and `save$rqscore[, 2]`, and used as the x- and y-coordinates in function `xyplot.tags` with the sample IDs from the source data frame. Appropriate plot and axis titling can be displayed by setting the function arguments ‘by hand’.

Note

The scaling for the PCA ensures that the loadings and scores may be plotted in the same numerical space to create a biplot. However, when robust covariance matrix and mean estimation is employed this is not possible due to the extreme values some scores may attain for outliers. In this case the variables are plotted in the same physical space but with different scaling and addition scaling provided for the top and right side of the plot frame.

Author(s)

Robert G. Garrett

References

Reimann, C., Filzmoser, P., Garrett, R. and Dutter, R., 2008. *Statistical Data Analysis Explained: Applied Environmental Statistics with R*. John Wiley & Sons, Ltd., 362 p.

Venables, W.N. and Ripley, B.D., 2001. *Modern Applied Statistics with S-Plus*, 3rd Edition, Springer, 501 p.

See Also

`gx.mva`, `gx.mva.closed`, `gx.robmva`, `gx.robmva.closed`, `gx.rotate`, `xyplot.tags`

Examples

```
## Make test data available
data(sind)
data(sind.mat2open)
```

```

attach(sind)

## Save PCA results and display biplots
sind.save <- gx.mva.closed(sind.mat2open)
gx.rqpca.plot(sind.save)
gx.rqpca.plot(sind.save,
main = "Howarth & Sinding Larsen Stream Sediments\nclr transform",
pch = 4, rcol =2, qcol = 4)
gx.rqpca.plot(sind.save, rplot = TRUE, qplot = FALSE, rowids = NULL)
gx.rqpca.plot(sind.save, rplot = FALSE, qplot = TRUE, rowids = NULL)
gx.rqpca.plot(sind.save, rplot = FALSE, qplot = TRUE, rowids = TRUE)
gx.rqpca.plot(sind.save, rplot = TRUE, qplot = TRUE, rowids = FALSE,
rcol = 2, qcol = 4)

##
## Alternately
xyplot.tags(sind.save$rqscores[, 1],sind.save$rqscores[, 2], ID, cex = 0.9)

## Clean-up and detach test data
rm(sind.save)
detach(sind)

```

gx.rqpca.print

Function to Display PCA Loadings and Scores

Description

Function to display PCA matrices following computations by functions [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#) or [gx.rotate](#). The user may optionally display the loadings (default), the percentage contribution of the variables to the loadings, i.e. communalities (not default), and the scores on the PCs (default).

Usage

```
gx.rqpca.print(save, ifload = TRUE, ifcntrb = FALSE, ifscore = TRUE)
```

Arguments

save	a saved object from any of functions gx.mva , gx.mva.closed , gx.robmva , gx.robmva.closed or gx.rotate .
ifload	if ifload = TRUE the PC loadings are displayed. The default is to display the PC loadings.
ifcntrb	if ifcntrb = TRUE the percentage contribution of each variable (communality) to each PC is displayed. The default is not to display this table.
ifscore	if ifscore = TRUE the scores on the PCs are displayed. The default is to display the PC loadings.

Details

By default the PCA loadings and scores on the PCs are displayed on the current device. Optionally the percentage contribution, communality, of each variable to each PC may also be displayed. Additionally a table of cumulative percent contributions, communalities, is displayed to assist in deciding how many components to retain for rotation or further study. When the saved object from `gx.rotate` is the input object both the original and Varimax loadings and scores will be displayed by default.

Value

The last displayed table, `table.rows`, is returned and may be saved as an object if required.

Note

For large tables, particularly scores, it may be preferable to save them as '.csv' files using `gx.rqpca.save` for inspection and later use.

Author(s)

Robert G. Garrett

See Also

[gx.rqpca.save](#), [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.rotate](#).

Examples

```
## Make test data available
data(sind.mat2open)

## Estimate and display robust PCA loadings and scores
sind.save <- gx.robmva.closed(sind.mat2open)
gx.rqpca.print(sind.save, ifcntrb = TRUE)

## Clean-up
rm(sind.save)
```

gx.rqpca.save

Function to Save PCA Loadings and Scores as .csv files

Description

Function to save PCA matrices following computations by functions [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#) or [gx.rotate](#) as '.csv' files. By default appropriate files names are generated and the files saved in the Working Directory.

Usage

```
gx.rqpca.save(save, ifload = TRUE, ifcntrb = FALSE, ifscore = TRUE,
file = NULL)
```

Arguments

save	a saved object from any of functions gx.mva , gx.mva.closed , gx.robmva , gx.robmva.closed or gx.rotate .
ifload	if ifload = TRUE the PC loadings will be saved. The default is to save the PC loadings.
ifcntrb	if ifcntrb = TRUE the percentage contribution of each variable (communality) to each PC will be saved. The default is not to save this table.
ifscore	if ifscore = TRUE the scores on the PCs will be saved. The default is to save the PC scores.
file	the name of the folder where the files are to be saved, a '/' is appended before the synthesized file name, details of which are displayed on the current device. If no file is specified, the files are saved to the Working Directory.

Details

By default the PCA matrices are saved to the Working Directory with a synthesized file name consisting of the name of the saved object from which the matrices are sourced and an appropriate identifying suffix and '.csv'. If a folder name has been provided in file, a '/' is appended prior to the synthesized file name. As each matrix is saved its file name is displayed on the current device. When the saved object is from [gx.rotate](#) and is the input object both the original and Varimax loadings and PC scores will be saved by default.

Value

The last saved table, `table.rows`, is returned and may be saved as an object if required.

Author(s)

Robert G. Garrett

See Also

[gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#), [gx.rotate](#).

Examples

```
## Make test data available
data(sind.mat2open)

## Estimate and save robust PCA loadings and scores
sind.save <- gx.robmva.closed(sind.mat2open)
gx.rqpca.save(sind.save)

## NOTE: the example below is commented out
```

```
## Save PCA scores for future use
## gx.rqpca.save(sind.save, ifcntrb = TRUE, file = "D:\R_work\Project3")

## Clean-up
rm(sind.save)
```

gx.rqpca.screepLOT *Display a Scree Plot*

Description

Function to display a scree plot arising from a Principal Components Analysis (PCA) from the saved object from [gx.mva](#), [gx.mva.closed](#), [gx.robmva](#) or [gx.robmva.closed](#). In addition to the screeplot the cumulative variability explained is also displayed.

Usage

```
gx.rqpca.screepLOT(save, main = "", ...)
```

Arguments

save	a saved object from the execution of function gx.mva , gx.mva.closed , gx.robmva or gx.robmva.closed .
main	an alternate plot title to that in the saved object, see Details below.
...	further arguments to be passed to methods concerning the plot. For example, if some colour other than black is required for the plotting characters, specify <code>col = 2</code> to obtain red (see display.lty for the default colour palette). If it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

If `main` is undefined the name of the matrix object from which the PCA was derived is passed to the function via the saved object. Using the matrix name is the recommended procedure in the source functions as it helps to track the progression of the data analysis, acting as a record of the data source. However, at a presentation stage an alternate plot title may be preferred and can be defined in this function, e.g., `main = "Plot Title Text"`. If no plot title is required set `main = ""`.

Author(s)

Robert G. Garrett

See Also

[gx.mva](#), [gx.mva.closed](#), [gx.robmva](#), [gx.robmva.closed](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Save PCA results and display scree plot
sind.save <- gx.mva(ilr(sind.mat2open))
gx.rqpca.screeplot(sind.save)
gx.rqpca.screeplot(sind.save,
  main = "Howarth & Sinding Larsen Stream Sediments\nilr transform",
  pch = 4, col = 2, cex.main = 0.9)

## Clean-up
rm(sind.save)
```

gx.runs

The Wald-Wolfowitz, 'Runs', Test

Description

The 'runs' test is used to infer whether two states, e.g., > and < some threshold are mutually independent along a traverse. In applied geochemical terms, it tests for pattern coherence. If the pattern of runs is not coherent at the scale of the sampling it will be difficult to identify any spatially consistent dispersion processes.

Usage

```
gx.runs(n1, n2, u)
```

Arguments

n1	the number of < threshold sites along a traverse.
n2	the number of > threshold sites along a traverse.
u	the number of runs of > and < threshold sites along the traverse.

Note

Given a priori information on the location of a mineral occurrence, the [gx.hypergeom](#) function provides a far more insightful test. The 'runs' test is better suited for evaluating patterns due to lithological or environmental changes along a traverse when some 'threshold' can be selected that differentiates between two patterns

Author(s)

Robert G. Garrett

References

Stanley, C.R., 2003. Statistical evaluation of anomaly recognition performance. *Geochemistry: Exploration, Environment, Analysis*, 3(1):3-12.

See Also

[gx.hypergeom](#)

Examples

```
## From Stanley (2003) Table 2

gx.runs(27, 4, 7)
gx.runs(25, 6, 7)
gx.runs(28, 3, 5)
```

gx.scores

Function to Compute Scores on the Basis of Threshold Estimates

Description

Computes scores for a user selected group of variables based on the ratio of variable value to the variable threshold, i.e. the upper limit of background variation. The user must provide thresholds for the variables contributing to the scores. Optionally a set of relative weights may be provided that are applied to the scores. If above threshold values occur for a variable whose influence is indicative of a 'false' anomaly the relative weight for that variable may be set '-ve', which will result in a reduction of the computed score. This function is a useful alternative to weighted sums when the variable data contains so many below DL values that summary statistics cannot be estimated. An object is created containing all the provided parameters and the scores for later reference and use.

Usage

```
gx.scores(xx, tholds, rwts = NULL, setna = FALSE)
```

Arguments

xx	name of the n by p matrix containing the data.
tholds	a vector of the threshold estimates for the p variables.
rwts	an optional vector of the relative weights for the p variables, negative weights are permissible to indicate that high levels of the variable should have a negative impact on the scores.
setna	if it is required to set any '<=0' scores to NAs then set setna = TRUE. See note below.

Details

If the data for only some of the variables available in an attached matrix or data frame are to be processed use the cbind construct. Thus, `temp.mat <- cbind(vname1, vname3, vname6, vname8)`, or the cbind may be used directly, see Example below. All computed scores with values less than 1 are set to zero, optionally these may be replaced by NAs to facilitate their removal from subsequent plots, see Note below.

Value

The following are returned as an object to be saved for further use:

input	the name of the input data set
tholds	the vector of thresholds used for the computations
rwts	the vector of relative weights provided by the user
scores	the computed scores

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with with NAs are removed prior to computing the weighted sums.

In general, it is not advisable to replace zeros with NAs as the overall data or spatial structure of the data will not be discerned in subsequent plots as records with NAs will have been removed.

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Compute scores - 1
sind.scores1 <- gx.scores(cbind(Cu, Zn, Cd), tholds = c(100, 200, 2))
map.z(E, N, sind.scores1$scores)

## Compute scores - 2
sind.scores2 <- gx.scores(cbind(Cu, Zn, Cd, Fe), tholds = c(100, 200, 2, 2),
rwts = c(1, 1, 1, -1))
xyplot.z(Fe, Mn, sind.scores2$scores, col = 2)

## Clean up and detach test data
rm(sind.scores1)
rm(sind.scores2)
detach(sind)
```

`gx.sm`*Display Robust ilr Stabilities and log-ratio Medians for Compositional Data*

Description

The function computes and displays a matrix of Robust ilr Stabilities (Filzmoser et al., 2010) and medians of log-ratios in the upper and lower triangles, respectively.

Usage

```
gx.sm(xx, ifwarn = TRUE)
```

Arguments

`xx` a matrix, or sub-matrix, of parts from a compositional data set.

`ifwarn` by default `ifwarn = TRUE` which generates a reminder/warning that when carrying out analyses of compositional data all data must be in the same measurement units. The message can be suppressed by setting `ifwarn = FALSE`.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors (rows) containing NAs are removed prior to computation.

This function is for used with closed compositional data sets, i.e. geochemical analyses. For the 'classical' Aitchison (1984, 1986) approach see [gx.vm](#).

Author(s)

Robert G. Garrett

References

Aitchison, J., 1984. The statistical analysis of geochemical compositions. *Mathematical Geology*, 16(6):531-564.

Aitchison, J., 1986. *The Statistical Analysis of Compositional data*. Chapman and Hall, London, U.K., 416 p.

Filzmoser, P, Hron, K. and Reimann, C., 2010. The bivariate statistical analysis of environmental (compositional) data. *Science of the Total Environment*, 408(19):4230-4238.

See Also

[ltdl.fix.df](#), [remove.na](#), [ilr.stab](#), [gx.vm](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Compute Robust ilr Stabilities and log-ratio medians
gx.sm(sind.mat2open)
```

gx.sort

Function to Single Column Sort a Matrix or Data Frame

Description

Function to sort a matrix or data frame by the value in a column. On exit the function displays the sorted data. Any NAs in the sort column are sorted to beyond the greatest value. If the function is run as `temp <- gx.sort(x, ncol)` the sorted data are not displayed, but retained in `temp` for subsequent use or display.

Usage

```
gx.sort(x, col = 1, reverse = FALSE)
```

Arguments

<code>x</code>	the matrix or data frame to be sorted.
<code>col</code>	a column number, the value of which will be used to sort the matrix or data frame
<code>reverse</code>	the default is to sort in ascending order of the value in column <code>col</code> . If a descending order sort is required, set <code>reverse = TRUE</code> .

Author(s)

Robert G. Garrett

See Also

[gx.sort.df](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Sort data frame sind into ascending order on the value
## of column 4, Zn
gx.sort(sind, 4)
```

```
## Sort data frame sind into descending order on the value
## of column 4, Zn
gx.sort(sind, 4, reverse = TRUE)

## Sort only the geochemical data in data frame sind into
## descending order on the value of column 4, Zn. Note
## that sind[, -c(1:3)] moves the old column 4 to
## position 1
gx.sort(sind[, -c(1:3)], 1, reverse = TRUE)

## Detach test data
detach(sind)
```

gx.sort.df

Function to Multi-Column Sort a Data Frame

Description

Function to sort a data frame on any combination of numerical values or factors in any combination of ascending or descending orders. If the function is run as `temp <- gx.sort.df(formula, dfname)` the sorted data are not displayed, but retained in `temp` for subsequent use or display.

Usage

```
gx.sort.df(formula, dfname)
```

Arguments

formula	a 'formula' defining the variables to be used in the sort and whether the sort for each is to be in ascending or descending order. The sort order is from left to right in the formula. See Details and Examples below.
dfname	the name of the data frame to be sorted.

Details

The sort is controlled by a text string in the form of a 'formula', so `~var1+var2` will sort in ascending order of `var1`, and then within equal values for `var1` in ascending order of `var2`. A preceding `+` or `-` before a column name indicates a sort in ascending or descending order, respectively.

The function also works if `formula` and `dfname` are reversed in the function call.

Author(s)

Kevin Wright with some ideas from Andy Liaw
Shared on S-News and R-help in September 2004.

See Also

[gx.sort](#)

Examples

```
## Make test data available
data(kola.c)
attach(kola.c)
names(kola.c)

## Create a small test data set for ID (1), COUNTRY (4),
## As (17), Co (21), Cu (23) and Ni (28)
test<-kola.c[1:25, c(1,4,17,21,23,28)]

## Sort test data into ascending order on the value of Ni
gx.sort.df(~Ni, test)
temp <- gx.sort.df(test, ~Ni)
temp

## Sort test data by Country and descending order of As
gx.sort.df(test, ~COUNTRY-As)

## Sort test data by Country and descending order of both
## As and Ni
gx.sort.df(test, ~COUNTRY-As-Ni)

## Clean-up and detach test data
rm(test)
rm(temp)
detach(kola.c)
```

gx.spearman

Display Spearman Correlation Coefficients and their Significances

Description

The function computes Spearman rank correlation coefficients and places them in the upper triangle of a printed matrix displayed on the current device, the probabilities that the coefficients are not due to chance (Ho: Coefficient = 0) are printed in the lower triangle. The diagonal is filled with NAs to visually split the two triangles.

Usage

```
gx.spearman(xx, ifclr = FALSE, ifwarn = TRUE)
```

Arguments

xx	a matrix of numeric data.
ifclr	if ifclr = TRUE the data are Centred Log-Ratio transformed prior to the computation of the Pearson Coefficients. The default is no transformation.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out a centred log-ratio transformation all the data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors (rows) containing NAs are removed prior to computation.

This function is not recommended for use with closed compositional data sets, i.e. geochemical analyses, unless correlations are sought between a non-compositional variable and individual compositional variables. If it is used with compositional data, it is highly recommended that `ifclr` be set to TRUE to remove the effects of closure and display the ‘true’ inter-element variability. However, different groups of elements, subsets, of a data set will yield different inter-element correlations for the same pair of elements due to the nature of the `clr` transform. When carrying out a centred log-ratio transformation it is essential that the data are all in the same measurement units, and by default a reminder/warning is display if the data are centred log-ratio transformed, see `ifwarn` above.

For working with compositional data sets functions [gx.vm](#) and [gx.sm](#) are recommended. For visual displays see [gx.pairs4parts](#) and [gx.plot2parts](#).

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [remove.na](#), [clr](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Compute Spearman correlation coefficients
gx.spearman(sind.mat2open)

## Note, unlike gx.pearson there is no example with a log
## transformation. The log transformation is monotonic
## and does not change the ranks

## Compute Spearmann correlation coefficients following
## a centred log-ratio transformation
gx.spearman(sind.mat2open, ifclr = TRUE)
```

gx.stats

Function to Compute and Display Summary Statistics

Description

Function to compute summary statistics for a ‘one-page’ report and display in [inset](#). Function may be used stand-alone, and is used as an ‘engine’ for the `gx.summary.*` series of functions

Usage

```
gx.stats(xx, xlab = deparse(substitute(xx)), display = TRUE,
iftell = TRUE)
```

Arguments

xx	name of the variable to be processed.
xlab	by default the character string for xx is used for the table title. An alternate title can be displayed with xlab = "text string", see Examples.
display	if display = TRUE the summary statistics are displayed on the current device. If display = FALSE output is suppressed.
iftell	by default the NA count is displayed by na.remove prior to the table of results from this function. When the function is used as a 'stats' engine the NA count display may be suppressed by the calling function when the NA count is to be displayed by that calling function.

Details

The summary statistics comprise the data minimum, maximum and percentile values, robust estimates of standard deviation, the Median Absolute Deviation (MAD) and the Inter Quartile Standard Deviation (IQSD), and the mean, variance, standard deviation (SD), coefficient of variation (CV%), and the 95% confidence bounds on the median. When the minimum data value is > 0 summary statistics are computed after a log10 data transformation and exported back to the calling function.

Value

stats	the computed summary statistics to be used in function inset , and by <code>gx.summary.*</code> functions. The list returned, stats, is a 32-element vector, see below:
[1:10]	the minimum value, and the 1st, 2nd, 5th, 10th, 20th, 25th (Q1), 30th, 40th and 50th (Q2) percentiles.
[11:19]	the 60th, 70th, 75th (Q3), 80th 90th, 95th, 98th and 99th percentiles and the maximum value.
[20]	the sample size, N.
[21]	the Median Absolute Deviation (MAD).
[22]	the Inter-Quartile Standard Deviation (IQSD).
[23]	the data (sample) Mean.
[24]	the data (sample) Variance.
[25]	the data (sample) Standard Deviation (SD).
[26]	the Coefficient of Variation as a percentage (CV%).
[27]	the Lower 95% Confidence Limit on the Median.
[28]	the Upper 95% Confidence Limit on the Median.
[29]	the log10 transformed data (sample) Mean.
[30]	the log10 transformed data (sample) Variance.

```
[31]          the log10 transformed data (sample) SD.
[32]          the log10 transformed data (sample) CV%.
If the minimum data value is <= 0, then stats[29:32] <- NA.
```

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to computation. Depending on the value of `iftell`, the NA count will be displayed, `iftell = TRUE`, or suppressed, `iftell = FALSE`.

The confidence bounds on the median are estimated via the binomial theorem, not by normal approximation.

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display
gx.stats(Cu)

## Provides a more appropriate labelled display
gx.stats(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil")

## Detach test data
detach(kola.o)
```

gx.subset

Extracts a Subset of Rows from a Data Frame

Description

The function extracts a subset of rows, and columns if required, from a data frame and returns the subset as a new data frame based on the criterion provided by the user. Unused factor names are dropped.

Usage

```
gx.subset(dfname, subset = TRUE)
```


Arguments

dfname name of the data frame from which rows are to be extracted.
subset the criterion for selecting the subset (rows).

Details

The subset criterion can be 'complex' and be a combination of conditions, see Examples below.

Value

data a data frame only containing the rows of the input data frame where the criterion is met.

Note

This function is based on a script shared by Bill Venables on S-News, October 10, 1997. As such it may pre-date the time that [subset](#) was added to the S-Plus library. It is simple to use and has been retained.

Author(s)

William N. Venables

See Also

[subset](#)

Examples

```
## Make test data available
data(kola.c)

## Make a subset of the data for Finland
finland.c <- gx.subset(kola.c, COUNTRY == "FIN")

## Make a subset of the data for rock type, LITHO 82 occurring
## in Russia. Note that both COUNTRY and LITHO are factor variables
russia.82 <- gx.subset(kola.c, COUNTRY == "RUS" & LITHO == 82)

## Make a subset of the data for Cu exceeding 50(ppm) in Norway
norway.cugt50 <- gx.subset(kola.c, COUNTRY == "NOR" & Cu >50)

## Make single element subsets, e.g. for use with function gx.cnpplts
## First locate the column in the data frame where the element of
## interest is stored using dimnames(kola.c)[[2]], we find that Be is
## the 19th column in the data frame
dimnames(kola.c)[[2]]
Norway <- gx.subset(kola.c, COUNTRY=="NOR")[,19]
Russia <- gx.subset(kola.c, COUNTRY=="RUS")[,19]
Finland <- gx.subset(kola.c, COUNTRY=="FIN")[,19]
```

```
## Clean-up
rm(finland.c)
rm(russia.82)
rm(norway.cugt50)
rm(Norway)
rm(Russia)
rm(Finland)
```

gx.summary

Compiles a Table of Summary Statistics

Description

This function is a ‘sub-engine’ between the main summary statistics engine, ‘gx.stats’, and the `gx.summary.*` display functions, it does generate any output to the current device. Its ‘sub-engine’ function is to select the required results from the `gx.stats` computations, and additionally compute 95% confidence bounds on means.

