Package ‘analyzer’

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anderson.test

Description

anderson.test performs Anderson-Darling test

Usage

anderson.test(x)

Arguments

x a numeric vector. Length must be greater than 7. Missing values are allowed.

Details

Performs the Anderson-Darling test for the composite hypothesis of normality, see e.g. Thode (2002, Sec. 5.1.4).

Value

A list with following elements:

statistic the value of Anderson-Darling test statistic
p.value p-value of the test
method Test name
data.name Vector name
See Also

norm_test_fun

Examples

anderson.test(mtcars$mpg)

association

Find association between variables

Description

association finds association among all the variables in the data.

Usage

association(
  tb,
  categorical = NULL,
  method1 = c("auto", "pearson", "kendall", "spearman"),
  method3 = c("auto", "parametric", "non-parametric"),
  methodMats = NULL,
  use = "everything",
  normality_test_method = c("ks", "anderson", "shapiro"),
  normality_test_pval = 0.05,
  ...
)

Arguments

  tabular data
categorical a vector specifying the names of categorical (character, factor) columns
method1 method for association between continuous-continuous variables. values can be
  "auto", "pearson", "kendall", "spearman". See details for more information.
method3 method for association between continuous-categorical variables. Values can be
  "auto", "parametric", "non-parametric". See details of CQassociation
  for more information. Parametric does t-test while non-parametric does 'Mann-
  Whitney' test.
methodMats This parameter can be used to define the methods for calculating correlation and
  association at variables pair level. The input is a square data.frame of dimension
  number of columns in tb. The row names and column names of methodMats
  are the column names of tb. The values in the data.frame can be:

  between continuous-continuous variables from parameter method1 - "auto",
    "pearson", "kendall", "spearman"
between continuous-categorical variables from parameter method3 - "auto", "parametric", "non-parametric"

between categorical-categorical variables can be anything

Default is NULL. In that case the method used for calculating correlation and association will be the inputs from parameters.

This parameter can also take some other values. See example for more details. But its advisable to use like mentioned above.

use

an optional character string giving a method for computing association in the presence of missing values. This must be (complete or an abbreviation of) one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs". If use is "everything", NAs will propagate conceptually, i.e., a resulting value will be NA whenever one of its contributing observations is NA. If use is "all.obs", then the presence of missing observations will produce an error. If use is "complete.obs" then missing values are handled by case wise deletion (and if there are no complete cases, that gives an error). "na.or.complete" is the same unless there are no complete cases, that gives NA.

normality_test_method

method for normality test for a variable. Values can be shapiro for Shapiro-Wilk test or 'anderson' for 'Anderson-Darling' test of normality or ks for 'Kolmogorov-Smirnov'

normality_test_pval

significance level for normality tests. Default is 0.05

... other parameters passed to cor, CCassociation, CQassociation and QQassociation

Details

This function calculates association value in three categories -

- between continuous variables (using CCassociation function)
- between categorical variables (using QQassociation function)
- between continuous and categorical variables (using CQassociation function)

For more details, look at the individual documentation of CCassociation, QQassociation, CQassociation

Value

A list of three tables:

continuous_corr correlation among all the continuous variables
continuous_pvalue Table containing p-value for the correlation test

categorical_cramers Cramer’s V value among all the categorical variables
categorical_pvalue Chi Sq test p-value

continuous_categorical association value among continuous and categorical variables

method_used A data.frame showing the method used for all pairs of variables
CCassocation

See Also

CCassocation for Correlation between Continuous variables, QQassocation for Association between Categorical variables, CQassocation for Association between Continuous-Categorical variables

Examples

```r
tb <- mtcars
tb$cyl <- as.factor(tb$cyl)
tb$vs <- as.factor(tb$vs)
out <- association(tb, categorical = c("cyl", "vs"))

# To use the methodMats parameter, create a matrix like this
methodMats <- out$method_used

# the values can be changed as per requirement
# NOTE: in addition to the values from parameters method1 and method3,
# the values in methodMats can also be the values returned by
# association function. But its advisable to use the options from
# method1 and method3 arguments
methodMats["mpg", "disp"] <- methodMats["disp", "mpg"] <- "spearman"
out <- association(tb, categorical = c("cyl", "vs"), methodMats = methodMats)
rm(tb)
```