Usage

```
gx.summary(xx, log = log, iftell = iftell)
```

Arguments

<code>xx</code>	name of the variable to be processed.
<code>log</code>	set <code>log = TRUE</code> if it is required to compute statistics following a log10 transformation of the data.
<code>iftell</code>	passes the value of <code>iftell</code> for controlling the display of the NA count to function <code>remove.na</code> from the calling function.

Value

<code>table</code>	a 15-element vector containing summary statistics, see below:
[1]	the sample size, N.
[2]	the number of NAs removed from the data passed for processing.
[3:7]	the minimum value, Q1, Median, Q3 and maximum value.
[8]	the Median Absolute Deviation (MAD).
[9]	the Inter-Quartile Standard Deviation (IQSD).
The contents of elements [10:15] depend on the ‘value’ of <code>log</code>	
[10]	the data (sample) Mean.
[11]	the data (sample) Standard Deviation (SD).
[12]	the Coefficient of Variation as a percentage (CV%).
[13]	the Standard Error (S.E.) of the Mean.
[14]	the Lower 95% Confidence Limit on the Mean.

[15] the Upper 95% Confidence Limit on the Mean.

If `log = TRUE`, the results for the mean, [13], and confidence limits, [14:15], are backtransformed to the natural scale.

The returned table is rounded to 4 significant figures.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing `gx.summary.*` functions that call this function, see `ltdl.fix.df`.

Any NAs in the data vector will be removed prior to computation in function `gx.stats`. Depending on the value of `iftell`, the NA count will be displayed, `iftell = TRUE`, or suppressed, `iftell = FALSE`.

There are no examples for this function. To display the summary statistics use one of the summary statistics display functions: `gx.stats`, `gx.summary1`; `gx.summary2`; `gx.summary.mat`; `gx.summary.groups`; `framework.stats`; or `framework.summary`.

Author(s)

Robert G. Garrett

See Also

`gx.stats`, `ltdl.fix.df`, `remove.na`,

`gx.summary.groups` *Displays Summary Statistics for a Variable Grouped by a Factor*

Description

Displays the same concise one-line summary statistics report as `gx.summary1` but with the data grouped by the value of a factor variable. The table consists of a heading line and a line of summary statistics for each 'group', value of the factor variable. Optionally the data may be logarithmically (base 10) transformed.

Usage

```
gx.summary.groups(group, x, xname = deparse(substitute(x)),
log = FALSE)
```

Arguments

group	the name of the factor variable the data are to be grouped by.
x	name of the variable to be processed.
xname	by default the character string for x is used for the title. An alternate title can be displayed with <code>xname = "text string"</code> , see Examples.
log	if the summary statistics are required following a log10 transformation, set <code>log = TRUE</code> .

Details

Setting `log = TRUE` results in a log transformation for the parametric statistical estimates. The maximum, minimum, quartiles and robust estimates of spread are estimated and reported in natural measurement units. Of the parametric statistics, the mean (the geometric mean) and 95% confidence are reported backtransformed into natural measurement units.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation in function `gx.stats`. Display of the number of NAs found by function `remove.na` is suppressed in `remove.na` as the information is included in the display from this function.

Alternately, function [framework.summary](#) generates grouped summary statistics that are exported in a file format that can be directly imported into a spreadsheet, e.g., MS Excel, for inspection, or into other software, e.g., a Geographical Information System (GIS) where the spatial information concerning the 'framework' units is available, e.g., ecoclassification units.

For more extensive summary statistics displaying one variable at a time, see [gx.summary2](#) using a construct like `gx.summary2(var[factor == "value"])` or use function [inset](#) with a similar construct.

For summary graphical presentations see functions [bwplots](#) or [tbplots](#).

Author(s)

Robert G. Garrett

See Also

[gx.summary1](#), [gx.summary](#), [gx.stats](#), [ltdl.fix.df](#), [remove.na](#), [gx.summary2](#)

Examples

```
data(kola.c)
attach(kola.c)

## Generates an initial display
gx.summary.groups(COUNTRY, Cu)
```

```
## Provide a more informative display
gx.summary.groups(COUNTRY, Cu, xname = "Cu (mg/kg) in <2 mm 0-horizon soil")

## As above but with a log10 transformation to display
## the geometric mean, etc.
gx.summary.groups(COUNTRY, Cu, xname = "Cu (mg/kg) in <2 mm 0-horizon soil",
log = TRUE)

## Detach test data
detach(kola.c)
```

gx.summary.mat

Displays Summary Statistics for a Matrix or Data Frame

Description

Displays the same concise one-line summary statistics report as [gx.summary1](#) for two or more columns of a matrix or data frame. The table consists of a heading line and a line of summary statistics for each ‘variable’, column of the matrix or data frame. Optionally the data may be logarithmically (base 10) transformed.

Usage

```
gx.summary.mat(xmat, vars, banner = deparse(substitute(xmat)),
log = FALSE)
```

Arguments

xmat	name of the matrix or data frame.
vars	the indices, or names (see Example), of the columns of the matrix or data frame for the variables whose summary statistics are to be displayed.
banner	by default the character string for xmat, the input matrix, is used for the title. An alternate title can be displayed with banner = "text string", see Examples.
log	if the summary statistics are required following a log10 transformation, set log = TRUE.

Details

Setting log = TRUE results in a log transformation for the parametric statistical estimates. The maximum, minimum, quartiles and robust estimates of spread are estimated and reported in natural measurement units. Of the parametric statistics, the mean (the geometric mean) and 95% confidence are reported backtransformed into natural measurement units.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation in function `gx.stats`. Display of the number of NAs found by function `remove.na` is suppressed in `remove.na` as the information is included in the display from this function.

For a more extensive summary statistics display a variable at a time, see [gx.summary2](#), and for a summary with graphical displays see [inset](#).

For summary graphical presentations see functions [bwplots.by.var](#) or [tbplots.by.var](#).

Author(s)

Robert G. Garrett

See Also

[gx.summary1](#), [gx.summary](#), [gx.stats](#), [ltdl.fix.df](#), [remove.na](#), [gx.summary2](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display for As [6], Co [13], Cu [15],
## Ni [24] and Zn [38]
gx.summary.mat(kola.o, c(6, 13, 15, 24, 38))

## Alternately
gx.summary.mat(kola.o, c("As", "Co", "Cu", "Ni", "Zn"))

## Provide a more informative display for Be [9], La [19], P [25],
## Th [33], U [35] and Y[37]
gx.summary.mat(kola.o, c(9, 19, 25, 33, 35, 37),
  banner = "Kola Project, <2 mm 0-horizon soils")

## As above but with a log10 transformation to display
## the geometric mean, etc.
gx.summary.mat(kola.o, c("Be", "La", "P", "Th", "U", "Y"),
  log = TRUE, banner = "Kola Project, <2 mm 0-horizon soils")

## Detach test data
detach(kola.o)
```

`gx.summary1`*Display a one-line Summary Statistics Report*

Description

Displays a concise one-line summary statistics report, below a heading line, consisting of sample size, number of NAs in the input vector; minimum, maximum and quartiles; robust estimates of the standard deviation (MAD and interquartile based measure); mean, standard deviation and coefficient of variation (%); and the standard error, and lower and upper 95% confidence limits on the mean. See Details for the results of setting `log = TRUE`. Optionally the data may be logarithmically (base 10) transformed.

Usage

```
gx.summary1(xx, xname = deparse(substitute(xx)), log = FALSE)
```

Arguments

<code>xx</code>	name of the variable to be processed.
<code>xname</code>	by default the character string for <code>xx</code> is used for the title. An alternate title can be displayed with <code>xname = "text string"</code> , see Examples.
<code>log</code>	if the summary statistics are required following a <code>log10</code> transformation, set <code>log = TRUE</code> .

Details

Setting `log = TRUE` results in a log transformation for the parametric statistical estimates. The maximum, minimum, quartiles and robust estimates of spread are estimated and reported in natural measurement units. Of the parametric statistics, the mean (the geometric mean) and 95% confidence are reported backtransformed into natural measurement units. If all the results are required following a `log10`, or some other transformation, this can be achieved by executing the transformation in the call, e.g., `gx.summary1(log10(Cu))` or `gx.summary1(sqrt(Cu))`, and setting `log = FALSE`.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation in function `gx.stats`.

For a more extensive summary statistics display, see [gx.summary2](#). For summary graphical displays see [shape](#) or [inset](#).

Author(s)

Robert G. Garrett

See Also

[gx.summary](#), [gx.stats](#), [ltdl.fix.df](#), [remove.na](#), [gx.summary2](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display
gx.summary1(Cu)

## Provide a more informative display
gx.summary1(Cu, xname = "Cu (mg/kg) in <2 mm Kola 0-horizon soil")

## As above but with a log10 transformation to display
## the geometric mean, etc.
gx.summary1(Cu, xname = "Cu (mg/kg) in <2 mm Kola 0-horizon soil", log = TRUE)

## Detach test data
detach(kola.o)
```

gx.summary2

Display a ten-line Summary Statistics Report

Description

Displays a more extensive report than `gx.summary1`. The report includes sample size, number of NAs in the input vector; arithmetic mean and 95% confidence limits, standard deviation and CV%; geometric mean and 95% confidence limits, with standard deviation and CV% in log10 units; median and 95% confidence limits robust estimates of spread (MAD and interquartile based measure); and minimum, maximum, quartiles, and 2nd, 5th, 10th, 90th, 95th and 98th percentiles.

Usage

```
gx.summary2(xx, xname = deparse(substitute(xx)))
```

Arguments

xx	name of the variable to be processed.
xname	by default the character string for xx is used for the report title. An alternate title can be displayed with <code>xname = "text string"</code> , see Examples.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector will be removed prior to computation in function `gx.stats`.

For a less extensive summary statistics display, see [gx.summary1](#). For summary graphical displays see [shape](#) or [inset](#).

Author(s)

Robert G. Garrett

See Also

[gx.summary](#), [gx.stats](#), [ltdl.fix.df](#), [remove.na](#), [gx.summary1](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display
gx.summary2(Cu)

## Provide a more informative display
gx.summary2(Cu, xname = "Cu (mg/kg) in <2 mm Kola O-horizon soil")

## Detach test data
detach(kola.o)
```

gx.t.test

t-test from Means, Standard Deviations and Sample Sizes

Description

Function to test hypothesis of equality of two means given the mean, standard deviation and size (N) of each population.

Usage

```
gx.t.test(xbar1, s1, n1, xbar2, s2, n2)
```

Arguments

xbar1	mean of population 1.
s1	standard deviation of population 1.
n1	size of population 1.
xbar2	mean of population 2.
s2	standard deviation of population 2.
n2	size of population 2.

Details

Function written to support QA/QC studies where data for sets of Control Reference Material analyses are to be compared, e.g., between two projects, or when comparing published statistics with those from a project.

The hypothesis, $H_0: \text{mean}_1 = \text{mean}_2$, is tested at the 95% confidence level and the user informed whether the hypothesis is accepted or rejected. Additionally the difference, t-value, degrees of freedom, and p-value are displayed.

Author(s)

Robert G. Garrett

Examples

```
gx.t.test(22.3, 4.6, 25, 24.2, 3.8, 22)
gx.t.test(22.3, 4.6, 25, 26.2, 3.8, 22)
```

gx.triples.aov *Carries out a 3-Level Staggered ANOVA and Estimates Variance Components*

Description

Function to undertake an ANOVA for the unbalanced triplicates from a GSC NGR or Tri-National survey. The data must be in the following order for each triplicate: Analytical Duplicate, Field Duplicate for the Analytical Duplicate Split, other Field Duplicate. The results replicate those generated by the UANOVA (Garrett and Goss, 1980) computer program. Optionally the data may be logarithmically (base 10) transformed.

Usage

```
gx.triples.aov(x, xname = deparse(substitute(x)), log = FALSE,
table = FALSE)
```

Arguments

x	a file of triplicate determinations, the order is critical, see Details below.
xname	by default the character string for the data file name, x, is used for the table title. An alternate title can be displayed with xname = "text string", see Examples.
log	if a logarithmic transformation of the data is required to meet homogeneity of variance considerations (i.e. severe heteroscedasticity) set log = TRUE. This is also advisable if the range of the observations exceeds 1.5 orders of magnitude.
table	set table = TRUE if the input data file is to be displayed. the default is no display.

Details

As noted above, the order of the data is critical and must be as follows for each triplicate: Analytical Duplicate, Field Duplicate for the Analytical Duplicate Split, other Field Duplicate. The 'other Field Duplicate' is equivalent to a regular regional-coverage sample, but is at a 'Field Duplicate' site. Thus below, x[i,1] will contain the Analytical Duplicates, x[i,2] the Field Duplicates from which the Analytical Duplicates were split, and x[i,3] the other analytically unduplicated Field Duplicates. See Details in [triples.test1](#) for additional information.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data must also be removed prior to running the `triples.aov` function. This requires care as the data must be in complete triplicate sets.

Author(s)

Robert G. Garrett

References

- Bainbridge, T.R., 1963. Staggered, nested designs for estimating variance components. American Society for Quality Control, Convention Transactions, pp. 93-103.
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- Garrett, R.G. & Goss, T.I., 1980. UANOVA: A Fortran IV program for unbalanced nested analysis of variance. *Mathematical Geology*, 6(1):35-60.
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- Snee, R.D., 1974. Computation and use of expected mean squares in Analysis of Variance. *Journal of Quality Technology*, 6(3):128-137.

See Also

[ltdl.fix.df](#), [remove.na](#), [triples.test1](#), [gx.triples.fgx](#), [triples.test2](#)

Examples

```
## Make test data available
data(triples.test1)
attach(triples.test1)

## Carry out unbalanced ANOVA
gx.triples.aov(Ba_ppm, xname =
"Ba (mg/kg - Aqua Regia digestion) in <2 mm unmilled C-horizon soil")

## Detach test data
detach(triples.test1)
```

gx.triples.fgx

ANOVA to Estimate if 'Triples' are a Valid Subset

Description

Function to execute a simple ANOVA to determine if the Field Duplicates are a valid subset of the regional coverage samples, and if the Field Duplicates pairs have 'equivalent' variability. Optionally the data may be logarithmically (base 10) transformed.

Usage

```
gx.triples.fgx(x, RepStat, xname = deparse(substitute(x)),
log = FALSE)
```

Arguments

x	a file of regional coverage and field duplicate data.
RepStat	the Replicate Status code.
xname	by default the character string for the data file name, x, is used for the table title. An alternate title can be displayed with xname = "text string", see Examples.
log	if a logarithmic transformation of the data is required to meet homogeneity of variance considerations (i.e. severe heteroscedasticity) set log = TRUE. This is also advisable if the range of the observations exceeds 1.5 orders of magnitude.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data must also be removed prior to running the `triples.fgx` function. This requires care as the data must be in complete duplicate sets.

Author(s)

Robert G. Garrett

References

Garrett, R.G., 2013. Assessment of local spatial and analytical variability in regional geochemical surveys with a simple sampling scheme. *Geochemistry: Exploration, Environment, Analysis*, 13(4):349-354, doi 10.1144/geochem2011-085.

See Also

[ltdl.fix.df](#), [remove.na](#), [triples.test1](#), [gx.triples.fgx](#), [triples.test2](#)

Examples

```
## Make test data available
data(triples.test2)
attach(triples.test2)

## Carry out ANOVAs for equivalence of variances
gx.triples.fgx(Ba_ppm, RS, xname =
"Ba (mg/kg - Aqua Regia digestion) in <2 mm unmilled C-horizon soil")

## Detach test data
detach(triples.test2)
```

gx.v_m

Display an Aitchison Variation Matrix for Compositional Data

Description

The function computes and displays an Aitchison Variation Matrix, with the variances and means of the log-ratios in the upper and lower triangles, respectively.

Usage

```
gx.vm(xx, ifwarn = TRUE)
```

Arguments

xx	a matrix, or sub-matrix, of parts from a compositional data set.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out analyses of compositional data all data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data vector, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors (rows) containing NAs are removed prior to computation.

This function is for used with closed compositional data sets, i.e. geochemical analyses. For an alternate approach see [gx.sm](#), where a robust ilr stability measure (Filzmoser et al., 2010) is used rather than the log-ratio variance, and the median of log-ratios is used rather than the mean.

Author(s)

Robert G. Garrett

References

Aitchison, J., 1984. The statistical analysis of geochemical compositions. *Mathematical Geology*, 16(6):531-564.

Aitchison, J., 1986. *The Statistical Analysis of Compositional data*. Chapman and Hall, London, U.K., 416 p.

Filzmoser, P, Hron, K. and Reimann, C., 2010. The bivariate statistical analysis of environmental (compositional) data. *Science of the Total Environment*, 408(19):4230-4238.

See Also

[ltdl.fix.df](#), [remove.na](#), [gx.sm](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Compute Aitchison Variation Matrix
gx.vm(sind.mat2open)
```

ilr

Isometric Log-Ratio (ilr) transformation

Description

Undertakes an isometric log-ratio transformation to remove the effects of closure in a data matrix.

Usage

```
ilr(xx, ifclose = FALSE, ifwarn = TRUE)
```

Arguments

xx	a n by p matrix to be isometrically log-ratio transformed. It is essential that a single unit of measurement is used. Thus it may be required to convert, for example, determinations in percent to ppm (mg/kg) so that all measurements are in ppm prior to executing this function. Natural logarithms are used.
ifclose	if it is required to close a data set prior to transformation set ifclose = TRUE.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out a centred log-ratio transformation all the data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.

Details

Most analytical chemical data for major, minor and trace elements are of a closed form, i.e. for a physical individual sample they sum to a constant, whether it be percent, ppm (mg/kg), or some other units. It does not matter that only some components contributing to the constant sum are present in the matrix, the data are closed. As a result, as some elements increase in concentration others must decrease, this leads to correlation measures and graphical presentations that do not reflect the true underlying relationships. However, isometrically transformed data are not suitable for univariate EDA inspection as the new synthetic variables bear a complex relationship to the original measurements. Other procedures for removing closure effects are additive log-ratios (alr) and centred log-ratios (clr).

Value

x	a n by (p-1) matrix of isometric log-ratio values. The names of the new (p-1) synthetic variables, iso1 through to isop, where the p in isop equals p-1, are entered as column names in the matrix.
---	---

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows containing NAs in the data matrix are removed prior to undertaking the transformation.

The `ilr` transform is recommended for the calculation of Mahalanobis distances, a procedure which requires matrix inversion. When a Principal Component or Factor Analysis is required use of the `ilr` transform may be preferable, see also the notes in [clr](#). In that instance back transformation from the isometrically transformed variables to the original variables is required. Interested R users should refer to the papers by Filzmoser et al. (see below).

Author(s)

Peter Filzmoser and Karel Hron, with additions by Robert G. Garrett

References

Aitchison, J. and Egozcue, J.J., 2005. Compositional data analysis; where are we and where should we be heading. *Mathematical Geology*, 37(7):829-850.

Buccianti, A., Mateu-Figueras, G, and Pawlowsky-Glahn, V. (eds.), 2006. Compositional data analysis in the geosciences: from theory to practice. The Geological Society Publishing House, Bath, U.K. Special Publication 264, 224 p.

Filzmoser, P. and Hron, K., 2008. Outlier detection for compositional data using robust methods. *Mathematical Geosciences*, 40(3):234-248.

Filzmoser, P., Hron, K. and Reimann, C., 2009. Principal component analysis for compositional data with outliers. *Environmetrics*, 20(6):621-633.

Filzmoser, P., Hron, K., Reimann, C. and Garrett, R.G., 2009. Robust factor analysis for compositional data. *Computers & Geosciences*, 35(9):1854-1861.

See Also

[alr](#), [clr](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data sind available
data(sind.mat2open)

## Undertake ilr transform
temp <- ilr(sind.mat2open)
temp

## Clean-up
rm(temp)
```

ilr.stab

Compute the Robust ilr Stability for Two Parts of a Composition

Description

Function computes the Robust ilr Stability for two parts of a composition following the procedure in Filzmoser et al. (2010), see details below.

Usage

```
ilr.stab(xx1, xx2, ifwarn = T)
```

Arguments

xx1	a column vector from a matrix or data frame of compositional data, xx1[1], ..., xx1[n].
xx2	another column vector from the matrix or data frame of compositional data, xx2[1], ..., xx2[n]. xx1 and xx2 must be of identical length, n.
ifwarn	by default ifwarn = TRUE which generates a reminder/warning that when carrying out analyses of compositional data all data must be in the same measurement units. The message can be suppressed by setting ifwarn = FALSE.

Details

The ilr transform of two parts of a composition is $\text{ilr.xy} = 1/(\sqrt{2}) * \log(x1/x2)$. The Robust ilr Stability (Filzmoser et al., 2010) is computed from the MAD of the `ilr.xy` values. This is normalized into the $(0, 1)$ interval as $\exp(-\text{ilr.MAD} * \text{ilr.MAD})$, following the procedure of Buccianti and Pawlowsky-Glahn (2005).

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors (rows) containing NAs are removed prior to computation.

Author(s)

Robert G. Garrett

References

Buccianti, a. and Pawlowsky-Glahn, V., 2005. New perspectives on water chemistry and compositional data analysis. *Mathematical Geology*, 37(7):703-727.

Filzmoser, P, Hron, K. and Reimann, C., 2010. The bivariate statistical analysis of environmental (compositional) data. *Science of the Total Environment*, 408(19):4230-4238.

See Also

[ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Compute Robust ilr Stability
ilr.stab(Cu, Zn)

## Detach test data
detach(sind)
```

Description

Plots a three panel graphical distributional summary for a data set, comprising a histogram and a cumulative normal percentage probability (CPP) plot, together with a table of selected percentiles of the data and summary statistics between them. Optionally the EDA graphics may be plotted with base 10 logarithmic scaling.

Usage

```
inset(xx, xlab = deparse(substitute(xx)), log = FALSE, xlim = NULL,
      nclass = NULL, colr = NULL, ifnright = TRUE, table.cex = 0.7, ...)
```

Arguments

<code>xx</code>	name of the variable to be plotted.
<code>xlab</code>	by default the character string for <code>xx</code> is used for the x-axis plot titles. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
<code>log</code>	to display the data with logarithmic (x-axis) scaling, set <code>log = TRUE</code> .
<code>xlim</code>	default limits of the x-axis are determined in the function. However when used stand-alone the limits may be user-defined by setting <code>xlim</code> , see Note below.
<code>nclass</code>	the default procedure for preparing the histogram depends on sample size. Where $N \leq 500$ the Scott (1979) rule is used, and when $N > 500$ the Freedman-Diaconis (1981) rule; both these rules are resistant to the presence of outliers, and usually provide informative histograms. Alternately, the user may define the histogram binning by setting <code>nclass</code> , i.e. <code>nclass = "scott"</code> , <code>nclass = "fd"</code> or <code>nclass = "sturges"</code> ; the latter being designed for normal distributions (Scott, 1992). See Venables and Ripley (2001) for details.
<code>colr</code>	by default the histogram is infilled in grey, <code>colr = 8</code> . If no infill is required, set <code>colr = 0</code> . See function <code>display.lty</code> for the range of available colours.
<code>ifnright</code>	controls where the sample size is plotted in the histogram display, by default this in the upper right corner of the plot. If the data distribution is such that the upper left corner would be preferable, set <code>ifnright = FALSE</code> . If neither option generates an acceptable plot, setting <code>ifnright = NULL</code> suppresses the display of the data set size.
<code>table.cex</code>	controls the size of the text in the central panel of summary statistics table, the default is <code>table.cex = 0.7</code> , which has been found to be optimal. If the entire table does not display, just parts of the columns, see Note below.
<code>...</code>	further arguments to be passed to methods. For example, by default individual data points in the CPP plot are marked by a plus sign, <code>pch = 3</code> , if a cross or open circle is desired, then set <code>pch = 4</code> or <code>pch = 1</code> , respectively. See <code>display.marks</code> for all available symbols. Adding <code>ifqs = TRUE</code> results in horizontal and vertical dotted lines being plotted at the three central quartiles and their values, respectively, in the CPP plot.

Details

A histogram is displayed on the left, and a cumulative normal percentage probability plot on the right. Between the two is a table of simple summary statistics, computed by `gx.stats`, including minimum, maximum and percentile values, robust estimates of standard deviation, and the mean, standard deviation and coefficient of variation. The plots may be displayed with logarithmic axes, however, the summary statistics are not computed with a logarithmic transform.

Note

Sometimes the table between the two plots may be left and/or right truncated, or incompletely displayed. Reducing the size of the graphics window will lead to a complete display. If this needs to be done the function needs to be repeated to ensure a correctly dimensioned display is saved. Once as a complete saved graphics file the image may be resized in the receiving document.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plot.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`. If the defined limits lie within the observed data range a truncated plot will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations.

The purpose of this function is to prepare publication quality graphics (`.emf` or `.ps`) files that can be included in reports or used as inset statistical summaries for maps. If a series of these are to be prepared the function [inset.exporter](#) can be used to advantage as it saves a graphics file as part of its procedure.

For summary statistics tables to complement the graphical display see, [gx.stats](#), [gx.summary1](#), [gx.summary2](#) and [gx.ngr.summary](#).

In some R installations the generation of multi-panel displays and the use of function `eqsplot` from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a 'first' function prepared by the user that loads the 'rgr' package, etc.

Author(s)

Robert G. Garrett

References

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p. See pp. 119 for a description of histogram bin selection computations.

See Also

[gx.hist](#), [cnpplt](#), [gx.stats](#), [inset.exporter](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display
inset(Cu)

## Provides a more appropriate display for publication
inset(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE)
```

```
## NOTE: The example statistics table may not display correctly

## Detach test data
detach(kola.o)
```

inset.exporter *Saves an EDA Graphical and Statistical Summary*

Description

Saves the output from function `inset` as a graphics file in the R Working Directory for use in report or map preparation. Optionally the EDA graphics may be plotted with base 10 logarithmic scaling.

Usage

```
inset.exporter(x, xlab = deparse(substitute(x)), log = FALSE,
xlim = NULL, nclass = NULL, ifnright = TRUE, file = NULL,
table.cex = 0.7, gtype = "emf", ...)
```

Arguments

<code>x</code>	name of the variable to be plotted.
<code>xlab</code>	a label for the x-axis. It is often desirable to replace the default x-axis label of the input variable name text string with a more informative label, e.g., <code>xlab = "Cu (mg/kg) in <2 mm 0-horizon soil"</code> .
<code>log</code>	to display the data with logarithmic (x-axis) scaling, set <code>log = TRUE</code> .
<code>xlim</code>	default limits of the x-axis are determined in the function. However when used stand-alone the limits may be user-defined by setting <code>xlim</code> , see Note below.
<code>nclass</code>	the default procedure for preparing the histogram is to use the Scott (1979) rule. This usually provides an informative histogram, other optional rules are <code>nclass = "sturges"</code> or <code>nclass = "fd"</code> ; the later standing for Freedman-Diaconis (1981), a rule that is resistant to the presence of outliers in the data. See Venables and Ripley (2001) for details, or <code>inset</code> or <code>gx.hist</code> .
<code>ifnright</code>	controls where the sample size is plotted in the histogram display, by default this in the upper right corner of the plot. If the data distribution is such that the upper left corner would be preferable, set <code>ifnright = FALSE</code> . If neither option generates an acceptable plot, setting <code>ifnright = NULL</code> suppresses the display of the data set size.
<code>file</code>	the name of the folder where the files are to be saved, a <code>'/'</code> is appended before the synthesized file name, details of which are displayed on the current device. If no file is specified, the files are saved to the Working Directory, see Details below.
<code>table.cex</code>	controls the size of the text in the central panel of summary statistics table, the default is <code>table.cex = 0.7</code> , which has been found to be optimal. If the entire table does not display, just parts of the columns, see Note below.

gtype	the format of the graphics file to be saved. By default gtype = "emf" for a Windows extended metafile. Other alternatives are gtype = "jpg" for a jpeg file, gtype = "png" for a portable network graphics file, gtype = "ps" for a postscript file, or gtype = "pdf" for a pdf file.
...	further arguments to be passed to methods. For example, by default individual data points in the CPP plot are marked by a plus sign, pch = 3, if a cross or open circle is desired, then set pch = 4 or pch = 1, respectively. See display.marks for all available symbols. Adding ifqs = TRUE results in horizontal and vertical dotted lines being plotted at the three central quartiles and their values, respectively, in the CPP plot.

Details

See [inset](#) for details concerning the inset parameters.

The graphics file will be saved, by default, in the Working Directory. Alternately a folder name may be provided in file. The file name is synthesized from the data frame name, `'_inset_'`, the variable name and `' .gtype'`. This is prefixed with a `'\'` and concatenated with the folder location.

Note

The synthesis of the file name uses the data frame name which it is assumed is located in `search()` position `[[2]]`.

Sometimes the table between the two plots may be left and/or right truncated, or incompletely displayed. Reducing the size of the graphics window will lead to a complete display. If this needs to be done the function needs to be repeated to ensure a correctly dimensioned display is saved. Once as a complete saved graphics file the image may be resized in the receiving document.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying and saving the plot.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range a truncated plot will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations.

In some R installations the generation of multi-panel displays and the use of function `eqsplot` from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a 'first' function prepared by the user that loads the 'rgr' package, etc.

Author(s)

Robert G. Garrett

References

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p. See pp. 119 for a description of histogram bin selection computations.

See Also

[inset](#), [ltdl.fix.df](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Usage is as follows
## Saves the graphics as a pdf file to the defined folde.r
## inset.exporter(Cu, xlab = "Cu (mg/kg) in\n<2 mm 0-horizon soil",
## log = TRUE, gtype = "pdf", file = "D://R_work//Project3")

## Detach test data
detach(kola.o)
```

kola.c

Kola Project C-horizon Soil Data

Description

These data arise from an ecogeochemical survey undertaken by the Central Kola Expedition of Russia (CKE), the Geological Survey of Finland (GTK) and the Norwegian Geological Survey (NGU). In 1995 a variety of soil and biological materials were collected from almost 700 sites lying between the Arctic Circle and the Barents Sea, and Longitudes 35.5 and 40 East. This specific data set is for C-horizon soils found at 606 of the sites visited. The data consist of an integer identifier, Universal Transverse Mercator (m) eastings and northings coordinates, the country the site was located in as a 3 character string, the lithology of the underlying bedrock as an integer code, 36 chemical measurements (total or near-total geochemical analyses), and soil pH for the <2 mm fraction of the C-horizon soils. The data reflect the natural geochemical variations in the parent material of the overlying soils. Further details concerning the project, methods of sampling and analysis can be found in Reimann et al. (1998) and the numerous papers published by the co-authors in international scientific journals.

Usage

kola.c

Format

A data frame containing 44 observations for 617 sites.

Source

These data are a subset of the full Kola C-horizon data set available from: <http://doi.pangaea.de/10.1594/PANGAEA.56227>

However, it should be noted that the spatial coordinates are recorded as Latitudes and Longitudes in the full data set.

References

Reimann, C., Ayras, M., Chekushin, V., Bogatyrev, I., Boyd, R., de Caritat, P., Dutter, R., Finne, T.E., Halleraker, J.H., Jaeger, O., Kashulina, G., Niskavaara, H., Pavlov, V., Raisanen, M.L., Strand, T. and Volden, T., 1998. A geochemical atlas of the central parts of the Barents Region. Norges Geologiske Undersokelse (NGU) Geological Survey of Norway, Trondheim, Norway. ISBN 82-7385-176-1. 745 p.

kola.o

Kola Project O-horizon Soil Data

Description

These data arise from an ecogeochemical survey undertaken by the Central Kola Expedition of Russia (CKE), the Geological Survey of Finland (GTK) and the Norwegian Geological Survey (NGU). In 1995 a variety of soil and biological materials were collected from almost 700 sites lying between the Arctic Circle and the Barents Sea, and Longitudes 35.5 and 40 East. This specific data set is for O-horizon soils found at 617 of the sites visited. The data consist of an integer identifier, Universal Transverse Mercator (m) eastings and northings coordinates, 38 chemical measurements (total or near-total geochemical analyses), Loss on Ignition, soil pH and specific conductivity for the <2 mm fraction of the O-horizon (humus) soils. The data reflect both natural biogeochemical variations and the presence of heavy industry. Further details concerning the project, and methods of sampling and analysis can be found in Reimann et al. (1998) and the numerous papers published by the co-authors in international scientific journals.

Usage

kola.o

Format

A data frame containing 44 observations for 617 sites.

Source

These data are the same as in the R package 'mvoutlier'. However, note that the names of the spatial coordinates have been changed from XCOO and YCOO to UTME and UTMN, respectively, and COND (specific conductivity) to SC.

The full data set is available from: <http://doi.pangaea.de/10.1594/PANGAEA.56279>

However, it should be noted that this is a superset containing all geochemical analyses and the spatial coordinates are recorded as Latitudes and Longitudes in the full data set.

References

Reimann, C., Ayras, M., Chekushin, V., Bogatyrev, I., Boyd, R., de Caritat, P., Dutter, R., Finne, T.E., Halleraker, J.H., Jaeger, O., Kashulina, G., Niskavaara, H., Pavlov, V., Raisanen, M.L., Strand, T. and Volden, T., 1998. A geochemical atlas of the central parts of the Barents Region. Norges Geologiske Undersokelse (NGU) Geological Survey of Norway, Trondheim, Norway. ISBN 82-7385-176-1. 745 p.

logit

Logit transformation

Description

Undertakes a logit transformation on a vector of proportions.

Usage

`logit(pp)`

Arguments

`pp` a vector of proportions in the range zero to one. The function may be used with a single proportion. Natural logarithms are used.

Details

Most analytical chemical data for major, minor and trace elements are of a closed form, i.e. for a sample they sum to a constant, whether it be percent, ppm (mg/kg), or some other units. It does not matter that only some components contributing to the constant sum are present in the matrix, the data are closed. As a result, as some elements increase in concentration others must decrease, this leads to statistics and graphical presentations that do not reflect the true underlying situation even in situations of univariate data analysis and display. The logit transformation is an appropriate transformation for univariate compositional data. However, for concentrations below 10% a logarithmic transform is sufficient. The inverse logit transform is the [expit](#). Procedures for removing closure effects for multivariate data are additive log-ratios ([alr](#)), centred log-ratios ([clr](#)), and isometric log-ratios ([ilr](#)).

Value

`z` a vector of the logit transformations of the proportions `p`.

Note

If a value outside the range zero to one is encountered as a proportion the function displays an error message and halts.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data matrix, must be removed prior to executing this function, see [ltdl.fix.df](#).

If any NAs exist in the vector, pp, they are removed by function `remove.na` and the number removed is displayed.

Author(s)

Robert G. Garrett

References

Filzmoser, P., Hron, K. and Reimann, C., 2009. Univariate statistical analysis of environmental (compositional) data: Problems and possibilities. *Science of the Total Environment*, 407(1/3):6100-6108.

See Also

`expit`, `alr`, `clr`, `ilr`, `ltdl.fix.df`

Examples

```
## Generate test data
p <- c(0.1, 0.5, 0.9)

## Undertake and display logit transformations
z <- logit(p)
z

## Clean-up
rm(p)
rm(z)
```

`ltdl.fix`

Replace Negative Values Representing Less Than Detects in a Vector

Description

Function to process a numeric vector to replace negative values representing less than detects (<DLs) with positive half that value. This permits processing of these effectively categorical data as real numbers and their display on logarithmically scaled axes. Some software packages replace blank fields that should be interpreted as NAs, i.e. no information, with zeros, the facility is provided to replace any zero values with NAs. In other instances data files have been built using an integer code, e.g., -9999, to indicate 'no data', i.e. the equivalent of NAs. The facility is provided to replace any so coded values with NAs. If required, all <values and may be replaced with NAs, e.g. when estimating analytical precision with `anova1` using only duplicate analyses with >DL values.

A report of the changes made is displayed on the current device.

For processing data matrices or data frames, see `ltdl.fix.df`.

Usage

```
ltdl.fix(x, negs2na = FALSE, zero2na = FALSE, coded = NA)
```

Arguments

x	name of the vector to be processed.
negs2na	to replace any -ve values with NAs, set negs2na = TRUE.
zero2na	to replace any zero values with NAs, set zero2na = TRUE.
coded	to replace any numeric coded values, e.g., -9999 with NAs, set coded = -9999.

Value

A numeric vector identical to that input but where any negative values have been replaced by half their positive values, or optionally by NAs, and optionally any zero or numeric coded values have been replaced by NAs.

Note

If data are being accessed through an ODBC link to a database, rather than from a data frame that can be processed by `ltdl.fix.df`, it may be important to run this function on the retrieved vector prior to any subsequent processing. The necessity for such vector processing can be ascertained using the `range` function, e.g., `range(na.omit(x))` or `range(x, na.rm = TRUE)`, where `x` is the variable name, to determine the presence of any negative values. The presence of any NAs in the vector will return NAs by `range` if they are not removed.

Great care needs to be taken when processing data where a large proportion of the data are less than detects (<value). In such cases parametric statistics have limited value, and can be misleading. Records should be kept of variables containing <values, and the fixed replacement values changed in tables for reports to the appropriate <values. Thus, in tables of percentiles the <value should replace the fixed value computed from $\text{absolute}(-\text{value})/2$. Various rules have been proposed as to how many less than detects treated in this way can be tolerated before means, variances, etc. become biased and of little value. Less than 5% in a large data set is usually tolerable, with greater than 10% concern increases, and with greater than 20% alternate procedures for processing the data should be sought, for example, the procedures outlined in Helsel (2005). Alternately replacement non-detect values may be imputed with R packages `robCompositions` (Hron et al., 2010; Templ et al., 2011) or `zCompositions` (Martin-Fernandez et al., 2012; Palarea-Albaladejo and Martin-Fernandez, 2013), specifically function `lrEM` in the latter, both may be used with closed, constant-sum, data.

Author(s)

Robert G. Garrett

References

- Helsel, D.R., 2005. *Nondetects and Data Analysis: Statistics for Censored Data*. John Wiley & Sons, Ltd., 250 p.
- Hron, K., Templ, M. and Filzmoser, P., 2010. Imputation of missing values for compositional data using classical and robust methods. *Computational Statistics and Data Analysis*, 54(12):3095-3107. 3107.
- Templ, M., Hron, K. and Filzmoser, P., 2010. `robCompositions`: An R-package for Robust Statistical Analysis of Composition data. In: *Compositional Data Analysis: Theory and Applications*, V. Pawowsky and A. Buccianti (Eds.), Chapter 25, pp. 342-355. John Wiley & Sons, Ltd.

Martin-Fernandez, J.A., Hron, K., Templ, M., Filzmoser, P. and Palarea-Albaladejo, J., 2012. Model-based replacement of rounded zeros in compositional data: classical and robust approaches. *Computational Statistics and Data Analysis*, 56(9):2688-2704.

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See Also

[ltdl.fix.df](#)

Examples

```
## Replace any missing data coded as -9999 with NAs and any remaining
## negative values representing less than detects with Abs(value)/2
data(fix.test)
x.fixed <- ltdl.fix(fix.test[, 3], coded = -9999)
x.fixed

## As above, and replace any zero values with NAs
x.fixed <- ltdl.fix(fix.test[, 3], coded = -9999, zero2na = TRUE)
x.fixed

## As above, but replace any negative values with NAs
x.fixed <- ltdl.fix(fix.test[, 3], negs2na = TRUE, coded = -9999, zero2na = TRUE)
x.fixed

## Make test data kola.o available, setting a -9999, indicating a
## missing pH measurement, to NA
data(kola.o)
attach(kola.o)
pH.fixed <- ltdl.fix(pH, coded = -9999)

## Display relationship between pH in one pH unit intervals and Cu in
## 0-horizon (humus) soil, extending the whiskers to the 2nd and 98th
## percentiles, finally removing the temporary data vector pH.fixed
bwplots(split(Cu, trunc(pH.fixed+0.5)), log = TRUE, wend = 0.02,
xlab = "Soil pH to the nearest pH unit",
ylab = "Cu (mg/kg) in < 2 mm Kola 0-horizon soil")
rm(pH.fixed)

## Or directly
bwplots(split(Cu, trunc(ltdl.fix(pH, coded = -9999)+0.5)), log = TRUE,
wend = 0.02, xlab = "Soil pH to the nearest pH unit",
ylab = "Cu (mg/kg) in < 2 mm Kola 0-horizon soil")

## Clean-up and detach test data
rm(x)
rm(x.fixed)
rm(pH.fixed)
detach(kola.o)
```

ltdl.fix.df

*Replace Negative Values Representing Less Than Detects in a Data Frame***Description**

Function to process a matrix or data frame to replace negative values representing less than detects (<value) with positive half that value. This permits processing of these effectively categorical data as real numbers and their display on logarithmically scaled axes. Some software packages replace blank fields that should be interpreted as NAs, i.e. no information, with zeros, the facility is provided to replace any zero values with NAs. In other instances data files have been built using an integer code, e.g., -9999, to indicate 'no data', i.e. the equivalent of NAs. The facility is provided to replace any so coded values with NAs. If required, all <values and may be replaced with NAs, e.g. when estimating analytical precision with `anova1` using only duplicate analyses with >DL values. Any factor variables in the input matrix or data frame are passed to the output matrix or data frame.

If a single vector is to be processed, use `ltdl.fix`

A report of the changes made is displayed on the current device.

Usage

```
ltdl.fix.df(x, negs2na = FALSE, zero2na = FALSE, coded = NA)
```

Arguments

x	name of the matrix or data frame to be processed.
negs2na	to replace any -ve values with NAs, set negs2na = TRUE.
zero2na	to replace any zero values with NAs, set zero2na = TRUE.
coded	to replace any numeric coded values, e.g., -9999 with NAs, set coded = -9999.

Value

A matrix or data frame identical to that input but where any negative values have been replaced by half their positive values, or NAs, and optionally any zero values or numeric coded values have been replaced by NAs.