---

CCassocation

**Association (Correlation) between Continuous (numeric) Variables**

Description

CCassocation finds correlation between all the variables in data with only numeric columns

Usage

```r
CCassocation(
  numtb,
  use = "everything",
  normality_test_method,
  normality_test_pval,
  method1 = c("auto", "pearson", "kendall", "spearman"),
  methodMat1 = NULL,
  methods_used
)
```

Arguments

- `numtb` a data frame with all the numerical columns. This should have at least two columns
use an optional character string giving a method for computing association in the presence of missing values. This must be (complete or an abbreviation of) one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs". If use is "everything", NAs will propagate conceptually, i.e., a resulting value will be NA whenever one of its contributing observations is NA. If use is "all.obs", then the presence of missing observations will produce an error. If use is "complete.obs" then missing values are handled by case wise deletion (and if there are no complete cases, that gives an error). "na.or.complete" is the same unless there are no complete cases, that gives NA

normality_test_method method for normality test for a variable. Values can be shapiro for Shapiro-Wilk test or 'anderson' for 'Anderson-Darling' test of normality or ks for 'Kolmogorov-Smirnov'  

normality_test_pval significance level for normality tests. Default is 0.05  

method1 method for association between continuous-continuous variables. values can be "auto", "pearson", "kendall", "spearman". See details for more information. 

methodMat1 method dataframe like methodMats from the function association 

methods_used a square data.frame which will store the type of association used between the variables. Dimension will be number of variables * number of variables. 

Details 

This function calls cor function to calculate the correlation values. The difference is that this doesn’t take method as parameter, instead it decides the methods itself using normality tests. If the variables satisfy the assumption of Pearson correlation, then pearson correlation is calculated. Otherwise Spearman is calculated. To learn more, check the cor 

Value 

a list of two tables with number of rows and column equal to number of columns in numtb:  

r  Table containing correlation values  

r_pvalue  Table containing p-value for the correlation test 

See Also  

association for association between any type of variables, QQassociation for Association between Categorical variables, CQassociation for Association between Continuous-Categorical variables
consoleBoxplot  

**Description**

consoleBoxplot prints the boxplot on console.

**Usage**

```r
consoleBoxplot(x)
```

**Arguments**

- `x` a numeric vector of length at least 3

**Details**

This function is for the numeric vectors. It prints a boxplot in a single line on the console. It automatically adjusts for the width of the console. The input vector must have a length of three, otherwise the function will throw a warning and not print any plot.

In case of any potential outliers (based on $1.5 \times IQR$ criteria), this will give a warning. This function is used in the explainer.

**Value**

prints a boxplot on the console which has:

- `|` at start and end means the minimum and maximum value respectively
- `<==*==>` The IQR region
- `*` shows the median
- `...` everything else in between

Gives a warning of potential outliers (if present)

**Examples**

```r
consoleBoxplot(mtcars$mpg)
```
CQassociation  

Association (Correlation) between Continuous-Categorical Variables

Description

CQassociation finds Association measure between one categorical and one continuous variable.

Usage

CQassociation(
  numtb,          
  factb,          
  method3 = c("auto", "parametric", "non-parametric"),
  use = "everything",
  normality_test_method = c("ks", "anderson", "shapiro"),
  normality_test_pval, 
  methodMat3 = NULL, 
  methods_used 
)