Note

Great care needs to be taken when processing data where a large proportion of the data are less than detects (<value). In such cases parametric statistics have limited value, and can be misleading. Records should be kept of variables containing <values, and the fixed replacement values changed in tables for reports to the appropriate <values. Thus, in tables of percentiles the <value should replace the fixed value computed from $\text{absolute}(-\text{value})/2$. Various rules have been proposed as to how many less than detects treated in this way can be tolerated before means, variances, etc. become biased and of little value. Less than 5% in a large data set is usually tolerable, with greater than 10% concern increases, and with greater than 20% alternate procedures for processing the data should be sought. For example, the procedures outlined in Helsel (2005). Alternately

replacement non-detect values may be imputed with R packages `robCompositions` (Hron et al., 2010; Templ et al., 2011) or `zCompositions` (Martin-Fernandez et al., 2012; Palarea-Albaladejo and Martin-Fernandez, 2013), specifically function `lrEM` in the latter, both may be used with closed, constant-sum, data.

Author(s)

Robert G. Garrett and David Lorenz

References

- Helsel, D.R., 2005. *Nondetects and Data Analysis: Statistics for Censored Data*. John Wiley & Sons, Ltd., 250 p.
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- Templ, M., Hron, K. and Filzmoser, P., 2010. `robCompositions`: An R-package for Robust Statistical Analysis of Composition data. In: *Compositional Data Analysis: Theory and Applications*, V. Pawowsky and A. Buccianti (Eds.), Chapter 25, pp. 342-355. John Wiley & Sons, Ltd.
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See Also

[ltdl.fix](#)

Examples

```
## Replace any missing data coded as -9999 with NAs and any remaining
## negative values representing less than detects with Abs(value)/2
data(fix.test)
fix.test
fix.test.fixed <- ltdl.fix.df(fix.test, coded = -9999)
fix.test.fixed

## As above, and replace any zero values with NAs
fix.test.fixed <- ltdl.fix.df(fix.test, coded = -9999, zero2na = TRUE)
fix.test.fixed

## As above, but replace any negative values with NAs
fix.test.fixed <- ltdl.fix.df(fix.test, negs2na = TRUE, coded = -9999, zero2na = TRUE)
fix.test.fixed

## Clean-up
rm(fix.test.fixed)
```

Description

Displays a simple map where the data are represented at their spatial locations by symbols using Tukey boxplot-based symbology. Tukey boxplots divide data into 7 groups, the middle 50%, and three lower and higher groupings, see Details below. The computation of the fences used to subdivide the data may be carried out following a logarithmic transformation of the data. The colours of the symbols may be optionally changed. Optionally a legend may be added to the map.

Usage

```
map.eda7(xx, yy, zz, sfact = 1, logz = FALSE, xlab = "Easting",
         ylab = "Northing", zlab = deparse(substitute(zz)), main = "",
         ifgrey = FALSE, symcolr = NULL, tol = 0.04, iflgnd = FALSE,
         title = deparse(substitute(zz)), cex.lgnd = 0.8, ...)
```

Arguments

xx	name of the x-axis spatial coordinate, the eastings.
yy	name of the y-axis spatial coordinate, the northings.
zz	name of the variable to be plotted.
sfact	controls the absolute size of the plotted symbols, by default <code>sfact = 1</code> . Increasing <code>sfact</code> results in larger symbols.
xlab	a title for the x-axis, defaults to <code>Easting</code> .
ylab	a title for the y-axis, defaults to <code>Northing</code> .
zlab	by default, <code>zlab = deparse(substitute(zz))</code> , a map title is generated by appending the input variable name text string to "EDA Tukey Boxplot Map for ". Alternative titles may be generated, see Details below.
main	an alternative map title, see Details below.
logz	if it is required to undertake the Tukey Boxplot computations after a logarithmic data transform, set <code>logz = TRUE</code> .
ifgrey	set <code>ifgrey = TRUE</code> if a grey-scale map is required, see Details below.
symcolr	the default is a colour map and default colours are provided, deeper blues for lower values, green for the middle 50% of the data, and oranges and reds for higher values. A set of alternate symbol colours can be provided by defining <code>symcolr</code> , see Details below.
tol	a parameter used to ensure the area included within the neatline around the map is larger than the distribution of the points so that the plotted symbols fall within the neatline. By default <code>tol = 0.04</code> , if more clearance is required increase the value of <code>tol</code> .

<code>iflgnd</code>	the default is no legend. If a legend is required set <code>iflgnd = TRUE</code> , following the plotting of the data the cursor will be activated, locate that at the top left corner of the desired legend position and ‘left button’ on the pointing device.
<code>title</code>	a short title for the legend, e.g., <code>title = "Cu (mg/kg)"</code> . The default is the character string for <code>zz</code> .
<code>cex.lgnd</code>	controls the scaling of the legend box and text, but not the symbols so that they match those on the map, the default is <code>cex.lgnd = 0.8</code> .
<code>...</code>	further arguments to be passed to methods. For example, if it is required to make the map title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

Tukey boxplots divide data into 7 groups, the middle 50%, and three lower and higher groupings: within the whisker, near outliers and far outliers, respectively. Symbols for values below the first quartile (Q1) are plotted as increasingly larger circles, while symbols for values above the third quartile are plotted as increasingly larger squares, a ‘+’ is used to plot the data falling in the middle 50%. For the higher groupings, the whisker contains values $>Q3$ and $<(Q3 + 1.5 * HW)$, where $HW = (Q3 - Q1)$, the interquartile range; near outliers lie between $(Q3 + 1.5 * HW)$ and $(Q3 + 3 * HW)$; and far outliers have values $>(Q3 + 3 * HW)$. For the lower groupings the group boundaries, fences, fall similarly spaced below Q1. The computation of the fences used to subdivide the data may be carried out following a logarithmic transformation of the data, set `logz = TRUE`.

A summary table of the values of the symbol intervals, the number of values plotting as each symbol, and symbol shapes, sizes and colours is displayed on the current device.

If `zlab` and `main` are undefined a default a map title is generated by appending the input variable name text string to "EDA Tukey Boxplot-Based Map for ". If no map title is required set `zlab = ""`, and if some user defined map title is required it should be defined in `main`, e.g. `main = "Map Title Text"`.

If the grey-scale option is chosen the symbols are plotted 100% black for the far outliers, 85% black for the near outliers, 70% black for values within the whiskers, and 60% black for values falling within the middle 50% of the data.

The default colours, `symcolr = c(25, 22, 20, 13, 6, 4, 1)`, are selected from the `rainbow(36)` palette, and alternate colour schemes need to be selected from the same palette. See [display.rainbow](#) for the available colours. It is essential that 7 colours be provided, e.g., `symcolr = c(27, 24, 22, 12, 5, 3, 36)`, if exactly 7 are not provided the default colours will be displayed.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors including NAs are removed prior to displaying the plot.

In some R installations the generation of multi-panel displays and the use of function `eqsplot` from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a ‘first’ function prepared by the user that loads the ‘rgr’ package, etc.

Author(s)

Robert G. Garrett

See Also[display.rainbow](#), [ltdl.fix.df](#), [remove.na](#)**Examples**

```
## Make test data available
data(kola.o)
attach(kola.o)

## Plot a default symbol map
map.eda7(UTME, UTMN, Cu)

## Plot with logarithmically scaled boxplot fences and more
## appropriate axis labelling
map.eda7(UTME/1000, UTMN/1000, Cu, logz = TRUE,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)")

## Plot a grey-scale equivalent of the above map
map.eda7(UTME/1000, UTMN/1000, Cu, logz = TRUE, ifgrey = TRUE,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)")

## Plot the same map with an alternate colour scheme
map.eda7(UTME/1000, UTMN/1000, Cu, logz = TRUE,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)",
symcolr = c(27, 24, 22, 12, 5, 3, 36))

## Detach test data
detach(kola.o)
```

`map.eda8`*Plot a Symbol Map of Data Based on their Percentiles*

Description

Displays a simple map where the data are represented at their spatial locations by symbols indicating within which group defined by the data's 2nd, 5th, 25th, 50th, 75th, 95th and 98th percentiles plotted a data value falls. The colours of the symbols may be optionally changed. Optionally a legend (two options) may be added to the map.

Usage

```
map.eda8(xx, yy, zz, sfact = 1, xlab = "Easting", ylab = "Northing",
zlab = deparse(substitute(z)), main = "", ifgrey = FALSE,
symcolr = NULL, tol = 0.04, iflgnd = FALSE, pctile = FALSE,
title = deparse(substitute(z)), cex.lgnd = 0.8, ...)
```

Arguments

xx	name of the x-axis spatial coordinate, the eastings.
yy	name of the y-axis spatial coordinate, the northings.
zz	name of the variable to be plotted.
sfact	controls the absolute size of the plotted symbols, by default <code>sfact = 1</code> . Increasing <code>sfact</code> results in larger symbols.
xlab	a title for the x-axis, defaults to <code>Easting</code> .
ylab	a title for the y-axis, defaults to <code>Northing</code> .
zlab	by default, <code>zlab = deparse(substitute(z))</code> , a map title is generated by appending the input variable name text string to "EDA Percentile Based Map for ". Alternative titles may be generated, see Details below.
main	an alternative map title, see Details below.
ifgrey	set <code>ifgrey = TRUE</code> if a grey-scale map is required, see Details below.
symcolr	the default is a colour map and default colours are provided, deeper blues for lower values, green for the middle 50% of the data, and oranges and reds for higher values. A set of alternate symbol colours can be provided by defining <code>symcolr</code> , see Details below.
tol	a parameter used to ensure the area included within the newline around the map is larger than the distribution of the points so that the plotted symbols fall within the newline. By default <code>tol = 0.04</code> , if more clearance is required increase the value of <code>tol</code> .
iflgnd	the default is no legend. If a legend is required set <code>iflgnd = TRUE</code> , following the plotting of the data the cursor will be activated, locate that at the top left corner of the desired legend position and 'left button' on the pointing device. There are two legends to choose from, see <code>pctile</code> below.
pctile	the default legend displays the range of values each symbol represents. Alternately, the percentiles may be displayed rather than their values by setting <code>pctile = TRUE</code> .
title	a short title for the legend, e.g., <code>title = "Cu (mg/kg)"</code> . The default is the character string for <code>zz</code> .
cex.lgnd	controls the scaling of the legend box and text, but not the symbols so that they match those on the map, the default is <code>cex.lgnd = 0.8</code> .
...	further arguments to be passed to methods. For example, if it is required to make the map title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

The selected percentiles, 2nd, 5th, 25th, 50th, 75th, 95th and 98th, divide the data into 8 groups. Values below the median are represented by increasingly larger deeper blue circles below the 25th percentile (Q1), and values above the 75th percentile (Q3) by increasingly larger orange and red squares. The mid 50% of the data are represented by green symbols, circles for the median (Q2) to Q1, and squares for the median (Q2) to Q3.

A summary table of the values of the symbol intervals, the number of values plotting as each symbol, and symbol shapes, sizes and colours is displayed on the current device.

If `zlab` and `main` are undefined a default a map title is generated by appending the input variable name text string to "EDA Percentile Based Map for ". If no map title is required set `zlab = ""`, and if some user defined map title is required it should be defined in `main`, e.g. `main = "Map Title Text"`.

If the grey-scale option is chosen the symbols are plotted 100% black for the far outliers, 85% black for the near outliers, 70% black for values within the whiskers, and 60% black for values falling within the middle 50% of the data.

The default colours, `symcolr = c(25, 22, 20, 13, 13, 6, 4, 1)`, are selected from the `rainbow(36)` palette, and alternate colour schemes need to be selected from the same palette. See [display.rainbow](#) for the available colours. It is essential that 8 colours be provided, e.g., `symcolr = c(27, 24, 22, 12, 12, 5, 3, 36)`, if exactly 8 are not provided the default colours will be displayed.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors including NAs are removed prior to displaying the plot.

In some R installations the generation of multi-panel displays and the use of function `eqscplot` from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a 'first' function prepared by the user that loads the 'rgr' package, etc.

Author(s)

Robert G. Garrett

See Also

[display.rainbow](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)
```

```

## Plot a default symbol map
map.eda8(UTME, UTMN, Cu)

## Plot a more appropriately labelled map
map.eda8(UTME/1000, UTMN/1000, Cu,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)")

## Plot a grey-scale equivalent of the above map
map.eda8(UTME/1000, UTMN/1000, Cu, ifgrey = TRUE,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)")

## Plot the same map with an alternate colour scheme
map.eda8(UTME/1000, UTMN/1000, Cu,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)",
symcolr = c(27, 24, 22, 12, 12, 5, 3, 36))

## Detach test data
detach(kola.o)

```

map.tags

Plot a Map of Posted Values

Description

Displays a simple map where the data are represented by the ‘written’ values of the data at their spatial locations.

Usage

```
map.tags(xx, yy, tag, xlab = "Easting", ylab = "Northing",
taglab = deparse(substitute(tag)), main = "", tol = 0.04, ...)
```

Arguments

xx	name of the x-axis spatial coordinate, the eastings.
yy	name of the y-axis spatial coordinate, the northings.
tag	name of the variable to be plotted as a map.
xlab	a title for the x-axis, defaults to Easting.
ylab	a title for the y-axis, defaults to Northing.
taglab	by default, taglab = deparse(substitute(tag)), a map title is generated by appending the input variable name text string to "Map of Values for ". Alternative titles may be generated, see Details below.
main	an alternative map title, see Details below.

`tol` a parameter used to ensure the area included within the neatline around the map is larger than the distribution of the points so that the plotted symbols fall within the neatline. By default `tol = 0.04`, if more clearance is required increase the value of `tol`.

`...` further arguments to be passed to methods. For example, if smaller plotting characters are required, specify `cex = 0.8`; or if some colour other than black is required for the plotting characters, specify `col = 2` to obtain red (see [display.lty](#) for the default colour palette). If it is required to make the map title smaller, add `cex.main = 0.9` to reduce the font size by 10%.

Details

If `taglab` and `main` are undefined a default a map title is generated by appending the input variable name text string to "Map of Values for ". If no map title is required set `taglab = ""`, or if an alternative to the variable name `taglab` is required it may be specified, `taglab = "Alternative"`. If some user defined map title is required it should be defined in `main`, e.g. `main = "Map Title Text"`, in which instance `taglab` is ignored.

If a map of sample numbers, 'IDs', is required and they are not explicitly in the data frame, a map of data frame row numbers may be displayed by specifying `dimnames(dfname)[[1]]` as the value of `tags`.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors including NAs for spatial coordinates are removed prior to displaying the map, thus those 'sites' are not plotted. However, where coordinates are present any NAs in the variable to be plotted are replaced with a '+' sign to indicate sites with 'missing data'.

In some R installations the generation of multi-panel displays and the use of function `eqscplot` from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a 'first' function prepared by the user that loads the 'rgr' package, etc.

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [remove.na](#), [display.lty](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Plot a sample site number map
```

```

map.tags(E, N, ID)

## Plot a sample site number map with smaller numbers
## and a wider internal map margin
map.tags(E, N, ID, cex = 0.8, tol = 0.06)

## Plot the data frame row numbers rather than the sample
## numbers
map.tags(E, N, dimnames(sind)[[1]], cex = 0.8, tol = 0.06)

## Plot the values for Zn in red, providing enough internal
## map margin so the values do not overprint the map neat-line
map.tags(E, N, Zn, cex = 0.8, tol = 0.1, col = 2)

## Plot as above but with an informative title spread over
## two lines and with a slightly smaller font
map.tags(E, N, Zn, cex = 0.8, tol = 0.1, col = 2, main =
"Horwarth & Sinding-Larsen\nStream Sediment Zn Data",
cex.main = 0.9)

## Detach test data
detach(sind)

```

map.z

Plot a Map of Data using Proportional Symbols

Description

Displays a simple map where the data are represented by open circles whose diameters are proportional to the value of the data at their spatial locations. The rate of change of symbol diameter with value and the absolute size of the symbols are defined by the user. Optionally a legend may be displayed on the map.

Usage

```

map.z(xx, yy, zz, p = 0.5, sfact = 2.5, zmin = NA, zmax = NA,
xlab = "Easting", ylab = "Northing",
zlab = deparse(substitute(zz)), main = "", tol = 0.04,
col = 1, iflgnd = FALSE, title = deparse(substitute(zz)),
cex.lgnd = 0.8, ifparams = FALSE, cex.params = 0.8, ...)

```

Arguments

xx	name of the x-axis spatial coordinate, the eastings.
yy	name of the y-axis spatial coordinate, the northings.
zz	name of the third variable, z, to be plotted as a map.
p	a parameter that controls the rate of change of symbol diameter with changing value. A default of $p = 0.5$ is provided. See Details below.

<code>sfact</code>	controls the absolute size of the plotted symbols, by default <code>sfact = 2.5</code> . Increasing <code>sfact</code> results in larger symbols.
<code>zmin</code>	a value below which all symbols will be plotted at the same minimum size. By default <code>zmin = NA</code> which results in the minimum value of the variable defining the minimum symbol size. See Details below.
<code>zmax</code>	a value above which all symbols will be plotted at the same maximum size. By default <code>zmax = NA</code> which results in the maximum value of the variable defining the maximum symbol size. See Details below.
<code>xlab</code>	a title for the x-axis, defaults to <code>Easting</code> .
<code>ylab</code>	a title for the y-axis, defaults to <code>Northing</code> .
<code>zlab</code>	by default, <code>zlab = deparse(substitute(z))</code> , a map title is generated by appending the input variable name text string to "Proportional Symbol Map for ". Alternative titles may be provided, see Details below.
<code>main</code>	an alternative map title, see Details below.
<code>tol</code>	a parameter used to ensure the area included within the neatline around the map is larger than the distribution of the points so that the plotted symbols fall within the neatline. By default <code>tol = 0.04</code> , if more clearance is required increase the value of <code>tol</code> .
<code>col</code>	the colour of the symbols, the default is black, <code>col = 1</code> . This may be changed if required, see display.lty for the default colour palette. For example, <code>col = 2</code> will cause the symbols to be plotted in red.
<code>iflgnd</code>	the default is no legend. If a legend is required set <code>iflgnd = TRUE</code> , following the plotting of the data the cursor will be activated, locate that at the top left corner of the desired legend position and 'left button' on the pointing device. See Notes below.
<code>title</code>	a short title for the legend, e.g., <code>title = "Zn (mg/kg)"</code> . The default is the character string for <code>zz</code> .
<code>cex.lgnd</code>	controls the scaling of the legend box and text, but not the symbols so that they match those on the map, the default is <code>cex.lgnd = 0.8</code> .
<code>ifparams</code>	if <code>ifparams = TRUE</code> on completion of plotting and after the legend has been plotted, if requested, the cursor is activated, locate that at the top left corner of the desired text position and 'left button' on the pointing device. This text comprises three lines: the values of <code>p</code> to three significant figures and <code>sfact</code> ; the maximum value of <code>z</code> to 3 significant figures and <code>zmax</code> ; and the minimum value of <code>z</code> to 3 significant figures and <code>zmin</code> . The default is no text display.
<code>cex.params</code>	controls the size of the text for the parameters display, defaults to <code>cex.params = 0.8</code> .
<code>...</code>	further arguments to be passed to methods. For example, if it is required to make the map title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

The symbol diameter is computed as a function of the value `z` to be plotted:

$$\text{diameter} = \text{dmin} + (\text{dmax} - \text{dmin}) * \{(z - \text{zmin}) / (\text{zmax} - \text{zmin})\}^p$$

where `dmin` and `dmax` are defined as 0.1 and 1 units, so the symbol diameters range over an order

of magnitude (and symbol areas over two); z_{\min} and z_{\max} are the observed range of the data, or the range over which the user wants the diameters to be computed; and p is a power defined by the user. The value of $(z - z_{\min}) / (z_{\max} - z_{\min})$ is the value of z normalized, 0 - 1, to the range over which the symbol diameters are to be computed. After being raised to the power p , which will result in a number in the range 0 to 1, this value is multiplied by the permissible range of diameters and added to the minimum diameter. This results in a diameter between 0.1 and 1 units that is proportional to the value of z .

A p value of 1 results in a linear rate of change. Values of p less than unity lead to a rapid initial rate of change with increasing value of z which is often suitable for displaying positively skewed data sets, see the example below. In contrast, values of p greater than unity result in an initial slow rate of change with increasing value of z which is often suitable for displaying negatively skewed data sets. To obtain a logarithmic scaling set $p = 0.3679$, i.e. $1/e$, and a value of $p = 0.5$ is equivalent to a square root transformation, the default, that is appropriate for a linear-area relationship. These values of p are equivalent to plotting $\log(x)$ or $\log_{10}(x)$ and \sqrt{x} , respectively. Experimentation is often necessary to obtain a satisfactory visual effect. See [syms.pfunc](#) for a graphic demonstrating the effect of varying the p parameter.

If z_{\min} or z_{\max} are defined this has the effect of setting a minimum or maximum value of z , respectively, beyond which changes in the value of z do not result in changes in symbol diameter. This can be useful in limiting the effect of one, or a few, extreme outlier(s) while still plotting them, they simply plot at the minimum or maximum symbol size and are not involved in the calculation of the range of z over which the symbol diameters vary. **Note:** If the variable z includes a transform, e.g., $\log_{10}(z)$, the values of z_{\min} and/or z_{\max} must be in those transform units.

If z_{lab} and $main$ are undefined a default a map title is generated by appending the input variable name text string to "Proportional Symbol Map for ". If no map title is required set $z_{\text{lab}} = ""$, and if some user defined map title is required it should be defined in $main$, e.g. $main = "Map Title Text"$.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

The legend consists of five proportional symbols and their corresponding z values: z_{\min} ; the three quartiles; and z_{\max} . If z_{\min} and z_{\max} have been user defined it is over their range that the symbol sizes are computed and displayed. When defining z_{\min} and/or z_{\max} it is useful to set $ifparams = TRUE$ as a reminder, whilst developing the required display.

Any NAs in the data vector are removed prior to displaying the plot.

In some R installations the generation of multi-panel displays and the use of function `eqsplot` from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a 'first' function prepared by the user that loads the 'rgr' package, etc.

Author(s)

Robert G. Garrett

See Also

[syms](#), [syms.pfunc](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Plot a default symbol map, p = 0.5 and sfact = 2.5
map.z(UTME, UTMN, Cu)

## Plot a map where the symbols are logarithmically scaled,
## and more appropriately labelled axes
map.z(UTME/1000, UTMN/1000, Cu, p = 0.3679,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)" )

## Plot with differently scaled symbols and more appropriately
## labelled axes
map.z(UTME/1000, UTMN/1000, Cu, p = 0.3, sfact = 2.0,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)" )

## Plot a map as above but where outliers above a value of 1000 are
## displayed with the same symbol
map.z(UTME/1000, UTMN/1000, Cu, p = 0.3, sfact = 2.0, zmax = 1000,
xlab = "Kola Project UTM Eastings (km)",
ylab = "Kola Project UTM Northings (km)" )

## Detach test data
detach(kola.o)
```

ms.data1

Measurement Variability Test Data

Description

A data frame of magnetic susceptibility data used by Stanley (2003) to demonstrate the Thompson-Howarth procedure for estimating analytical variability (Thompson and Howarth, 1973 & 1978). They are used in the `rg` package examples for the duplicate analysis ANOVA and Thompson-Howarth plot functions, [anova1](#) and [thplot1](#), respectively. See also Garrett and Grunsky (2003).

Usage

```
ms.data1
```


Format

A data frame containing 2 measurements of magnetic susceptibility for each of 16 rock samples in 16 records.

Source

Stanley (2003), see below.

References

Garrett, R.G. and Grunsky, E.C., 2003. S and R functions for the display of Thompson-Howarth plots. *Computers & Geosciences*, 29(2):239-242.

Stanley, C.R., 2003. THPLOT.M: a MATLAB function to implement generalized Thompson-Howarth error analysis using replicate data. *Computers & Geosciences*, 29(2):225-237.

Thompson, M. and Howarth, R.J., 1973. The rapid estimation and control of precision by duplicate determinations. *The Analyst*, 98(1164):153-160.

Thompson, M. and Howarth, R.J., 1978. A new approach to the estimation of analytical precision. *Journal of Geochemical Exploration*, 9(1):23-30.

ms.data2

Measurement Variability Test Data

Description

A data frame of magnetic susceptibility data used by Stanley (2003) to demonstrate the Thompson-Howarth procedure for estimating analytical variability (Thompson and Howarth, 1973 & 1978). They are used in the `rgr` package examples for the duplicate analysis ANOVA and Thompson-Howarth plot functions, `anova2` and `thplot2`, respectively, with `ifalt = FALSE`. See also Garrett and Grunsky (2003).

Usage

ms.data2

Format

A data frame containing 2 measurements of magnetic susceptibility for each of 16 rock samples in 32 records. The measurements for the original analyses are in records 1 to 16, and the duplicate measurements are in records 17 to 32 in the same order.

Source

Stanley (2003), see below.

References

- Garrett, R.G. and Grunsky, E.C., 2003. S and R functions for the display of Thompson-Howarth plots. *Computers & Geosciences*, 29(2):239-242.
- Stanley, C.R., 2003. THPLOT.M: a MATLAB function to implement generalized Thompson-Howarth error analysis using replicate data. *Computers & Geosciences*, 29(2):225-237.
- Thompson, M. and Howarth, R.J., 1973. The rapid estimation and control of precision by duplicate determinations. *The Analyst*, 98(1164):153-160.
- Thompson, M. and Howarth, R.J., 1978. A new approach to the estimation of analytical precision. *Journal of Geochemical Exploration*, 9(1):23-30.

ms.data3

Measurement Variability Test Data

Description

A data frame of magnetic susceptibility data used by Stanley (2003) to demonstrate the Thompson-Howarth procedure for estimating analytical variability (Thompson and Howarth, 1973 & 1978). They are used in the `rgr` package examples for the duplicate analysis ANOVA and Thompson-Howarth plot functions, `anova2` and `thplot2`, respectively. See also Garrett and Grunsky (2003).

Usage

ms.data3

Format

A data frame containing 2 measurements of magnetic susceptibility for each of 16 rock samples in 32 records. The measurements for the original and duplicate analyses alternate. So the first duplicate pair are in records 1 and 2, and the last in records 31 and 32.

Source

Stanley (2003), see below.

References

- Garrett, R.G. and Grunsky, E.C., 2003. S and R functions for the display of Thompson-Howarth plots. *Computers & Geosciences*, 29(2):239-242.
- Stanley, C.R., 2003. THPLOT.M: a MATLAB function to implement generalized Thompson-Howarth error analysis using replicate data. *Computers & Geosciences*, 29(2):225-237.
- Thompson, M. and Howarth, R.J., 1973. The rapid estimation and control of precision by duplicate determinations. *The Analyst*, 98(1164):153-160.
- Thompson, M. and Howarth, R.J., 1978. A new approach to the estimation of analytical precision. *Journal of Geochemical Exploration*, 9(1):23-30.

Description

A data frame of major, minor and trace element data for the O'Grady pluton, NWT, (NTS map sheet 105I) from a regional lithogeochemical survey undertaken by the Geological Survey of Canada between 1969 and 1972 of Cretaceous-age granitoid plutons northeast of the Tintina Trench in the Yukon and adjoining NWT. Samples were collected in pairs from each site and a sub-sample ground to <100 mesh. Major and, some trace, element analyses were undertaken by direct reading optical spectroscopy (OES) either after a Li-Tetraborate (Li-T) fusion, or directly, other major elements were determined by AAS following HNO₃ dissolution of the fusion product. Other trace-elements were determined by atomic absorption spectrophotometry after a HF-HClO₄ digestion, or colorimetry (Col) after an alkaline fusion (AF). For Na, K, Fe, Mo and W detection limits (DLs) were 0.05, 0.1 and 0.1 %, and 0.5 and 2 mg/kg, respectively; <DL observations are represented by values of 0.02, 0.05, 0.05 % and 0.2 and 1 mg/kg, respectively.

Usage

data(ogrady)

Format

A data frame with 110 observations for the following 24 variables:

ID a numeric vector, part of the unique GSC sample number.

E UTM Eastings (m) for the sample site (UTM Zone 9).

N UTM Northings (m) for the sample site (UTM Zone 9).

Lith field name for the sampled lithology.

Si silicon (%) in granitoid (Li-T OES).

Al aluminium (%) in granitoid (Li-T OES).

Fe iron (%) in granitoid (Li-T HNO₃ AAS).

Mg magnesium (%) in granitoid (Li-T OES).

Ca calcium (%) in granitoid (Li-T OES).

Na sodium (%) in granitoid (Li-T HNO₃ AAS).

K potassium (%) in granitoid (Li-T HNO₃ AAS).

Ti titanium (mg/kg) in granitoid (Li-T OES).

Mn manganese (mg/kg) in granitoid (Li-T OES).

Ba barium (mg/kg) in granitoid (Li-T OES).

Zn zinc (mg/kg) in granitoid (HF-HClO₄ AAS).

Cu copper (mg/kg) in granitoid (HF-HClO₄ AAS).

Pb lead (mg/kg) in granitoid (HF-HClO₄ AAS).

- Mo** molybdenum (mg/kg) in granitoid (AF Col).
W tungsten (mg/kg) in granitoid. (AF Col).
U uranium (mg/kg) in granitoid (HF-HClO₄ Fluorimetry).
Be beryllium (mg/kg) in granitoid (OES).
V vanadium (mg/kg) in granitoid (OES).
Sn tin (mg/kg) in granitoid (OES).
Zr zirconium (mg/kg) in granitoid (OES).

Source

Garrett, R.G., 1992. Lithogeochemical data release, major and trace elements in Cretaceous granitoid rocks in the Yukon Territory and adjoining parts of the N.W.T. (95E, L, 105H, I, J, K, L, M, N, O, P, 106D, 115P, 116D). Geological Survey of Canada Open File 2479, digital data.

References

Garrett, R.G., 1972. Regional geochemical study of Cretaceous acidic igneous rocks in the northern Canadian Cordillera as a tool for broad mineral exploration. in Proc. 4th International Geochemical Exploration Symp., Geochemical Exploration 1972 (Ed. M.J. Jones). Institute of Mining and Metallurgy, London, pp. 203-219.

ogrady.mat2open

Lithogeochemical Data Set from the O'Grady Pluton, NWT, 1970

Description

A matrix of major and minor element data (see below) for the O'Grady pluton, NWT, (NTS map sheet 105I), see [ogrady](#) for further details. Additionally, the UTM coordinates and lithology classifications have been removed from the ogrady data frame. This data set is provided so that users can investigate different approaches to the closure problem using the various functions in package 'rgr'.

Usage

```
data(ogrady.mat2open)
```

Format

A matrix with 110 observations for the following 10 variables:

- Si** silicon (mg/kg) in granitoid (Li-T OES).
Al aluminium (mg/kg) in granitoid (Li-T OES).
Fe iron (mg/kg) in granitoid (Li-T HNO₃ AAS).
Mg magnesium (mg/kg) in granitoid (Li-T OES).
Ca calcium (mg/kg) in granitoid (Li-T OES).

- Na** sodium (mg/kg) in granitoid (Li-T HNO₃ AAS).
K potassium (mg/kg) in granitoid (Li-T HNO₃ AAS).
Ti titanium (mg/kg) in granitoid (Li-T OES).
Mn manganese (mg/kg) in granitoid (Li-T OES).
Ba barium (mg/kg) in granitoid (Li-T OES).

Source

Garrett, R.G., 1992. Lithochemical data release, major and trace elements in Cretaceous granitoid rocks in the Yukon Territory and adjoining parts of the N.W.T. (95E, L, 105H, I, J, K, L, M, N, O, P, 106D, 115P, 116D). Geological Survey of Canada Open File 2479, digital data.

References

Garrett, R.G., 1972. Regional geochemical study of Cretaceous acidic igneous rocks in the northern Canadian Cordillera as a tool for broad mineral exploration. in Proc. 4th International Geochemical Exploration Symp., Geochemical Exploration 1972 (Ed. M.J. Jones). Institute of Mining and Metallurgy, London, pp. 203-219.

See Also

[ogradey](#)

orthonorm

Computation of an Orthonormal Basis Matrix

Description

Computes an orthonormal basis matrix to be used for the back-transformation of ilr-based data and statistics to clr-based data and statistics.

Usage

orthonorm(p)

Arguments

p the dimension of the p-space, the number of original variables.

Value

V the p by (p-1) orthonormal basis matrix.

Author(s)

Based on a function by Peter Filzmoser and Karel Hron

References

Filzmoser, P., Hron, K., Reimann, C. and Garrett, R., 2009. Robust factor analysis for compositional data. *Computers & Geosciences*, 35(9):1854-1861.

See Also

[ilr](#), [clr](#), [gx.mva.closed](#), [gx.robmva.closed](#), [gx.md.gait.closed](#)

Examples

```
## Make test data available
data(sind.mat2open)

## Compute and display clr transformed data
prmatrix(clr(sind.mat2open))

## Compute and display ilr transformed data
sind.ilr <- ilr(sind.mat2open)
prmatrix(sind.ilr)

## Compute and display orthonormal basis matrix
## sind.mat2open is a 25 by 6 matrix (data set)
V <- orthonorm(6)
prmatrix(V)

## Back-transform ilr transformed data to clr form and display
temp <- sind.ilr %*% t(V)
dimnames(temp)[[2]] <- dimnames(sind.mat2open)[[2]]
prmatrix(temp)

## Clean-up
rm(sind.ilr)
rm(V)
rm(temp)
```

remove.na

Remove and Count NAs

Description

Function to remove rows containing NAs from a data vector or matrix. Also counts the number of rows remaining, the number of rows deleted, and in the case of a matrix the number of columns. The results are returned in a list for subsequent processing in the calling function.

Usage

```
remove.na(xx, iftell = TRUE)
```

Arguments

`xx` name of the vector or matrix to be processed.
`iftell` if `iftell = TRUE`, the default, the number of removed records is displayed.

Details

This function is called by many of the procedures in the 'rgr' package. If one or more NAs are found the user is informed of how many. In general a data frame will have been cleared of any <values represented by negative values or zeros prior to executing the procedure calling this function, see [ltdl.fix.df](#), or [ltdl.fix](#) if a single vector is being processed.

Value

`x` a data vector or matrix containing the elements in the vector or rows of the matrix `xx` without NAs.
`n` the length of `x`.
`m` the number of columns in the matrix `xx`, if `xx` is a vector the value 1 is returned.
`nna` the number of rows removed from `xx`.

Note

The `iftell` 'switch' is used to suppress the display of the NA count in some summary statistics tables as the information is included in the table.

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [where.na](#)

Examples

```
## remove NAs
xx <- c(15, 39, 18, 16, NA, 53)
temp.x <- remove.na(xx)
x <- temp.x$x[1:temp.x$n]

## to recover the other values returned
n <- temp.x$n
m <- temp.x$m
nna <- temp.x$nna

## to remove NA replacing a -9999 in kola.o
data(kola.o)
kola.o.fixed <- ltdl.fix.df(kola.o, coded = -9999)
temp.x <- remove.na(kola.o.fixed$pH)
x <- temp.x$x[1:temp.x$n]
```

```
## Clean-up
rm(xx)
rm(temp.x)
rm(x)
rm(n)
rm(m)
rm(nna)
rm(kola.o.fixed)
```

rng

Undertakes a Range Transformation on the Columns of a Matrix

Description

Function to undertake a range transformation on a data matrix in order that each column is scaled zero-one between the minimum and maximum values.

Usage

```
rng(xx)
```

Arguments

xx a n by p matrix to be range transformed.

Value

x a n by p matrix of range-transformed values.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows containing NAs in the data matrix are removed prior to undertaking the transformation.

A range transform may be appropriate for cluster analysis, including 2-d projection displays, applications to ensure all measured variables have equal weight.

Author(s)

Robert G. Garrett

See Also

[remove.na](#)

Examples

```
## Make test data available
data(sind)
sind.mat <- as.matrix(sind[, -c(1:3)])

## Undertake range transform
temp <- rng(sind.mat)
temp

## Clean-up
rm(sind.mat)
rm(temp)
```

shape

An EDA Graphical Summary

Description

Plots a simple four panel graphical distributional summary for a dataset, comprising a histogram, a horizontal Tukey boxplot or box-and-whisker plot (Garrett, 1988), an empirical cumulative distribution function (ECDF), and a cumulative normal percentage probability (CPP) plot. The plots in all four panels will have identical x-axis scaling. Optionally the EDA graphics may be plotted with logarithmic (base 10) scaling.

Usage

```
shape(xx, xlab = deparse(substitute(xx)), log = FALSE,
      xlim = NULL, nclass = NULL, ifbw = FALSE, wend = 0.05,
      ifnright = TRUE, colr = 8, cex = 0.8, ...)
```

Arguments

<code>xx</code>	name of the variable to be plotted.
<code>xlab</code>	by default the character string for <code>xx</code> is used for the x-axis plot titles. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
<code>log</code>	to display the data with logarithmic (x-axis) scaling, set <code>log = TRUE</code> .
<code>xlim</code>	is determined by gx.hist and used to ensure all four panels in this function have the same x-axis scaling. <code>xlim</code> may be defined, see Note below.
<code>nclass</code>	the default procedure for preparing the histogram depends on sample size. Where $N \leq 500$ the Scott (1979) rule is used, and when $N > 500$ the Freedman-Diaconis (1981) rule; both these rules are resistant to the presence of outliers, and usually provide informative histograms. Alternately, the user may define the histogram binning by setting <code>nclass</code> , i.e. <code>nclass = "scott"</code> , <code>nclass = "fd"</code> or <code>nclass = "sturges"</code> ; the latter being designed for normal distributions (Scott, 1992). See Venables and Ripley (2001) for details.

<code>ifbw</code>	the default is to plot a horizontal Tukey boxplot, if a box-and-whisker plot is required set <code>ifbw = TRUE</code> .
<code>wend</code>	if <code>ifbw = TRUE</code> the locations of the whisker-ends have to be defined. By default these are at the 5th and 95th percentiles of the data, setting <code>wend = 0.02</code> plots the whisker ends at the 2nd and 98th percentiles.
<code>colr</code>	by default the histogram and Tukey boxplot, or box-and-whisker plot, are infilled in grey, <code>colr = 8</code> . If no infill is required, set <code>colr = 0</code> . See function display.lty for the range of available colours.
<code>ifnright</code>	controls where the sample size is plotted in the histogram display, by default this in the upper right corner of the plot. If the data distribution is such that the upper left corner would be preferable, set <code>ifnright = FALSE</code> . If <code>ifnright = NULL</code> there will be no display of the sample size.
<code>cex</code>	by default the size of the text sample size, N , is set to 80%, i.e. <code>cex = 0.8</code> , and may be changed if required.
<code>...</code>	further arguments to be passed to methods. For example, the size of the axis scale annotation can be changed by setting <code>cex.axis</code> , the size of the axis titles by setting <code>cex.lab</code> , and the size of the plot title by setting <code>cex.main</code> . For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%. By default individual data points in the ECDF and CPP plots are marked by a plus sign, <code>pch = 3</code> , if a cross or open circle is desired, then set <code>pch = 4</code> or <code>pch = 1</code> , respectively. See display.marks for all available symbols. Adding <code>ifqs = TRUE</code> results in horizontal and vertical dotted lines being plotted at the three central quartiles and their values, respectively, in the ECDF and CPP plots. By default the histogram and 'box' are infilled in grey, <code>colr = 8</code> . If no infill is required, set <code>colr = 0</code> . See display.lty for the range of available colours.

Details

A histogram is displayed upper left, an ECDF is displayed below it (lower left). To the right of the histogram a horizontal Tukey boxplot (default) or box-and-whisker plot (option) is displayed (upper right). In the lower right quadrant a cumulative normal percentage probability (CPP) plot is displayed. The x-axis scaling is identical in all four plots.

In a box-and-whisker plot there are two special cases. When `wend = 0` the whiskers extend to the observed minima and maxima that are not plotted with the plus symbol. When `wend = 0.25` no whiskers or the data minimum and maximum are plotted, only the median and box representing the span of the middle 50 percent of the data are displayed.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plots.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = TRUE`. If the defined limits lie within the observed data range truncated plots will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations in the various panels.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or `xx` may be defined in the function call as, for example, `Cu[Cu < some.value]` which would remove the influence of one or more outliers having values greater than `some.value`. In this case the number of data values displayed will be the number that are `<some.value`.

In some R installations the generation of multi-panel displays and the use of function `eqsplot` from package `MASS` causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering `options(warn = -1)` on the R command line, or that line may be included in a 'first' function prepared by the user that loads the 'rgr' package, etc.

For summary statistics displays to complement the graphics see, [gx.summary1](#), [gx.summary2](#) and [inset](#).

Author(s)

Robert G. Garrett

References

Garrett, R.G., 1988. IDEAS - An Interactive Computer Graphics Tool to Assist the Exploration Geochemist. In Current Research Part F, Geological Survey of Canada Paper 88-1F, pp. 1-13. See pp. 5 for a description of box-and-whisker plots.

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p. See pp. 119 for a description of histogram bin selection computations.

See Also

[gx.hist](#), [bxplot](#), [gx.ecdf](#), [cnpplt](#), [remove.na](#), [display.lty](#), [display.marks](#), [ltdl.fix.df](#), [inset](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display to have a first look at the data and
## decide how best to proceed
shape(Cu)

## Provides a more appropriate initial display and indicates the
## quartiles
shape(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE,
ifqs = TRUE)

## Causes the Friedman-Diaconis rule to be used to select the number of
## histogram bins and changes the ECDF and CPP plotting symbols to a
## cross/x
shape(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE,
nclass = "fd", pch = 4)
```

```
## Replaces the Tukey boxplot with a box-and-whisker plot where the
## whiskers extend to the 10th and 90th percentiles and the minimum
## and maximum observed values are marked with a plus sign.
shape(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE,
ifbw =TRUE, wend = 0.1)

## Detach test data
detach(kola.o)
```

shape.alt

An Alternate EDA Graphical Summary

Description

Plots a simple four panel graphical distributional summary for a data set, comprising a histogram, a cumulative normal percentage probability (CPP) plot, an empirical cumulative distribution function (ECDF), and a log-log concentration-number (C-N) plot for multifractality. Optionally the EDA graphics may be plotted with logarithmic (base 10) scaling, in which case all four plots have identical x-axis scaling.