Arguments

tumb      a data frame with all the numerical columns. This should have at least two columns
factb     a data frame with all the categorical columns. This should have at least two columns
method3   method for association between continuous-categorical variables. Values can be "auto", "parametric", "non-parametric". See details for more information. Parametric does t-test while non-parametric does ‘Mann-Whitney’ test.
use       an optional character string giving a method for computing association in the presence of missing values. This must be (complete or an abbreviation of) one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs". If use is "everything", NAs will propagate conceptually, i.e., a resulting value will be NA whenever one of its contributing observations is NA. If use is "all.obs", then the presence of missing observations will produce an error. If use is "complete.obs" then missing values are handled by case wise deletion (and if there are no complete cases, that gives an error). "na.or.complete" is the same unless there are no complete cases, that gives NA
normality_test_method  
takes values as 'shapiro' or 'anderson'. this parameter decides which test to perform for the normality test. See details of norm_test_fun for more information.
normality_test_pval     significance level for normality tests. Default is 0.05
methodMat3  method dataframe like methodMats from the function association
methods_used a square data.frame which will store the type of association used between the variables. Dimension will be number of variables * number of variables.
Details

This function measures the association between one categorical variable and one continuous variable present in different dataset. Two datasets are provided as input, one data has only numerical columns while other data has only categorical columns. This performs either t-test for the parametric case and 'Mann-Whitney’ test for the non-parametric case. If the method3 is passed as 'auto', the function defines the method itself based on different tests for equal variance and normality check which checks for assumptions for the t-test. If the assumptions are satisfied, then t-test (parametric) is performed, otherwise 'Mann-Whitney’ (non-parametric) test is performed.

Value

a table with number of rows equal to number of columns in numtb and number of columns equal to number of columns in factb. Table containing p-values of performed test

See Also

norm_test_fun for normality test association for association between any type of variables, CCassociation for Association between Continuous (numeric) variables, QQassociation for Association between Categorical variables

Description

This function is used by [plottr()] when independent variable is continuous. This function can be used as a template to define a custom function.

Usage

Cx(dat, xname, binwidth = NULL, inc.density = T, ...)

Arguments

dat a data.frame with only one column
xname name of independent (x) variable in dat
binwidth for the histograms (extra parameters can be added like this)
inc.density Binary. True to include the density plot on histogram
... required

Value

a grob of plot
**CxCy**

*Plots for Continuous independent and dependent variables*

**Description**

This function is used by `plottr()` when both the dependent and independent variables are continuous. This function can be used as a template to define a custom function.

**Usage**

```r
CxCy(dat, xname, yname, ...)
```

**Arguments**

- `dat`: a data.frame with two columns (including the dependent)
- `xname`: name of independent (x) variable in `dat`
- `yname`: name of dependent (y) variable in `dat`
- `...`: required

**Value**

a grob of plot

---

**explainer**

*Generic explainer*

**Description**

Generic function for printing the details of data. Based on the data type, this calls the appropriate method.

**Usage**

```r
explainer(X, xname = NULL, ...)
```

**Arguments**

- `X`: a data.frame or a vector
- `xname`: name of the data to be printed. If missing then the name of variable passed as `X` will be used
- `...`: other parameters required for other methods of explainer To see the parameters for numeric methods, use `?explainer.numeric` and similarly for other methods (`?explainer.factor` etc.)
**Details**

Current methods for `explainer` are for data.frame, numeric, integer, character and factor vectors. To get the list of all available methods type the command `methods(explainer)`.

**Value**

Prints the information on the console. For print information for the individual methods, see their documentation. Returns nothing.

**Examples**

```r
# for numeric
explainer(mtcars)
explainer(mtcars$mpg) # same as explainer.numeric(mtcars$mpg)
# for factor
explainer(as.factor(mtcars$cyl)) # same as explainer.factor(as.factor(mtcars$cyl))
```

---

**Description**

This is a `explainer` method for character vector.

**Usage**

```r
## S3 method for class 'character'
explainer(X, xname = NULL, ...)
```

**Arguments**

- `X` a vector of character data type
- `xname` a placeholder for variable name
- `...` other parameters required

**Details**

This method removes all the missing values in `x` before computing the summaries.

**Value**

Prints the following information on console:

- vector name
- type
- number of distinct values
- number of missing values
- a frequency table and histogram. If counts of all the factor levels are less than half of length of \( X \), then the histogram is scaled with maximum of 50

**Examples**

```r
alphabets <- sample(LETTERS[1:5], 50, replace = TRUE)
explainer(alphabets)
rm(alphabets)
```

### Description

`explainer` shows detail of all the columns of the data

### Usage

```r
## S3 method for class 'data.frame'
explainer(X, xname = NULL, ...)
```

### Arguments

- `X`: A data.frame
- `xname`: variable name
- `...`: parameters for `explainer` for other classes

### Details

This function uses `explainer` on each column.

### Value

Prints details of the dataset which includes: dataset name, type, number of columns, rows and unique rows. Also prints output of `explainer` for all the columns. Returns nothing.

### Examples

```r
explainer(mtcars)
```
**explainer.factor**  
*Explain method for factor data types*

**Description**
This is an explainer method for factor vector.

**Usage**

```r
## S3 method for class 'factor'
explaner(X, xname = NULL, ...)
```

**Arguments**
- `X`: a numeric (or integer) data type
- `xname`: a placeholder for variable name
- `...`: other parameters required

**Details**
This method removes all the missing values in `x` before computing the summaries. This calls the method `explainer.character`.

**Value**
Prints the following information on console:

- vector name
- type
- number of distinct values
- number of missing values
- a frequency table and histogram. If counts of all the factor levels are less than half of length of `X`, then the histogram is scaled with a maximum of 50.

**Examples**

```r
alphabets <- as.factor(sample(LETTERS[1:5], 50, replace = TRUE))
explaner(alphabets)
rm(alphabets)
```
**Descripción**

Este es un método `explainer` para tipos de datos numéricos.

**Uso**

```r
## S3 method for class 'numeric'
explainer(
  X,
  xname = NULL,
  include.numeric = NULL,
  round.digit = 2,
  quant.seq = seq(0, 1, 0.2),
  trim = 0.05,
  ...
)
```

**Argumentos**

- **X**: un tipo de datos numérico (o entero)
- **xname**: un placeholder para el nombre de la variable
- **include.numeric**: un vector de cadenas que también se requiere junto con el resultado predeterminado. Puede tener valores:
  - `trimmed.means` para imprimir la media desviada después de eliminar un porcentaje `trim` de los datos de cada extremo de `X`. `trim` puede pasarse como un parámetro
  - `skewness` para imprimir la asimetría de los datos. Use `?skewness` para más información
  - `kurtosis` para imprimir la curtosis de los datos. Use `?kurtosis` para más información
- **round.digit**: el número de decimales requeridos en el resultado
- **quant.seq**: un vector de fracciones (0 a 1) para las cuantiles que se requieren. 0.5 significa mediana, 0 significa observación más pequeña y 1 significa observación más grande
- **trim**: la fracción (0 a 0.5) de observaciones que se eliminarán a cada extremo de `X` antes de calcular el promedio. Valores de trim fuera de este rango se tomarán como el próximo menor extremo. Este solo funciona si `include.numeric` tiene una cadena 'trimmed.means'

**Detalles**

Este método elimina todos los valores faltantes en `X` antes de calcular las resúmenes.
freqTable

Value

Prints the following information on console:

• vector name
• type
• number of distinct values
• number of missing values
• mean
• sd (standard deviation)
• median
• quantiles based on quant.seq parameter
• other information based on include.numeric
• a box plot (only if number distinct numbers are > 2). If counts of all the factor levels are less than half of length of x, then the histogram is scaled with maximum of 50 ?consoleBoxplot for how to read the table and histogram)
• a frequency table and histogram (only if number of distinct numbers are < 11) (look at ?freqTable for how to read the table and histogram)

Examples

explainer(mtcars$mpg)
explainer(mtcars$mpg, include.numeric = c('trimmed.means', 'skewness', 'kurtosis'), round.digit = 1, quant.seq = seq(0,1,0.1), trim = 0.05)

Description

freqTable prints a frequency table and histogram of a vector.