Usage

```
shape.alt(xx, xlab = deparse(substitute(xx)), log = FALSE, xlim = NULL,
nclass = NULL, ifnright = TRUE, ifrev = FALSE, colr = 8, ...)
```

Arguments

xx	name of the variable to be plotted.
xlab	by default the character string for xx is used for the x-axis plot titles. An alternate title can be displayed with xlab = "text string", see Examples.
log	to display the data with logarithmic (x-axis) scaling, set log = TRUE.
xlim	is determined by gx.hist and used to ensure all four panels in this function have the same x-axis scaling. xlim may be defined, see Note below.
nclass	the default procedure for preparing the histogram depends on sample size. Where $N \leq 500$ the Scott (1979) rule is used, and when $N > 500$ the Freedman-Diaconis (1981) rule; both these rules are resistant to the presence of outliers, and usually provide informative histograms. Alternately, the user may define the histogram binning by setting nclass, i.e. nclass = "scott", nclass = "fd" or nclass = "sturges"; the latter being designed for normal distributions (Scott, 1992). See Venables and Ripley (2001) for details.
ifnright	controls where the sample size is plotted in the histogram display, by default this in the upper right corner of the plot. If the data distribution is such that the upper left corner would be preferable, set ifnright = FALSE. If ifnright = NULL there will be no display of the sample size.

ifrev	by default the empirical C-N function is plotted from highest value to lowest, ifrev = FALSE. As the C-N plot is a log-log display this provides greater detail for the highest values. The direction of accumulation can be key in detecting multifractal patterns, it is usually informative to also prepare a plot with ifrev = TRUE, i.e. accumulation from lowest to highest values. To see a dramatic example of this, run the Examples below.
colr	by default the histogram and Tukey boxplot, or box-and-whisker plot, are infilled in grey, colr = 8. If no infill is required, set colr = 0. See function display.lty for the range of available colours.
...	further arguments to be passed to methods. For example, the size of the axis scale annotation can be changed by setting cex.axis, the size of the axis titles by setting cex.lab, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%. By default individual data points in the ECDF and CPP plots are marked by a plus sign, pch = 3, if a cross or open circle is desired, then set pch = 4 or pch = 1, respectively. See display.marks for all available symbols. Adding ifqs = TRUE results in horizontal and vertical dotted lines being plotted at the three central quartiles and their values, respectively, in the ECDF and CPP plots. By default the histogram is infilled in grey, colr = 8. If no infill is required, set colr = 0. See display.lty for the range of available colours.

Details

A histogram is displayed upper left, and an ECDF is displayed below it (lower left). To the right of the histogram a cumulative normal percentage probability (CPP) plot is displayed. Below it (lower right) a log-log C-N plot is displayed to highlight any multifractality in the data, which will be revealed as 'lines' of data points with different slopes. When log scaling is selected the x-axis scaling is identical in all four plots.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to displaying the plots.

If the default selection for xlim is inappropriate it can be set, e.g., xlim = c(0, 200) or c(2, 200), the latter being appropriate for a logarithmically scaled plot, i.e. log = TRUE. If the defined limits lie within the observed data range truncated plots will be displayed. If this occurs the number of data points omitted is displayed below the total number of observations in the various panels.

If it is desired to prepare a display of data falling within a defined part of the actual data range, then either a data subset can be prepared externally using the appropriate R syntax, or xx may be defined in the function call as, for example, Cu[Cu < some.value] which would remove the influence of one or more outliers having values greater than some.value. In this case the number of data values displayed will be the number that are <some.value.

In some R installations the generation of multi-panel displays and the use of function eqsplot from package MASS causes warning messages related to graphics parameters to be displayed on the current device. These may be suppressed by entering options(warn = -1) on the R command

line, or that line may be included in a ‘first’ function prepared by the user that loads the ‘rgr’ package, etc.

For summary statistics displays to complement the graphics see, [gx.summary1](#), [gx.summary2](#) and [inset](#).

Author(s)

Robert G. Garrett

References

Venables, W.N. and Ripley, B.D., 2001. Modern Applied Statistics with S-Plus, 3rd Edition, Springer, 501 p. See pp. 119 for a description of histogram bin selection computations.

See Also

[gx.hist](#), [cnpplt](#), [gx.ecdf](#), [gx.mf](#), [remove.na](#), [display.lty](#), [display.marks](#), [ltdl.fix.df](#), [inset](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Generates an initial display to have a first look at the data and
## decide how best to proceed
shape.alt(Cu)

## Provides a more appropriate initial display and indicates the
## quartiles
shape.alt(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE,
ifqs = TRUE)

## Causes the C-N plot to be cumulated in reverse order. This will reveal
## any multifractal properties of the data at lower concentrations
shape.alt(Cu, xlab = "Cu (mg/kg) in <2 mm 0-horizon soil", log = TRUE,
ifrev = TRUE)

## Detach test data
detach(kola.o)
```

sind

Sinding-Larsen Norwegian Stream Sediment Test Data Set

Description

A data frame for a small subset of data from a regional geochemical stream sediment survey undertaken by the Norwegian Geological Survey.

Usage

```
data(sind)
```

Format

A data frame with 25 observations on the following 9 variables:

ID an arbitrary ID (numeric vector).

E an Eastings coordinate.

N a Northings coordinate.

Zn zinc (mg/kg) in stream sediment.

Fe iron (%) in stream sediment.

Mn manganese (mg/kg) in stream sediment.

Cd cadmium (mg/kg) in stream sediment.

Cu copper (mg/kg) in stream sediment.

Pb lead (mg/kg) in stream sediment.

Details

These data were used by Howarth and Sinding-Larsen (1983) to demonstrate the use of a number of multivariate statistical analysis techniques. Other authors, e.g., Howarth and Garrett (1986) and Garrett and Grunsky (2001) have also used the data set for demonstration purposes.

Source

Howarth and Sinding-Larson (1983), see below. Spatial coordinates added by digitizing Fig. 6-1 (op. cit.).

References

- Howarth, R.J. and Sinding-Larsen, R., 1983. Multivariate analysis. Chapter 6 of Handbook of Exploration Geochemistry, Vol. 2, Statistics and Data Analysis in Geochemical Prospecting (Ed. R.J. Howarth), Elsevier, pp. 207-289.
- Howarth, R.J. and Garrett, R.G., 1986. The role of computing in applied geochemistry. In Applied Geochemistry in the 1980s (Eds. I. Thornton and R.J. Howarth), Graham and Trotman, London, pp. 163-184.
- Garrett, R.G. and Grunsky, E.G., 2001. Weighted Sums - Knowledge based empirical indices for use in exploration geochemistry. Geochemistry: Exploration, Environment and Analysis, 1(2):135-141.

sind.mat2open

Sinding-Larsen Norwegian Stream Sediment Test Data Set

Description

A matrix for a small subset of data from a regional geochemical stream sediment survey undertaken by the Norwegian Geological Survey. Similar to data set [sind](#) but the ID, Eastings and Northings columns have been removed, and the measurements are all in mg/kg.

Usage

```
data(sind.mat2open)
```

Format

A matrix with 25 observations on the following 6 variables:

Zn zinc (mg/kg) in stream sediment.

Fe iron (mg/kg) in stream sediment.

Mn manganese (mg/kg) in stream sediment.

Cd cadmium (mg/kg) in stream sediment.

Cu copper (mg/kg) in stream sediment.

Pb lead (mg/kg) in stream sediment.

Details

These data were used by Howarth and Sinding-Larsen (1983) to demonstrate the use of a number of multivariate statistical analysis techniques. Other authors, e.g., Howarth and Garrett (1986) and Garrett and Grunsky (2001) have also used the data set for demonstration purposes.

Source

Howarth and Sinding-Larsen (1983), see below. Spatial coordinates added by digitizing Fig. 6-1 (op. cit.).

References

Howarth, R.J. and Sinding-Larsen, R., 1983. Multivariate analysis. Chapter 6 of Handbook of Exploration Geochemistry, Vol. 2, Statistics and Data Analysis in Geochemical Prospecting (Ed. R.J. Howarth), Elsevier, pp. 207-289.

Howarth, R.J. and Garrett, R.G., 1986. The role of computing in applied geochemistry. In Applied Geochemistry in the 1980s (Eds. I. Thornton and R.J. Howarth), Graham and Trotman, London, pp. 163-184.

Garrett, R.G. and Grunsky, E.G., 2001. Weighted Sums - Knowledge based empirical indices for use in exploration geochemistry. Geochemistry: Exploration, Environment and Analysis, 1(2):135-141.

See Also[sind](#)

syms

*Function to Compute the Diameters of Proportional Symbols***Description**

This function computes the diameters of the open circles to be plotted in a map or other display.

Usage

```
syms(z, zrange = c(NA, NA), p = 1)
```

Arguments

z	name of the variable to be plotted for which diameters are to be computed.
zrange	The minimum and maximum values of z to be used as the lower and upper limits, respectively, for the computed symbol diameters. By default the minimum and maximum values of the input data are used.
p	a parameter that controls the rate of change of symbol diameter with changing value. A default of $p = 1$ is provided that results in a linear rate of change. See Details below.

Details

The symbol diameter is computed as a function of the value z to be plotted:

$$\text{diameter} = \text{dmin} + (\text{dmax} - \text{dmin}) * \{(z - \text{zmin})/(\text{zmax} - \text{zmin})\}^p$$

where dmin and dmax are defined as 0.1 and 1 units, so the symbol diameters range over an order of magnitude (and symbol areas over two); zmin and zmax are the observed range of the data, or the range over which the user wants the diameters to be computed; and p is a power defined by the user. The value of $(z - \text{zmin})/(\text{zmax} - \text{zmin})$ is the value of z normalized, 0 - 1, to the range over which the symbol diameters are to be computed. After being raised to the power p, which will result in a number in the range 0 to 1, this value is multiplied by the permissible range of diameters and added to the minimum diameter. This results in a diameter between 0.1 and 1 units that is proportional to the value of z.

A p value of 1 results in a linear rate of change. Values of p less than unity lead to a rapid initial rate of change with increasing value of z which is often suitable for displaying negatively skewed data sets, see the example below. In contrast, values of p greater than unity result in an initial slow rate of change with increasing value of z which is often suitable for displaying positively skewed data sets. Experimentation is usually necessary to obtain a satisfactory visual effect. See [syms.pfunc](#) for a graphic demonstrating the effect of varying the p parameter.

If zmin or zmax are defined this has the effect of setting a minimum or maximum value of z, respectively, beyond which changes in the value of z do not result in changes in symbol diameter. This can be useful in limiting the effect of one or a few extreme outliers while still plotting them, they simply plot at the minimum or maximum symbol size and are not involved in the calculation of the range of z over which the diameter varies.

Value

zdiam the computed diameter of the symbol.

Author(s)

Robert G. Garrett

See Also

[syms.pfunc](#)

Examples

```
## Make test data available
data(kola.o)
attach(kola.o)

## Compute default symbol diameters
circle.diam <- syms(Cu, p = 0.3)
circle.diam

## Compute symbol diameters holding all symbols for values greater
## than 1000 to the same size
circle.diam <- syms(Cu, zrange = c(NA, 1000), p = 0.3)
circle.diam

## Clean-up and detach test data
rm(circle.diam)
detach(kola.o)
```

syms.pfunc

Function to Demonstrate the Effect of Different Values of p

Description

This function displays a plot demonstrating the effect of varying the value of p , for a range of p values from 0.2 to 5, on the 0 to 1 normalized values of a variable in order to compute corresponding circular symbol diameters.

Usage

```
syms.pfunc()
```

Author(s)

Robert G. Garrett

 tbplots

Plot Vertical Tukey Boxplots

Description

Plots a series of vertical Tukey boxplots where the individual boxplots represent the data subdivided by the value of some factor. Optionally the y-axis may be scaled logarithmically (base 10) and the values of the Tukey fences used to identify near and far outliers may also be optionally based on the logarithmically transformed data. A variety of other plot options are available, see Details and Note below.

Usage

```
tbplots(x, by, log = FALSE, logx = FALSE, notch = TRUE, xlab = "",
        ylab = deparse(substitute(x)), ylim = NULL, main = "",
        label = NULL, plot.order = NULL, xpos = NA, width, space = 0.25,
        las = 1, cex = 1, adj = 0.5, add = FALSE, ssl = 1, colr = 8,
        ...)
```

Arguments

x	name of the variable to be plotted.
by	the name of the factor variable to be used to subdivide the data. See Details below for when by is undefined.
log	to display the data with logarithmic (y-axis) scaling, set log = TRUE.
logx	if the position of the Tukey boxplot fences are to be computed on the basis of the log transformed data set logx = TRUE. When logx = TRUE it is ensured that log = TRUE.
notch	determines if the boxplots are to be “notched” such that the notches indicate the 95% confidence intervals for the medians. The default is to notch the boxplots, to suppress the notches set notch = FALSE. See Details below.
xlab	a title for the x-axis, by default none is provided.
ylab	by default the character string for x is used for the y-axis title. An alternate title can be displayed with xlab = “text string”, see Examples.
ylim	only for log = FALSE, defines the limits of the y-axis if the default limits based on the range of the data are unsatisfactory. It can be used to ensure the y-axis scaling in multiple sets of boxplots are the same to facilitate visual comparison.
main	a main title may be added optionally above the display by setting main, e.g., main = “Kola Project, 1995”.
label	by default the character strings defining the factors are used to label the boxplots along the x-axis. Alternate labels can be provided with label = c(“Alt1”, “Alt2”, “Alt3”), see Examples.
plot.order	provides an alternate order for the boxplots. Thus, plot.order = c(2, 1, 3) will plot the 2nd ordered factor in the 1st position, the 1st in the 2nd, and the 3rd in its 3rd ordered position, see Details and Examples below.

xpos	the locations along the x-axis for the individual vertical boxplots to be plotted. By default this is set to NA, which causes default equally spaced positions to be used, i.e. boxplot 1 plots at value 1 on the x-axis, boxplot 2 at value 2, etc., up to boxplot “n” at value “n”. See Details below for defining xpos.
width	the width of the boxes, by default this is set to the minimum distance between all adjacent boxplots times the value of space. With the default values of xpos this results in a minimum difference of 1, and with the default of space = 0.25 the width is computed as 0.25. To specify different widths for all boxplots use, for example, width = c(0.3). See Details below for changing individual boxplot widths.
space	the space between the individual boxplots, by default this is 0.25 x-axis units.
las	controls whether the x-axis labels are written parallel to the x-axis, the default las = 1, or are written down from the x-axis by setting las = 2. See also, Details below.
cex	controls the size of the font used for the factor labels plotted along the x-axis. By default this is 1, however, if the labels are long it is sometimes necessary to use a smaller font, for example cex = 0.8 results in a font 80% of normal size.
adj	controls the justification of the x-axis labels. By default they are centred, adj = 0.5, to left justify them if the labels are written downwards set adj = 0.
add	permits the user to plot additional boxplots into an existing display. It is recommended that this option is left as add = FALSE.
ssll	determines the minimum data subset size for which a subset will be plotted. By default this is set to 1, which leads to only a circle with a median bar being plotted, as the subset size increases additional features of the boxplot are displayed. If ssll results in subset boxplots not being plotted, a gap is left and the factor label is still plotted on the x-axis.
colr	by default the boxes are infilled in grey, colr = 8. If no infill is required, set colr = 0. See display.lty for the range of available colours.
...	further arguments to be passed to methods. For example, the size of the axis titles by setting cex.lab, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Details

There are two ways to execute this function. Firstly by defining `x` and `by`, and secondly by combining the two variables with the `split` function. See the first two examples below. The `split` function can be useful if the factors to use in the boxplot are to be generated at run-time, see the last example below. Note that when the `split` construct is used instead of by the whole `split` statement will be displayed as the default y-axis title. Also note that when using `by` the subsets are listed in the order that the factors are encountered in the data, but when using `split` the subsets are listed alphabetically. In either case they can be re-ordered using `plot.order`, see Examples.

The `width` option can be used to define different widths for the individual boxplots. For example, the widths could be scaled to be proportional to the subset population sizes as some function of the square root ($\text{const} * \sqrt{n}$) or logarithm ($\text{const} * \log_{10}(n)$) of those sizes (n). The constant, `const`, would need to be chosen so that on average the width of the individual boxes would be

approximately 0.25, see Example below. It may be desirable for cosmetic purposes to adjust the positions of the boxes along the x-axis, this can be achieved by specifying `xpos`.

Long subset (factor) names can lead to display problems, changing the `las` parameter from its default of `las = 1` which plots subset labels parallel to the axis to `las = 2`, to plot perpendicular to the axis, can help. It may also help to use `label` and `split` the character string into two lines, e.g., by changing the string "Granodiorite" that was supplied to replace the coded factor variable `GRDR` to "Grano-\ndiorite". If this, or setting `las = 2`, causes a conflict with the x-axis title, if one is needed, the title can be moved down a line by using `xlab = "\nLithological Units"`. In both cases the `\n` forces the following text to be placed on the next lower line.

If there are more than 7 labels (subsets) and no alternate labels are provided `las` is set to 2, otherwise some labels may fail to be displayed.

The notches in the boxplots indicate the 95% confidence intervals for the medians and can extend beyond the upper and lower limits of the boxes indicating the middle 50% of the data when subset population sizes are small. The confidence intervals are estimated using the binomial theorem. It can be argued that for small populations a normal approximation would be better. However, it was decided to remain with a non-parametric estimate despite the fact that the calculation of the Tukey fence values involves normality assumptions.

Note

This function is based on a script shared by Doug Nychka on S-News, April 28, 1992.

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the data vector are removed prior to preparing the boxplots.

For summary statistics displays to complement the graphics see [gx.summary.groups](#) or [framework.summary](#).

Author(s)

Douglas W. Nychka and Robert G. Garrett

See Also

[cat2list](#), [ltdl.fix.df](#)

Examples

```
## Make test data kola.c available
data(kola.c)
attach(kola.c)

## Display a simple Tukey boxplot
tbplots(Cu, by = COUNTRY)
tbplots(split(Cu,COUNTRY))

## Display a more appropriately labelled and scaled boxplot
tbplots(Cu, by = COUNTRY, log = TRUE, logx = TRUE, xlab = "Country",
ylab = "Ni (mg/kg) in <2 mm Kola C-horizon soil")

## Display a west-to-east re-ordered plot using the full country names
```

```

tbplots(split(Cu, COUNTRY), log = TRUE, logx = TRUE,
ylab = "Ni (mg/kg) in <2 mm Kola C-horizon soil",
label = c("Finland", "Norway", "Russia"),
plot.order = c(2, 1, 3))

## Detach test data kola.c
detach(kola.c)

## Make test data kola.o available, setting a -9999, indicating a
## missing pH measurement, to NA
data(kola.o)
kola.o.fixed <- ltdl.fix.df(kola.o, coded = -9999)
attach(kola.o.fixed)

## Display relationship between pH in one pH unit intervals and Cu in
## O-horizon (humus) soil
tbplots(split(Cu, trunc(pH+0.5)), log=TRUE, logx = TRUE,
xlab = "O-horizon soil pH to the nearest pH unit",
ylab = "Cu (mg/kg) in <2 mm Kola O-horizon soil")

## As above, but demonstrating the use of variable box widths and the
## suppression of 95% confidence interval notches. The box widths are
## computed as (Log10(n)+0.1)/5, the 0.1 is added as one subset has a
## population of 1. Note: paste is used in constructing xlab, below,
## as the label is long and overflows the text line length
table(trunc(pH+0.5))
tbplots(split(Cu, trunc(pH+0.5)), log=TRUE, logx = TRUE, notch = FALSE,
xlab = paste("O-horizon soil pH to the nearest pH unit,",
"\nbox widths proportional to Log(subset_size)"),
ylab = "Cu (mg/kg) in <2 mm Kola O-horizon soil",
width = c(0.26, 0.58, 0.24, 0.02))

## Detach test data kola.o.fixed
detach(kola.o.fixed)

```

tbplots.by.var

Plot Vertical Tukey Boxplots for Variables

Description

Plots a series of vertical Tukey boxplots where the individual boxplots represent the data subdivided by variables. Optionally the y-axis may be scaled logarithmically (base 10). A variety of other plot options are available, see Details and Note below.

Usage

```

tbplots.by.var(xmat, log = FALSE, logx = FALSE, notch = FALSE,
xlab = "Measured Variables", ylab = "Reported Values",
main = "", label = NULL, plot.order = NULL, xpos = NA,
las = 1, cex = 1, adj = 0.5, colr = 8, ...)

```

Arguments

xmat	the data matrix or data frame containing the data (variables).
log	to display the data with logarithmic (y-axis) scaling, set log = TRUE.
logx	if the positions of the Tukey boxplot fences are to be computed on the basis of log transformed data set logx = TRUE. When logx = TRUE it is ensured that log = TRUE.
notch	determines if the boxplots are to be “notched” such that the notches indicate the 95% confidence intervals for the medians. The default is not to notch the boxplots, to have notches set notch = TRUE.
xlab	a title for the x-axis, by default xlab = “Measured Variables”.
ylab	a title for the y-axis, by default ylab = “Reported Values”.
main	a main title may be added optionally above the display by setting main, e.g., main = “Kola Project, 1995”.
label	by default the character strings defining the variables are used to label the boxplots along the x-axis. Alternate labels can be provided with label = c(“Alt1”, “Alt2”, “Alt3”), see Examples.
plot.order	provides an alternate order for the boxplots. By default the boxplot are plotted in alphabetical order of the factor variables. Thus, plot.order = c(2, 1, 3) will plot the 2nd alphabetically ordered factor in the 1st position, the 1st in the 2nd, and the 3rd in its alphabetically 3rd ordered position.
xpos	the locations along the x-axis for the individual vertical boxplots to be plotted. By default this is set to NA, which causes default equally spaced positions to be used, i.e. boxplot 1 plots at value 1 on the x-axis, boxplot 2 at value 2, etc., up to boxplot “n” at value “n”. See Details below for defining xpos.
las	controls whether the x-axis labels are written parallel to the x-axis, the default las = 1, or are written down from the x-axis by setting las = 2. See also, Details below.
cex	controls the size of the font used for the factor labels plotted along the x-axis. By default this is 1, however, if the labels are long it is sometimes necessary to use a smaller font, for example cex = 0.8 results in a font 80% of normal size.
adj	controls the justification of the x-axis labels. By default they are centred, adj = 0.5, to left justify them if the labels are written downwards set adj = 0.
colr	by default the boxes are infilled in grey, colr = 8. If no infill is required, set colr = 0. See display.lty for the range of available colours.
...	further arguments to be passed to methods. For example, the size of the axis titles by setting cex.lab, and the size of the plot title by setting cex.main. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Details

There are two ways to provide data to this function. Firstly, if all the variables in a data frame are to be displayed, and there are no factor variables, the data frame name can be entered for xmat.

However, if there are factor variables, or only a subset of the variables are to be displayed, the data are entered via the `cbind` construct, see Examples below.

Long variable names can lead to display problems, changing the `las` parameter from its default of `las = 1` which plots subset labels parallel to the axis to `las = 2`, to plot perpendicular to the axis, can help. It may also help to use `label` and split the character string into two lines, e.g., by changing the string "Specific Conductivity" that was supplied to replace the variable name `SC` to "Specific\nConductivity". If this, or setting `las = 2`, causes a conflict with the x-axis title, if one is needed, the title can be moved down a line by using `xlab = "\nPhysical soil properties"`. In both cases the `\n` forces the following text to be placed on the next lower line.

If there are more than 7 labels (variables) and no alternate labels are provided `las` is set to 2, otherwise some variable names may fail to be displayed.

The notches in the boxplots indicate the 95% confidence intervals for the medians and can extend beyond the upper and lower limits of the boxes indicating the middle 50% of the data when subset population sizes are small. The confidence intervals are estimated using the binomial theorem. It can be argued that for small populations a normal approximation would be better. However, it was decided to remain with a non-parametric estimate despite the fact that the calculation of the Tukey fence values involves normality assumptions.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see `ltdl.fix.df`.

Any NAs in the data vectors are removed prior to preparing the boxplots.

For a summary statistics display to complement the graphics see `gx.summary.mat`.

Author(s)

Robert G. Garrett

See Also

`tbplots`, `var2fact`, `ltdl.fix.df`

Examples