Usage

freqTable(Value, limit = NULL)

Arguments

Value a vector of any type
limit Upper limit of the bars in histogram. Default is NULL, for which the function will automatically find the suitable limit. This value should be in fraction (between 0 to 1)
Details

This function works for all type of vector type. But calling `freqTable` for vector with many unique values will print a very long table. If the limit parameter is left blank, then the limit of histogram is adjusted automatically and is shown at the end in brackets (eg. 50 This function is used in the explainer.

Value

Prints a table with columns

- **Value** Value. Each row has a unique value in this table
- **Freq** The frequency count of the Value
- **Proportion** Proportion of the Value \((= \text{Freq} / \text{length}(x))\)

This table is followed by a histogram with bars for each of the unique values present in the data.

Examples

```r
freqTable(mtcars$cyl)
freqTable(mtcars$mpg, limit = 0.08)
```

Description

GenerateReport generates the markdown report in one command

Usage

```r
GenerateReport(
  dtpath,
  catVars,
  yvar = NULL,
  model = "linReg",
  title = "Report",
  output_format = "html_document",
  output_dir = tempdir(),
  normality_test_method = "ks",
  interactive.plots = FALSE,
  include.vars = NULL
)
```
Arguments

dtpath      dataset path as a string

catVars    vector of categorical variables names

yvar       y variable name if present else NULL

model      type of model - linReg for linear regression binClass for binary classification and multiClass for multiclass classification

title      Title of the generated report

output_format  output report format. 'html_document' for html file or pdf_document for pdf file output. OR c("html_document", "pdf_document") for both.

output_dir  Directory where the output files needs to be stored.

normality_test_method  method for normality test for a variable. Values can be shapiro for Shapiro-Wilk test or 'anderson' for 'Anderson-Darling' test of normality or ks for 'Kolmogorov-Smirnov'

interactive.plots  for interactive variable exploration

include.vars  include only these variables from the full data

Details

This function creates a rmarkdown report which can be converted to html or pdf format file.

Value

creates a rmarkdown and html/pdf file. Returns the output directory on successful run and FALSE in case of error

Examples

# Assigning the temporary folder using tempdir(). replace with required directory
GenerateReport(dtpath = mtcars,
             catVars = c("cyl", "vs", "am", "gear"),
             yvar = "vs", model = "binClass",
             output_format = NULL,
             title = "Report",
             output_dir = tempdir(),       # pass the output directory
             interactive.plots = FALSE)   # set TRUE for interactive
Description

kurtosis calculates the Kurtosis

Usage

kurtosis(x, na.rm = T)

Arguments

x a numeric vector, matrix or a data.frame
na.rm (logical) Should missing values be removed?

Details

This function calculates the kurtosis of data which is a measure of the "tailedness" of the probability distribution of a real-valued random variable. Like skewness, kurtosis describes the shape of a probability distribution. The formula used is:

\[
\frac{E[(X - \mu)^4]}{(E[(X - \mu)^2])^2}
\]

This formula is the typical definition used in many older textbooks and wikipedia

Value

returns a single value if x is a vector, otherwise a named vector of size = ncol(x).

Examples

# for a single vector
kurtosis(mtcars$mpg)

# for a dataframe
kurtosis(mtcars)
linedivider  

Description

Draws a horizontal line on console

Usage

linedivider(consolewidth = getOption("width"), st = "x")

Arguments

consolewidth  a integer  
st  a character or symbol of length to be used for creating the line

Value

Prints a horizontal line of width 'consolewidth'

Examples

linedivider(20)

mergeAnalyzer  

Description

mergeAnalyzer analyzes the data drop after merge

Usage

mergeAnalyzer(x, y, round.digit = 2, ...)

Arguments

x  left data to merge
y  right data to merge
round.digit  integer indicating the number of decimal places to be used
...  other parameters needs to be passed to merge function

Details

Prints the summary of data retained after merge and returns the merged data. This function uses data.table (if the package is installed) for faster data merge
Value

Returns merged data with same class as that of input data. Prints summary of data retained after merging. The summary has 6 columns:

- Column: Number of rows and union of column names of numeric columns in both the data
- x, y: Sum of columns in both table
- Merged: Sum of columns in merged data
- remainingWRTx: ratio of remaining data in merged data after merging. example - 0.5 means that 50 of inner join). 1.5 means value became 150 duplicates present in data
- remainingWRTy: same as above, but for y table

Examples

# Creating two tables to merge
A <- data.frame(id = c("A", "A", "B", "D", "B"),
                valA = c(30, 83, 45, 2, 58))
B <- data.frame(id = c("A", "C", "A", "B", "C", "C"),
                valB = c(10, 20, 30, 40, 50, 60))
mergeAnalyzer(A, B, allow.cartesian = TRUE, all = FALSE)

data.frame(id = c("A", "A", "B", "D", "B"),
            valA = c(30, 83, 45, 2, 58))
data.frame(id = c("A", "C", "A", "B", "C", "C"),
            valB = c(10, 20, 30, 40, 50, 60))
mergeAnalyzer(A, B, allow.cartesian = TRUE, all = FALSE)

norm_test_fun

Description

norm_test_fun checks for the normality assumption

Usage

norm_test_fun(x, method = "anderson", pval = 0.05, xn = "x", bin = FALSE)

Arguments

x          a numeric vector
method     shapiro for Shapiro-Wilk test or 'anderson' for 'Anderson-Darling' test of normality or ks for 'Kolmogorov-Smirnov'
pval       significance level for normality tests. Default is 0.05
xn          vector name
bin         TRUE if only TRUE/FALSE is required
Details

This function checks for normality assumption using shapiro, Kolmogorov-Smirnov or Anderson Darling test. If the parameter `bin` is TRUE, then TRUE is returned if vector is normal, otherwise FALSE. The significance level is passed through the parameter `pval`.

Value

Logical TRUE/FALSE based on the performed test and `pval`. If the vector follows the normality assumption, then TRUE is returned.

See Also

anderson.test for Anderson Darling test

Examples

```r
norm_test_fun(mtcars$mpg)
norm_test_fun(mtcars$mpg, method = "shapiro",
             pval = 0.05, xn = "mpg", bin = TRUE)
```

Description

This function plots the plot generated by the library analyzer.

Usage

```r
## S3 method for class 'analyzerPlot'
plot(x, ...)
```

Arguments

- `x` a plot of class `analyzerPlot`
- `...` extra arguments if required

Value

Displays the plot

Examples

```r
# creating the plot
p <- plottr(mtcars)
plot(p$mpg)
```
plotNA  

Description

plotNA returns a grob visualizing the missing values in data.

Usage

plotNA(tb, order = T, limit = T, add_percent = T, row.level = F)

Arguments

tb  a data.frame
order  (logical) Whether to order the variables based on missing values in plot
limit  (logical) Whether to limit the plot to maximum missing value. FALSE means the limit of axis will be [0, nrow(tb)]
add_percent  (logical) Whether to add percent as data labels on bar plot
row.level  (logical) Whether to create plot at rows and variables level

Details

This is a function which helps in visualizing the missing values in data using plots. By default a bar plot is generated which shows the count of missing values in each variable.

If order is set as TRUE then the bars are arranged in order of missing values. If limit is set as TRUE then limit of axis is set to [0, nrow(tb)]. If add_percent is set as TRUE then percent is added as text to the bars. If row.level is set to TRUE then an additional plot is generated which shows which rows have missing values and in which variable (reshape2 (https://CRAN.R-project.org/package=reshape2) library is required for this).

Value

This function returns a grob of class 'analyzePlot' which has a bar plot showing the count of missing value for each variable. order, limit, add_percent can be used to modify the bar plot. An additional plot will be created and added to the grob if row.level is set as TRUE.

Examples

p <- plotNA(airquality)  
# function to show the 'analyzePlot' class plot
plot(p)
p1 <- plotNA(airquality, order = FALSE)
plot(p1)
plottr can be used to create plots for all the variables in a dataframe or any one vector. The output is a list of plots for each variable of class 'analyzerPlot'.