```
## Make test data kola.c available
data(kola.c)
attach(kola.c)

## Display a simple Tukey boxplot for measured variables
tbplots.by.var(cbind(Co,Cu,Ni))

## Display a more appropriately labelled and scaled Tukey boxplot
tbplots.by.var(cbind(Co,Cu,Ni), log = TRUE, logx = TRUE,
ylab = "Concentrations (mg/kg) in <2 mm Kola C-horizon soil")

## Detach test data kola.c
detach(kola.c)
```



```
## Make test data ms.data1 available
data(ms.data1)

## Display variables in a data frame, remembering to omit the
## sample IDs
tbplots.by.var(ms.data1[, -1], log=TRUE, logx = TRUE)
```

thplot1

*Display a Thompson-Howarth Plot of Duplicate Measurements***Description**

Function displays a Thompson-Howarth (1973 & 1978) plot for a set of duplicate measurements to visually inspect them as a part of the QA/QC process. By inputting a target precision the data may be visually checked to determine if they meet that criterion. The user is prompted for the location of the two legend items.

Usage

```
thplot1(x1, x2, xname = "", ifzero = 0.01, xlow = NA, xhih = NA,
        yhih = NA, rsd = 5, ptile = 95, main = "", ...)
```

Arguments

x1	a column vector from a matrix or data frame, x1[1], ..., x1[n].
x2	another column vector from a matrix or data frame, x2[1], ..., x2[n]. x1, x2 must be of identical length, n, where x2 is a duplicate measurement of x1.
xname	by default the character string for x1 is used for the title. An alternate title can be displayed with xlab = "text string", see Examples.
ifzero	as the Thompson-Howarth plot is log-scaled values of zero cannot be displayed, therefore the parameter ifzero has to be specified. A suitable choice is a value one order of magnitude lower than the value of the detection limit. A default value of ifzero = 0.01 units is provided, corresponding to a detection limit of 0.1 units.
xlow	if is desired to produce plots with consistent scaling this may be achieved by defining xlow, xhih and yhih, the ylow value is set equal to ifzero. Enter an appropriate value of xlow to ensure all data are displayed on all plots.
xhih	enter an appropriate value of xhih to ensure all data are displayed on all plots.
yhih	enter an appropriate value of yhih to ensure all data are displayed on all plots.
rsd	to assist in QA/QC inspection a target precision may be defined as a RSD%, a default of rsd = 5 is provided. See comments concerning RSD in Details below.
ptile	defines the confidence interval for a line to be drawn on the plot above which only 100 - ptile% of the points should plot if the defined target RSD is being met. A default of ptile = 95 is provided. The function counts the number of points falling 'out of limits' and reports the probability that this number would have fallen 'out of limits' by chance alone.

main a title may be added optionally above the display, see Example.

... further arguments to be passed to methods. For example, the size of the axis scale annotation can be change by setting `cex.axis`, the size of the axis titles by setting `cex.lab`, and the size of the plot title by setting `cex.main`. For example, if it is required to make the plot title smaller, add `cex.main = 0.9` to reduce the font size by 10%.

Details

This function expects the RSD% as a measure of repeatability (precision), which is more familiar to the current generation of applied geochemists, rather than the precision at the 2 Standard Deviation level. The necessary calculations to conform with the Thompson and Howarth procedure are made internally.

Duplicate pairs containing any NAs are omitted from the calculations.

If the data are as a single concatenated vector from a matrix or data frame as `x[1]`, ..., `x[n]` followed by `x[n+1]`, ..., `x[2n]`, or alternated as `x[1]` and `x[2]` being a pair through to `x[2*i+1]` and `x[2*i+2]`, for the `i` in `1:n` duplicate pairs use function [thplot2](#).

The user is prompted for the location of the two legend items added to the plot, the number of duplicate pairs, and whether or not the duplicates have met the RSD% criterion. In both instances the user is prompted for the location of left end of the text line, or the top left corner of the text block. The probability that the plotted data have met the RSD criterion is displayed in the text block; if the probability exceeds 0.9999 it is displayed as 0.9999.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Duplicate pairs `x1`, `x2` containing any NAs are omitted from the calculations.

This script was published by Garrett and Grunsky (2003)

Author(s)

Robert G. Garrett

References

Garrett, R.G. & Grunsky, E.C., 2003. S and R functions to display Thompson-Howarth plots. *Computers & Geosciences*, 29(2):239-242.

Stanley, C.R., 2003. THPLOT.M: A MATLAB function to implement generalized Thompson-Howarth error analysis using replicate data. *Computers & Geosciences*, 29(2):225-237.

Thompson, M. and Howarth, R.J., 1973. The rapid estimation and control of precision by duplicate determinations. *The Analyst*, 98(1164):153-160.

Thompson, M. and Howarth, R.J., 1978. A new approach to the estimation of analytical precision. *Journal of Geochemical Exploration*, 9(1):23-30.

See Also

[thplot2](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## NOTE: the examples below are commented out as thplot1 makes a
## call to the locator function that fails when the examples are run
## during package checking and building

## Make the Stanley (2003) test data available
##data(ms.data1)
##attach(ms.data1)

## Display the default plot
##thplot1(MS.1, MS.2, xname = "Magnetic Susceptibility",
## main = "Stanley (2003) Test Data")

## Display a Thompson-Howarth plot for a RSD of 7.5% and a draw the limit
## for a confidence interval of 90%
##thplot1(MS.1, MS.2, xname = "Magnetic Susceptibility", rsd = 7.5,
## ptilt = 90, main = "Stanley (2003) Test Data")

## Detach test data
##detach(ms.data1)
```

thplot2

Display a Thompson-Howarth Plot of Duplicate Measurements, Alternate Input

Description

Function to prepare data stored in alternate forms from that expected by function [thplot1](#) for its use. For further details see 'x' in Arguments below. The user is prompted for the location of the two legend items.

Usage

```
thplot2(x, xname = deparse(substitute(x)), ifzero = 0.01,
xlow = NA, xhih = NA, yhih = NA, rsd = 5, ptilt = 95, main = "",
ifalt = FALSE, ...)
```

Arguments

x a column vector from a matrix or data frame, $x[1]$, ..., $x[2*n]$. The default is that the first n members of the vector are the first measurements and the second n members are the duplicate measurements. If the measurements alternate, i.e. duplicate pair 1 measurement 1 followed by measurement 2, etc., set `ifalt = TRUE`.

xname	by default the character string for x is used for the title. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
ifzero	as the Thompson-Howarth plot is log-scaled values of zero cannot be displayed, therefore the parameter <code>ifzero</code> has to be specified. A suitable choice is a value one order of magnitude lower than the value of the detection limit. A default value of <code>ifzero = 0.01</code> units is provided, corresponding to a detection limit of 0.1 units.
xlow	if is desired to produce plots with consistent scaling this may be achieved by defining <code>xlow</code> , <code>xhih</code> and <code>yhih</code> , the <code>ylow</code> , the <code>ylow</code> value is set equal to <code>ifzero</code> . Enter an appropriate value of <code>xlow</code> to ensure all data are displayed on all plots.
xhih	enter an appropriate value of <code>xhih</code> to ensure all data are displayed on all plots.
yhih	enter an appropriate value of <code>yhih</code> to ensure all data are displayed on all plots.
rsd	to assist in QA/QC inspection a target precision may be defined as a RSD%, a default of <code>rsd = 5</code> is provided. See comments concerning RSD in details below.
ptile	defines the confidence interval for a line to be drawn on the plot above which only <code>100 - ptile%</code> of the points should plot if the defined target RSD is being met. A default of <code>ptile = 95</code> is provided. The function counts the number of points falling 'out of limits' and reports the probability that this number would have fallen 'out of limits' by chance alone.
main	a title may be added optionally above the display, see Example.
ifalt	set <code>ifalt = TRUE</code> to accommodate alternating sets of paired observations.
...	further arguments to be passed to methods. For example, the size of the axis scale annotation can be change by setting <code>cex.axis</code> , the size of the axis titles by setting <code>cex.lab</code> , and the size of the plot title by setting <code>cex.main</code> . For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

This function expects the RSD% as a measure of repeatability (precision), which is more familiar to the current generation of applied geochemists, rather than the precision at the 2 Standard Deviation level. The necessary calculations to conform with the Thompson and Howarth procedure are made internally.

For further details see [thplot1](#).

Duplicate pairs containing any NAs are omitted from the calculations.

If the data are as n duplicate pairs, `x1` and `x2`, use function [thplot1](#).

The user is prompted for the location of the two legend items added to the plot, the number of duplicate pairs, and whether or not the duplicates have met the RSD% criterion. In both instances the user is prompted for the location of the left end of the text line, or the top left corner of the text block.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Author(s)

Robert G. Garrett

See Also[thplot1, ltdl.fix.df](#)**Examples**

```
## NOTE: the examples below are commented out as thplot1 makes a
## call to the locator function that fails when the examples are run
## during package checking and building

## Make test data available
##data(ms.data2)
##attach(ms.data2)

## Display the default plot
##thplot2(MS, xname = "Magnetic Susceptibility",
## main = "Stanley (2003) Test Data")

## Detach test data
##detach(ms.data2)

## Make test data available
##data(ms.data3)
##attach(ms.data3)

## Display a Thompson-Howarth plot for a RSD of 7.5% and draw
## the limit for a confidence interval of 90%
##thplot2(MS, xname = "Magnetic Susceptibility", rsd = 7.5, ptile = 90,
## main = "Stanley (2003) Test Data", ifalt = TRUE)

## Detach test data
##detach(ms.data3)
```

triples.test1*North American Geochemical Soil Landscapes Project QA/QC data*

Description

A data frame of QA/QC triplicates samples from the Maritimes 2007 North American Soil Geochemical Landscapes survey. The analyses are for <2 mm unmilled C-horizon soil samples, and the determinations were made by ICP-OES or -MS following an Aqua Regia digestion.

Usage

```
data(triples.test1)
```

Format

A data frame with 27 observations for the following 3 variables:

ID the NAmSGLs unique site identifier.

RS the Replicate Status code.

Ba_ppm the barium determinations, mg/kg.

Details

The Replicate Status code indicates the ‘position’ of the geochemical sample in the QA/QC structure. RS = 8 indicates analytical duplicate, RS = 2 indicates the field duplicate, and RS = 1 indicates a routine regional coverage site that was ‘duplicated’. All other routine regional coverage sites are coded RS = 0. The analytical duplicate may be split from either of the two field sites, this information being in the project database. Thus the ‘triples’ may occur in the sequence ‘8, 2, 1’ or ‘8, 1, 2’. For `gx.triples.aov` to estimate the variance components correctly the geochemical samples must occur in the file in correct sequence.

Source

The Geological Survey of Canada, see Open File 6433, from which this QA/QC subset for barium (Ba) was extracted.

References

Friske, P.W.B., Ford, K.L. and McNeil, R.J., 2012. Soil Geochemical, Mineralogical, Radon and Radiometric Data from the 2007 North American Soil Geochemical Landscapes Project in New Brunswick, Nova Scotia and Prince Edward Island. Geological Survey of Canada, Open File 6433.

triples.test2

North American Geochemical Soil Landscapes Project QA/QC data

Description

A data frame of regional and field duplicate samples from the Maritimes 2007 North American Soil Geochemical Landscapes survey. The analyses are for <2 mm unmilled C-horizon soil samples, and the determinations were made by ICP-OES or -MS following an Aqua Regia digestion.

Usage

```
data(triples.test2)
```

Format

A data frame with 186 observations for the following 3 variables:

ID the NAmSGLs unique site identifier.

RS the Replicate Status code.

Ba_ppm the barium determinations, mg/kg

Details

The Replicate Status code indicates the ‘position’ of the geochemical sample in the QA/QC structure. RS = 2 indicates the field duplicate, and RS = 1 indicates a routine regional coverage site that was ‘duplicated’. All other routine regional coverage sites are coded RS = 0.

Source

The Geological Survey of Canada, see Open File 6433, from which this data subset for barium (Ba) was extracted.

References

Friske, P.W.B., Ford, K.L. and McNeil, R.J., 2012. Soil Geochemical, Mineralogical, Radon and Radiometric Data from the 2007 North American Soil Geochemical Landscapes Project in New Brunswick, Nova Scotia and Prince Edward Island. Geological Survey of Canada, Open File 6433.

var2fact

Rearranges Data for Variables as Factors

Description

Rearranges data from a matrix or data frame into a matrix where data are tagged by their variables names as factors. Used to concatenate data for display with functions [tbplots.by.var](#) and [bwplots.by.var](#).

Usage

```
var2fact(xmat)
```

Arguments

xmat name of the n by p data matrix or data frame to be processed.

Details

If the data for only some of the variables available in an attached matrix or data frame are to be processed use the cbind construct. Thus, temp.mat <- cbind(vname1, vname3, vname6, vname8).

Value

xx a n * p by 2 matrix where each of the n * p rows contains a value that is paired with its variable name as a factor, see Note below.

Note

The p variables for n cases results in a n * p by 2 matrix, where [1:n, 1] contains the variable name for value[1] and [1:n, 2] contains the values for the n rows in the first column of xmat. Then rows [n+1:2n, 1] contain the variable name for value[2] and [n+1:2n, 2] contain the values for n rows in the second column, and so on.

Author(s)

Robert G. Garrett

Examples

```
## Display, convert data frame and display the result
data(ms.data1)
ms.data1
temp <- var2fact(ms.data1)
temp

## Clean-up
rm(temp)
```

where.na

Identify Vector Elements or Data Frame/Matrix Rows with NAs

Description

Function to display the positions of elements in a vector containing NAs, or the numbers of rows in a data frame or matrix containing one or more NAs. The function can also be used to remove NAs.

Usage

```
where.na(x)
```

Arguments

x name of the vector or matrix/data frame to be processed.

Value

whichna a vector containing the indices of the positions in x containing NAs.

Note

This function is based on the S-Plus function `which.na` and is useful in finding the location of NAs in a data set. While `remove.na` removes NAs it does not identify their positions. A vector is returned that can be used to remove NAs, see example below.

Remember, a matrix is also a vector with the columns occurring sequentially.

Author(s)

S-Plus team and Robert G. Garrett

See Also

[remove.na](#)

Examples

```
## Identify rows with NAs
xx <- c(15, 39, 18, 16, NA, 53)
where.na(xx)

## To use where.na to remove NAs, method 1
xx
temp <- where.na(xx)
temp
xxx <- xx[-temp]
xxx

## To use where.na to remove NAs, method 2
xx
xxx <- xx[-where.na(xx)]
xxx

## Clean-up
rm(xx)
rm(xxx)
rm(temp)
```

wtd.sums

Function to Compute Weighted Sums

Description

Computes weighted sums for a user selected group of variables (Garrett et al., 1980; Garrett and Grunsky, 2001). The user must provide the relative importances of the the variables contributing to the weighted sums. By default the median and MAD are estimated as measures of location and spread for the data. These may be replaced with alternate estimates if the user wishes, see Details below. An object is created containing all the estimated parameters and the weighted sums for later reference and use.

Usage

```
wtd.sums(xx, ri, xloc = NULL, xspread = NULL)
```

Arguments

xx	name of the n by p matrix containing the data.
ri	a vector of the relative weights for the p variables, negative weights are permissible to indicate that high levels of the variable should have a negative impact on the weighted sums.
xloc	the default procedure is to use the computed medians of the input variables. Alternately, a vector of p estimates of location may be provided.
xspread	the default procedure is to use the computed MADs of the input variables. Alternately, a vector of p estimates of spread may be provided.

Details

If the data for only some of the variables available in an attached matrix or data frame are to be processed use the `cbind` construct. Thus, `temp.mat <- cbind(vname1, vname3, vname6, vname8)`, or the `cbind` may be used directly, see Example below.

Value

The following are returned as an object to be saved for further use:

<code>input</code>	the name of the input data set
<code>xloc</code>	the vector of locations used for the computations
<code>xspread</code>	the vector of spreads used for the computations
<code>ri</code>	the vector of relative importances provided by the user
<code>w</code>	the vector of weights computed from the relative importances
<code>a</code>	the vector of coefficients - the normalized weights
<code>ws</code>	the computed weighted sums

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any rows in the data matrix with with NAs are removed prior to computing the weighted sums.

Author(s)

Robert G. Garrett

References

Garrett, R.G. and Grunsky, E.G., 2001. Weighted Sums - Knowledge based empirical indices for use in exploration geochemistry. *Geochemistry: Exploration, Environment and Analysis*, 1(2):135-141.

Garrett, R.G, Kane, V.E. and Zeigler, R.K., 1980. The management and analysis of regional geochemical data. *Journal of Geochemical Exploration*, 13(1/2):115-152.

See Also

[ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Compute weighted sums as in Garrett & Grunsky (2001)
## using medians and interquartile SDs
sind.ws.geea <- wtd.sums(cbind(Zn, Cd, Fe, Mn), ri = c(2, 1, -1, -1),
```

```

xloc = c(48, 0.6, 1.74, 590),
xspread = c(41.5128, 0.44478, 0.882147, 333.585))

## Compute weighted sums using the median and MAD defaults
sind.ws.def <- wtd.sums(cbind(Zn, Cd, Fe, Mn), ri = c(2, 1, -1, -1))

## Plot the two results against one-another, adding a constant,
## 3, to the weighted sums to make them positive and log-scale
## plottable
par(pty="s")
plot(sind.ws.geea$ws+3, sind.ws.def$ws+3, log = "xy",
xlim = c(2, 28), ylim = c(2, 28))
abline(0, 1, lty = 3)
abline(v =3, lty = 3)
abline(h = 3, lty = 3)

## Inspect the default weighted sums, adding a constant, 3, to the
## weighted sums to make them positive and log-scale plottable
shape(sind.ws.def$ws+3, log = TRUE)

## Plot EDA Tukey boxplot based map of default weighted sums
map.eda7(E, N, sind.ws.def$ws)

## Clean-up and detach test data
rm(sind.ws.geea)
rm(sind.ws.def)
par(pty = "m")
detach(sind)

```

xyplot.eda7

*Display a Third Variable in a X-Y Plot using Tukey Boxplot Symbology***Description**

Displays a third variable where the data are represented by symbols using Tukey boxplot-based symbology. Tukey boxplots divide data into 7 groups, the middle 50%, and three lower and higher groupings, see Details below. The computation of the fences used to subdivide the data may be carried out following a logarithmic transformation of the data. The colours of the symbols may be optionally changed. The x-y plot axes may be optionally displayed with logarithmic (base 10) scaling. Optionally a legend may be added to the plot.

Usage