Usage

```r
plottr(
  tb,
  yvar = NULL,
  xclasses = NULL,
  yclass = NULL,
  printall = F,
  callasfactor = 1,
  FUN1 = Cx,
  FUN2 = Qx,
  FUN3 = CxCy,
  FUN4 = QxCy,
  FUN5 = CxQy,
  FUN6 = QxQy,
  ...
)
```

Arguments

tb: a data.frame or a vector. If yvar argument is also passed, then this should be a data.frame including the response variable (yvar)
yvar: a string showing the response (dependent) variable name. Can be NULL if response variable is not present. Make sure that this variable is present in the tb
xclasses: a vector of length = ncol(tb) with the data type of all the columns. Can be NULL, in such case function assigns a class to each column. The values have to be either NULL, or a vector of either 'factor' or 'numeric'. The order should be same as the actual columns in tb. In case when tb is a vector, this can be a vector of length 1.
yclass: class of response variable. Can be NULL, but must have value when yvar is not NULL. Value can be 'factor' or 'numeric'
printall: (logical) Whether user wants to show the plots. Setting this as FALSE will only returns a list of plots silently.
callasfactor: minimum unique values needed for x to be considered as numeric. See details for more information
FUN1: an user-defined function for plotting 1 variables when the variable is Continuous. See details for more details on how to define these variables
FUN2
same as FUN1 but for categorical variable

FUN3
an user defined function for plotting 2 variables when both the independent variable (x) and dependent variable (y) are Continuous

FUN4
same as FUN3, but when independent variable (x) is Categorical and dependent variable (y) is Continuous

FUN5
same as FUN3, but when independent variable (x) is Continuous and dependent variable (y) is Categorical

FUN6
same as FUN3, but when both the independent variable (x) and dependent variable (y) are Categorical

... extra arguments passed to functions FUN1-FUN6

Details
This is a function which helps in understanding the data through multiple visualizations. This works either for a data.frame having multiple variables or a single x variable or for a combination of predictor x and response y variables. Based on class of x and y different types of plots are automatically generated.

Please note the following points:

Defining the class of variables: If yvar is not NULL, then yclass has to be passed (which can be 'factor' for classification type problem, or 'numeric' for regression). xclasses stores the class of all the variables in the dataframe in same order of columns. Note - if yvar is not NULL, then tb has to be a data.frame with at least 2 columns (including the yvar). In such case xclasses should also have the class of yvar although it is also passed through yclass. This can also be set as NULL, in such case the function assigns a class based on the contents. If variable is factor/character type, then xclasses will have 'factor' as the entry for that variable, else if x is numeric with number of unique values less than callasfactor parameter value, then xclasses will have 'factor', else 'numeric'.

DEFINING CUSTOM FUNCTIONS FOR THE PLOTS USING FUN1, FUN2, FUN3 ... FUN6:
Custom plots can be made using these functions passed as arguments. Following things must be followed while defining such functions:

- the return plot must be of type 'grob' or 'gtables' or 'ggplot'. Since these outputs will go to arrangeGrob, make sure the output plots are acceptable by arrangeGrob function. See code of CxCy for sample.
- not all 6 functions are required to be passed. Only pass those functions for which plots need to be changed.
- FUN1 and FUN2 must have 3 parameters: dat (of type data.frame for the data. Even if there is only one column, it should be passed as a data.frame of one column), xname name of column in dat and ... In addition to these three, any number of additional parameters can be added. Look into source of code of Cx for sample.
- FUN3, FUN4, FUN5 and FUN6 must have 4 parameters: dat (of type data.frame for the data. Must have two columns for independent and dependent variables), xname name of independent variable in dat, yname name of dependent variable in dat and ... In addition to these four, any number of additional parameters can be added. Look into source of code of CxCy for sample.
- ... must be added as an argument in all the functions.
To get a better idea, see the code for function CxCy and Cx

Default plots: If the y is NULL, then histogram with density is generated for numeric x. Boxplot is also shown in the same histogram using color and vertical lines. For factor x, a pie chart showing the distribution. This are the univariate plots which can be modified by using the FUN1 and FUN2 arguments.