```

xyplot.eda7(xx, yy, zz, sfact = 1, xlim = NULL, ylim = NULL,
log = NULL, logz = FALSE, xlab = deparse(substitute(xx)),
ylab = deparse(substitute(yy)), zlab = deparse(substitute(zz)),
main = "", ifgrey = FALSE, symcolr = NULL, iflgnd = FALSE,
title = deparse(substitute(zz)), cex.lgnd = 0.8, ...)

```

Arguments

<code>xx</code>	name of the x-axis variable.
<code>yy</code>	name of the y-axis variable.
<code>zz</code>	name of the third variable to be plotted.
<code>sfact</code>	controls the absolute size of the plotted symbols, by default <code>sfact = 1</code> . Increasing <code>sfact</code> results in larger symbols.
<code>xlim</code>	user defined limits for the x-axis, see Details below.
<code>ylim</code>	user defined limits for the y-axis, see Details below.
<code>log</code>	if it is required to display the data with logarithmic axis scaling, set <code>log = "x"</code> for a logarithmically scaled x-axis, <code>log = "y"</code> for a logarithmically scaled y-axis, and <code>log = "xy"</code> for both axes logarithmically scaled.
<code>logz</code>	if it is required to undertake the Tukey Boxplot computations after a logarithmic data transform, set <code>logz = TRUE</code> .
<code>xlab</code>	by default the character string for <code>xx</code> is used for the x-axis title. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
<code>ylab</code>	by default the character string for <code>yy</code> is used for the x-axis title. An alternate title can be displayed with <code>ylab = "text string"</code> , see Examples.
<code>zlab</code>	by default the character string for <code>zz</code> is appended to the text string "EDA Tukey Boxplot Based Plot for" for the plot title. An alternate title can be displayed with <code>zlab = "text string"</code> , see Details below.
<code>main</code>	an alternative plot title, see Details below.
<code>ifgrey</code>	set <code>ifgrey = TRUE</code> if a grey-scale plot is required, see Details below.
<code>symcolr</code>	the default is a colour plot and default colours are provided, deeper blues for lower values, green for the middle 50% of the data, and oranges and reds for higher values. A set of alternate symbol colours can be provided by defining <code>symcolr</code> , see Details below.
<code>iflgn</code>	the default is no legend. If a legend is required set <code>iflgn = TRUE</code> , following the plotting of the data the cursor will be activated, locate that at the top left corner of the desired legend position and 'left button' on the pointing device.
<code>title</code>	a short title for the legend, e.g., <code>title = "Zn (mg/kg)"</code> . The default is the character string for <code>zz</code> .
<code>cex.lgn</code>	controls the scaling of the legend box and text, but not the symbols so that they match those on the plot, the default is <code>cex.lgn = 0.8</code> .
<code>...</code>	further arguments to be passed to methods. For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

Tukey boxplots divide data into 7 groups, the middle 50%, and three lower and higher groupings: within the whisker, near outliers and far outliers, respectively. Symbols for values below the first quartile (Q1) are plotted as increasingly larger circles, while symbols for values above the third quartile are plotted as increasingly larger squares, a '+' is used to plot the data falling in the middle 50%. For the higher groupings, the whisker contains values $>Q3$ and $<(Q3 + 1.5 * HW)$, where

HW = (Q3 - Q1), the interquartile range; near outliers lie between (Q3 + 1.5 * HW) and (Q3 + 3 * HW); and far outliers have values >(Q3 + 3 * HW). For the lower groupings the group boundaries, fences, fall similarly spaced below Q1. The computation of the fences used to subdivide the data may be carried out following a logarithmic transformation of the data, set `logz = TRUE`.

A summary table of the values of the symbol intervals, the number of values plotting as each symbol, and symbol shapes, sizes and colours is displayed on the current device.

If `zlab` and `main` are undefined a default plot title is generated by appending the input variable name text string to "EDA Tukey Boxplot-Based Plot for ". If no plot title is required set `zlab = " "`, and if some user defined plot title is required it should be defined in `main`, e.g. `main = "Plot Title Text"`.

If the grey-scale option is chosen the symbols are plotted 100% black for the far outliers, 85% black for the near outliers, 70% black for values within the whiskers, and 60% black for values falling within the middle 50% of the data.

The default colours, `symcolr = c(25, 22, 20, 13, 6, 4, 1)`, are selected from the `rainbow(36)` palette, and alternate colour schemes need to be selected from the same palette. See [display.rainbow](#) for the available colours. It is essential that 7 colours be provided, e.g., `symcolr = c(27, 24, 22, 12, 5, 3, 36)`, if exactly 7 are not provided the default colours will be displayed.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors including NAs are removed prior to displaying the plot.

Author(s)

Robert G. Garrett

See Also

[display.rainbow](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Plot a default Tukey boxplot-based display
xyplot.eda7(Fe, Mn, Zn)

## Plot with logarithmically scaled boxplot fences and more
## appropriate axis scaling and labelling with a user specified title
xyplot.eda7(Fe, Mn, Zn, sfact = 2, log = "y", logz = TRUE,
xlab = "Fe (pct) in stream sediment",
ylab = "Mn (mg/kg) in stream sediment",
main = "Howarth & Sinding-Larsen Test Data\nLog10(Zn) (mg/kg)")
```

```
## Display a grey-scale equivalent of the above plot
xyplot.eda7(Fe, Mn, Zn, sfact = 2, log = "y", logz = TRUE, ifgrey = TRUE,
  xlab = "Fe (pct) in stream sediment",
  ylab = "Mn (mg/kg) in stream sediment",
  main = "Howarth & Sinding-Larsen Test Data\nLog10(Zn) (mg/kg)")

## Plot the same display with an alternate colour scheme
xyplot.eda7(Fe, Mn, Zn, sfact = 2, log = "y", logz = TRUE,
  xlab = "Fe (pct) in stream sediment",
  ylab = "Mn (mg/kg) in stream sediment",
  main = "Howarth & Sinding-Larsen Test Data\nLog10(Zn) (mg/kg)",
  symcolr = c(27, 24, 22, 12, 5, 3, 36))

## Detach test data
detach(sind)
```

xyplot.eda8

*Display a Third Variable in a X-Y Plot as Percentiles***Description**

Displays a third variable on a X-Y plot where the the third variable is represented by symbols indicating within which group defined by the data's 2nd, 5th, 25th, 50th, 75th, 95th and 98th percentiles plotted a data value falls. The colours of the symbols may be optionally changed. The x-y plot axes may be optionally displayed with logarithmic (base 10) scaling. Optionally a legend (two options) may be added to the plot.

Usage

```
xyplot.eda8(xx, yy, zz, sfact = 1, xlim = NULL, ylim = NULL,
  xlab = deparse(substitute(xx)), ylab = deparse(substitute(yy)),
  zlab = deparse(substitute(zz)), main = "", log = NULL,
  ifgrey = FALSE, symcolr = NULL, iflgnd = FALSE, pctile = FALSE,
  title = deparse(substitute(zz)), cex.lgnd = 0.8, ...)
```

Arguments

xx	name of the x-axis variable.
yy	name of the y-axis variable.
zz	name of the third variable to be plotted.
sfact	controls the absolute size of the plotted symbols, by default sfact = 1. Increasing sfact results in larger symbols.
log	if it is required to display the data with logarithmic axis scaling, set log = "x" for a logarithmically scaled x-axis, log = "y" for a logarithmically scaled y-axis, and log = "xy" for both axes logarithmically scaled.
xlim	user defined limits for the x-axis, see Details below.
ylim	user defined limits for the y-axis, see Details below.

xlab	by default the character string for xx is used for the x-axis title. An alternate title can be displayed with <code>xlab = "text string"</code> , see Examples.
ylab	by default the character string for yy is used for the y-axis title. An alternate title can be displayed with <code>ylab = "text string"</code> , see Examples.
zlab	by default the character string for zz is appended to the text string "EDA Percentile Based Plot for" for the plot title. An alternate title can be displayed with <code>zlab = "text string"</code> , see Details below.
main	an alternative plot title, see Details below.
ifgrey	set <code>ifgrey = TRUE</code> if a grey-scale plot is required, see Details below.
symcolr	the default is a colour plot and default colours are provided, deeper blues for lower values, green for the middle 50% of the data, and oranges and reds for higher values. A set of alternate symbol colours can be provided by defining <code>symcolr</code> , see Details below.
iflgnd	the default is no legend. If a legend is required set <code>iflgnd = TRUE</code> , following the plotting of the data the cursor will be activated, locate that at the top left corner of the desired legend position and 'left button' on the pointing device. There are two legends to choose from, see <code>pctile</code> below.
pctile	the default legend displays the range of values each symbol represents. Alternately, the percentiles may be displayed rather than their values by setting <code>pctile = TRUE</code> .
title	a short title for the legend, e.g., <code>title = "Zn (mg/kg)"</code> . The default is the character string for <code>zz</code> .
cex.lgnd	controls the scaling of the legend box and text, but not the symbols so that they match those in the plot, the default is <code>cex.lgnd = 0.8</code> .
...	further arguments to be passed to methods. For example, if it is required to make the plot title smaller, add <code>cex.main = 0.9</code> to reduce the font size by 10%.

Details

The selected percentiles, 2nd, 5th, 25th, 50th, 75th, 95th and 98th, divide the data into 8 groups. Values below the median are represented by increasingly larger deeper blue circles below the 25th percentile (Q1), and values above the 75th percentile (Q3) by increasingly larger orange and red squares. The mid 50% of the data are represented by green symbols, circles for the median (Q2) to Q1, and squares for the median (Q2) to Q3.

A summary table of the values of the symbol intervals, the number of values plotting as each symbol, and symbol shapes, sizes and colours is displayed on the current device.

If `zlab` and `main` are undefined a default plot title is generated by appending the input variable name text string to "EDA Percentile Based Plot for ". If no plot title is required set `zlab = ""`, and if some user defined plot title is required it should be defined in `main`, e.g. `main = "Plot Title Text"`.

If the grey-scale option is chosen the symbols are plotted 100% black for the far outliers, 85% black for the near outliers, 70% black for values within the whiskers, and 60% black for values falling within the middle 50% of the data.

The default colours, `symcolr = c(25, 22, 20, 13, 13, 6, 4, 1)`, are selected from the `rainbow(36)` palette, and alternate colour schemes need to be selected from the same palette.

See [display.rainbow](#) for the available colours. It is essential that 8 colours be provided, e.g., `symcolr = c(27, 24, 22, 12, 12, 5, 3, 36)`, if exactly 8 are not provided the default colours will be displayed.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any data vectors including NAs are removed prior to displaying the plot.

Author(s)

Robert G. Garrett

See Also

[display.rainbow](#), [ltdl.fix.df](#), [remove.na](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Plot a default percentile display
xyplot.eda8(Fe, Mn, Zn)

## Plot with more appropriate axis scaling and labelling
## with a user specified title
xyplot.eda8(Fe, Mn, Zn, sfact = 2.0, log = "y",
  xlab = "Fe (pct) in stream sediment",
  ylab = "Mn (mg/kg) in stream sediment",
  main = "Howarth & Sinding-Larsen Test Data\nZn (mg/kg)")

## Display a grey-scale equivalent of the above plot
xyplot.eda8(Fe, Mn, Zn, sfact = 2, log = "y", ifgrey = TRUE,
  xlab = "Fe (pct) in stream sediment",
  ylab = "Mn (mg/kg) in stream sediment",
  main = "Howarth & Sinding-Larsen Test Data\nZn (mg/kg)")

## Plot the same display with an alternate colour scheme
xyplot.eda8(Fe, Mn, Zn, sfact = 2, log = "y",
  xlab = "Fe (pct) in stream sediment",
  ylab = "Mn (mg/kg) in stream sediment",
  main = "Howarth & Sinding-Larsen Test Data\nZn (mg/kg)",
  symcolr = c(27, 24, 22, 12, 12, 5, 3, 36))

## Detach test data
detach(sind)
```


xyplot.tags

*Display a Plot of Posted Values for a Third Variable***Description**

Displays a x-y plot where the data for a third variable are represented by the ‘written’ values of the data at their x-y position. The x-y plot axes may be optionally displayed with logarithmic (base 10) scaling.

Usage

```
xyplot.tags(xx, yy, tag, log = NULL, xlim = NULL, ylim = NULL,
  xlab = deparse(substitute(xx)), ylab = deparse(substitute(yy)),
  taglab = deparse(substitute(tag)), main = "", ...)
```

Arguments

xx	name of the x-axis variable.
yy	name of the y-axis variable.
tag	name of the third variable to be displayed.
log	if it is required to display the data with logarithmic axis scaling, set log = "x" for a logarithmically scaled x-axis, log = "y" for a logarithmically scaled y-axis, and log = "xy" for both axes logarithmically scaled.
xlim	user defined limits for the x-axis, see Details below.
ylim	user defined limits for the y-axis, see Details below.
xlab	by default the character string for xx is used for the x-axis title. An alternate title can be displayed with xlab = "text string", see Examples.
ylab	by default the character string for yy is used for the y-axis title. An alternate title can be displayed with ylab = "text string", see Examples.
taglab	text to be inserted in the plot title, by default deparse(substitute(tag)) is used. See Details below.
main	an alternative plot title, see Details below.
...	further arguments to be passed to methods. For example, if smaller plotting characters are required, specify cex = 0.8; or if some colour other than black is required for the plotting characters, specify col = 2 to obtain red (see display.lty for the default colour palette). If it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Details

If taglab and main are undefined a default a plot title is generated by appending the input variable name text string to "Plot of Values for ". If no plot title is required set taglab = " ", or if an alternative to the variable name taglab is required it may be specified, taglab = "Alternative". If some

user defined plot title is required it should be defined in main, e.g., main = "Plot Title Text", in which instance taglab is ignored.

If the default selection for xlim is inappropriate it can be set, e.g., xlim = c(0, 200) or c(2, 200), the latter being appropriate for a logarithmically scaled plot, i.e. log = "x". If the defined limits lie within the observed data range a truncated plot will be displayed. The same procedure applies to setting ylim.

If a plot of sample numbers, 'IDs', is required and they are not explicitly in the data frame, a plot of data frame row numbers may be displayed by specifying dimnames(dfname)[[1]] as the value of tags.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

Any NAs in the x-y coordinate vectors are removed prior to displaying the plot, thus those 'data' are not plotted. However, any NAs in the third variable to be plotted are replaced with a '+' sign to indicate data for the third variable are 'missing'.

Author(s)

Robert G. Garrett

See Also

[ltdl.fix.df](#), [remove.na](#), [display.lty](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Plot the sample site IDs in the x-y space
xyplot.tags(Fe, Mn, ID)

## Plot the data frame row numbers in the x-y space and appropriately
## scale the y-axis
xyplot.tags(Fe, Mn, dimnames(sind)[[1]], log = "y")

## Plot the values for zinc (Zn) in smaller red text in the x-y
## space, providing more appropriate axis scaling and labelling,
## and adding a user specified title
xyplot.tags(Fe, Mn, Zn, log = "y", xlab = "Fe (pct) in stream sediment",
ylab = "Mn (mg/kg) in stream sediment",
main = "Howarth & Sinding-Larsen Test Data\nZn (mg/kg)", cex = 0.8, col = 2)

## Detach test data
detach(sind)
```

Description

Displays a third variable where the data are represented by open circles whose diameters are proportional to the value of the data at their x-y locations. The rate of change of symbol diameter with value and the absolute size of the symbols are defined by the user. The x-y plot axes may be optionally displayed with logarithmic (base 10) scaling. Optionally a legend may be displayed on the plot.

Usage

```
xyplot.z(xx, yy, zz, p = 0.5, sfact = 2.5, zmin = NA, zmax = NA, log = NULL,
xlim = NULL, ylim = NULL, xlab = deparse(substitute(xx)),
ylab = deparse(substitute(yy)), zlab = deparse(substitute(zz)),
main = "", col = 1, iflgnd = FALSE, title = deparse(substitute(zz)),
cex.lgnd = 0.8, ifparams = FALSE, cex.params = 0.8, ...)
```

Arguments

xx	name of the x-axis variable.
yy	name of the y-axis variable.
zz	name of the third variable, z, to be plotted.
p	a parameter that controls the rate of change of symbol diameter with changing value. A default of $p = 0.5$ is provided. See Details below.
sfact	controls the absolute size of the plotted symbols, by default $\text{sfact} = 2.5$. Increasing sfact results in larger symbols.
zmin	a value below which all symbols will be plotted at the same minimum size. By default $\text{zmin} = \text{NA}$ which results in the minimum value of the variable defining the minimum symbol size. See Details below.
zmax	a value above which all symbols will be plotted at the same maximum size. By default $\text{zmax} = \text{NA}$ which results in the maximum value of the variable defining the maximum symbol size. See Details below.
log	if it is required to display the data with logarithmic axis scaling, set $\text{log} = "x"$ for a logarithmically scaled x-axis, $\text{log} = "y"$ for a logarithmically scaled y-axis, and $\text{log} = "xy"$ for both axes logarithmically scaled.
xlim	user defined limits for the x-axis, see Details below.
ylim	user defined limits for the y-axis, see Details below.
xlab	by default the character string for xx is used for the x-axis title. An alternate title can be displayed with $\text{xlab} = "text\ string"$, see Examples.
ylab	by default the character string for yy is used for the y-axis title. An alternate title can be displayed with $\text{ylab} = "text\ string"$, see Examples.

zlab	by default, zlab = deparse(substitute(z)), a plot title is generated by appending the input variable name text string to "Proportional Symbol Plot for ". Alternative titles may be generated, see Details below.
main	an alternative plot title, see Details below.
col	the colour of the symbols, the default is black, col = 1. This may be changed if required, see display.lty for the default colour palette. For example, col = 2 will cause the symbols to be plotted in red.
iflgn	the default is no legend. If a legend is required set iflgn = TRUE, following the plotting of the data the cursor will be activated, locate that at the top left corner of the desired legend position and 'left button' on the pointing device. See Notes below.
title	a short title for the legend, e.g., title = "Zn (mg/kg)". The default is the character string for z.
cex.lgn	controls the scaling of the legend box and text, but not the symbols so that they match those on the plot, the default is cex.lgn = 0.8.
ifparams	if ifparams = TRUE on completion of plotting and after the legend has been plotted, if requested, the cursor is activated, locate that at the top left corner of the desired text position and 'left button' on the pointing device. This text comprises three lines: the values of p to three significant figures and sfact; the maximum value of z to 3 significant figures and zmax; and the minimum value of z to 3 significant figures and zmin. The default is no text display.
cex.params	controls the size of the text for the parameters display, defaults to cex.params = 0.8.
...	further arguments to be passed to methods. For example, if it is required to make the plot title smaller, add cex.main = 0.9 to reduce the font size by 10%.

Details

The symbol diameter is computed as a function of the value z to be plotted:

$$\text{diameter} = \text{dmin} + (\text{dmax} - \text{dmin}) * \{(z - \text{zmin})/(\text{zmax} - \text{zmin})\}^p$$

where dmin and dmax are defined as 0.1 and 1 units, so the symbol diameters range over an order of magnitude (and symbol areas over two); zmin and zmax are the observed range of the data, or the range over which the user wants the diameters to be computed; and p is a power defined by the user. The value of $(z - \text{zmin})/(\text{zmax} - \text{zmin})$ is the value of z normalized, 0 - 1, to the range over which the symbol diameters are to be computed. After being raised to the power p, which will result in a number in the range 0 to 1, this value is multiplied by the permissible range of diameters and added to the minimum diameter. This results in a diameter between 0.1 and 1 units that is proportional to the value of z.

A p value of 1 results in a linear rate of change. Values of p less than unity lead to a rapid initial rate of change with increasing value of z which is often suitable for displaying positively skewed data sets, see the example below. In contrast, values of p greater than unity result in an initial slow rate of change with increasing value of z which is often suitable for displaying negatively skewed data sets. To obtain a logarithmic scaling set $p = 0.3679$, i.e. $1/e$, and a value of $p = 0.5$ is equivalent to a square root transformation, the default, that is appropriate for a linear-area relationship. These values of p are equivalent to plotting $\log(x)$ or $\log_{10}(x)$ and \sqrt{x} , respectively. Experimentation is often necessary to obtain a satisfactory visual effect. See [syms.pfunc](#) for a graphic demonstrating the effect of varying the p parameter.

If `zmin` or `zmax` are defined this has the effect of setting a minimum or maximum value of `z`, respectively, beyond which changes in the value of `z` do not result in changes in symbol diameter. This can be useful in limiting the effect of one, or a few, extreme outlier(s) while still plotting them, they simply plot at the minimum or maximum symbol size and are not involved in the calculation of the range of `z` over which the symbol diameters vary. **Note:** If the variable `z` includes a transform, e.g., `log10(z)`, the values of `zmin` and/or `zmax` must be in those transform units.

If `zlab` and `main` are undefined a default plot title is generated by appending the input variable name text string to "Proportional Symbol Plot for ". If no plot title is required set `zlab = ""`, and if some user defined plot title is required it should be defined in `main`, e.g. `main = "Plot Title Text"`.

If the default selection for `xlim` is inappropriate it can be set, e.g., `xlim = c(0, 200)` or `c(2, 200)`, the latter being appropriate for a logarithmically scaled plot, i.e. `log = "x"`. If the defined limits lie within the observed data range a truncated plot will be displayed. The same procedure applies to setting `ylim`.

Note

Any less than detection limit values represented by negative values, or zeros or other numeric codes representing blanks in the data, must be removed prior to executing this function, see [ltdl.fix.df](#).

The legend consists of five proportional symbols and their corresponding `z` values: `zmin`; the three quartiles; and `zmax`. If `zmin` and `zmax` have been user defined it is over their range that the symbol sizes are computed and displayed. When defining `zmin` and/or `zmax` it is useful to set `ifparams = TRUE` as a reminder, whilst developing the required display.

Any data vectors containing NAs are removed prior to displaying the plot.

Author(s)

Robert G. Garrett

See Also

[syms](#), [syms.pfunc](#), [ltdl.fix.df](#), [remove.na](#), [display.lty](#)

Examples

```
## Make test data available
data(sind)
attach(sind)

## Display a default symbol plot, p = 0.5 and sfact = 2.5
xyplot.z(Fe, Mn, Zn)

## Plot with logarithmically scaled symbols and more appropriately
## labelled axes
xyplot.z(Fe, Mn, log10(Zn), p = 1, log = "y",
xlab = "Fe (pct) in stream sediment",
ylab = "Mn (mg/kg) in stream sediment")
```

```
## Plot with differently scaled symbols in red and more appropriate
## scaling and labelling with a user specified title
xyplot.z(Fe, Mn, Zn, p = 0.3, sfact = 2.0, log = "y",
xlab = "Fe (pct) in stream sediment",
ylab = "Mn (mg/kg) in stream sediment",
main = "Howarth & Sinding-Larsen Test Data\nZn (mg/kg)", symcolr = 2)

## Plot as above but where outliers above a value of 1000 displayed
## with the same symbol
xyplot.z(Fe, Mn, Zn, p = 0.3, sfact = 2.0, zmax = 1000, log = "y",
xlab = "Fe (pct) in stream sediment",
ylab = "Mn (mg/kg) in stream sediment",
main = "Howarth & Sinding-Larsen Test Data\nZn (mg/kg)", symcolr = 2)

## Detach test data
detach(sind)
```

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