If y is not NULL, then additional plots are added which can be modified by using the FUN3, FUN4, FUN5, FUN6 arguments:

- **factor x, factor y**: Crosstab with heatmap (modified by using FUN6)
- **factor x, numeric y**: histogram and boxplot of y for different values of x (modified by using FUN4)
- **numeric x, factor y**: histogram and boxplot of x for different values of y (modified by using FUN5)
- **numeric x, numeric y**: Scatter plot of x and y with rug plot included (modified by using FUN3)

**Value**

A list of plots for all the variables. Each plot will have the class analyzerPlot and can be displayed using plot(). If printall = TRUE, then all plots will also be displayed.

**Examples**

```r
# simple use for one variable
p <- plottr(mtcars$mpg)
# To display the plot
plot(p$x)

# With complete dataframe and assuming 'mpg' as a dependent variable
p <- plottr(mtcars, yvar = "mpg", yclass = "numeric")
plot(p$disp)
```

**print.analyzerPlot**  
**Method for print generic**

**Description**

This function plots the plot generated by the library analyzer

**Usage**

```r
## S3 method for class 'analyzerPlot'
print(x, ...)
```
Arguments

\[ \text{x} \quad \text{a plot of class analyzerPlot} \]
\[ \ldots \quad \text{other parameters} \]

Value

Displays the plot

Examples

```r
# creating the plot
p <- plottr(mtcars$mpg)
p$x
```

## QQassociation

**Association (Correlation) between Categorical Variables**

**Description**

QQassociation finds Association measure between all the variables in data with only categorical columns.

**Usage**

```r
QQassociation(factb, use = "everything", methods_used)
```

**Arguments**

- **factb** a data frame with all the categorical columns. This should have at least two columns
- **use** an optional character string giving a method for computing association in the presence of missing values. This must be (complete or an abbreviation of) one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs". If use is "everything", NAs will propagate conceptually, i.e., a resulting value will be NA whenever one of its contributing observations is NA. If use is "all.obs", then the presence of missing observations will produce an error. If use is "complete.obs" then missing values are handled by case wise deletion (and if there are no complete cases, that gives an error). "na.or.complete" is the same unless there are no complete cases, that gives NA
- **methods_used** a square data.frame which will store the type of association used between the variables. Dimension will be number of variables * number of variables.
skewness

Details
This function measures the association between categorical variables using Chi Square test. This also returns Cramers V value which is a measure of association between two nominal variables, giving a value between 0 and +1 (inclusive). Higher number indicates higher association. Note that, unlike Pearson correlation this doesn’t give negative value.

The relation between Cramer’s V and Chi Sq test is

\[ \sqrt{\frac{\chi^2}{n \times \min(k - 1, r - 1)}} \]

where:

X is derived from Pearson’s chi-squared test
n is the grand total of observations
k being the number of columns
r being the number of rows

The p-value for the significance of Cramer’s V is the same one that is calculated using the Pearson’s chi-squared test.

Value
a list of two tables with number of rows and column equal to number of columns in factb:

chisq Table containing p-values of chi-square test
cramers Table containing Cramer’s V

See Also
association for association between any type of variables, CCassociation for Association between Continuous (numeric) variables, CQassociation for Association between Continuous-Categorical variables

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skewness

Description
skewness calculates the skewness

Usage
skewness(x, na.rm = T)
skewness

Arguments

x  a numeric vector, matrix or a data.frame
na.rm (logical) Should missing values be removed?

Details

This function calculates the skewness of data which is a measure of the asymmetry of the probability distribution of a real-valued random variable about its mean. The formula used is:

\[
\frac{E[(X - \mu)^3]}{(E[(X - \mu)^2])^{\frac{3}{2}}}
\]

This formula is the typical definition used in many older textbooks and wikipedia.

Value

returns a single value if x is a vector, otherwise a named vector of size = ncol(x).

Examples

# for a single vector
skewness(mtcars$mpg)

# for a dataframe
skewness(mtcars)
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