Package 'scitb'

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Type Package

Title Provides Some Useful Functions for Making Statistical Tables

Version 0.1.7

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Description You can use the functions provided by the package to make various statistical tables, such as baseline data tables. Creates 'Table 1', i.e., a description of the baseline patient characteristics, which is essential in every medical research. Supports both continuous and categorical variables, as well as p-values and standardized mean differences. This method was described by Mary L McHugh (2013) <doi:10.11613/bm.2013.018>.

License GPL-3

Encoding UTF-8

LazyData true

Depends R (>= 3.50)

RoxygenNote 7.2.1

Imports ggplot2, MASS, nortest, reshape2, stringi

NeedsCompilation no

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prematurity

plotsmd

Description

You can use it to draw a baseline table of data.

Usage

```
plotsmd(vars,unmatchdata,matchdata,refline=NULL,title=NULL,xlab='SMD',ylab='variable')
```

Arguments

vars	List the variables you need to compare.
unmatchdata	Data before conducting propensity matching.
matchdata	The data after propensity score matching.
refline	Set a reference line with a default value of 0.1.
title	The title of the image.
xlab	The name of the X-axis.
ylab	The name of the Y-axis.

Details

The differences between variables can be represented using SMD. This program can draw SMD graphs of variable differences.

Value

A picture.

prematurity

A data on indicators for premature newborns.

Description

A data on indicators for premature newborns.

Usage

```
data(prematurity)
```

Format

An object of class data.frame with 189 rows and 11 columns.

sci1freq

Examples

data(prematurity)

sci1freq

scilfreq

Description

You can use it to draw a baseline table of data.Creates 'Table 1', i.e., description of baseline patient characteristics, which is essential in every medical research. Supports both continuous and categorical variables, as well as p-values and standardized mean differences.

Arguments

mvars	The full range of variables you don't want to compare.
x	Enter the variables to be layered. If you fill in consecutive variables, by default they will be split into 3 layers.
data	Enter your data.
dec	The precision of the data, which defaults to 2 decimal places.
nonnormal	When the data belongs to a non-normal distribution, this parameter is needed to indicate which is variable is non-normally distributed.
statistic	Statistical effect values. Usually, it is the default F, and selecting T will return a statistical effect value.
fisher	Fisher's exact test. The default is FALSE.
correct	Chi square test for continuity correction. The default is FALSE.
Overall	Generate summary data. The default is FALSE.
smd	The default is FALSE. If it is true, return the SMD value.

Details

Table 1 represents the relationship between the baseline values of the data. This function can be easily done.

Value

A data frame.

sci1mean

Description

You can use it to draw a baseline table of data.Creates 'Table 1', i.e., description of baseline patient characteristics, which is essential in every medical research. Supports both continuous and categorical variables, as well as p-values and standardized mean differences.

Arguments

mvars	The full range of variables you don't want to compare.
x	Enter the variables to be layered. If you fill in consecutive variables, by default they will be split into 3 layers.
data	Enter your data.
dec	The precision of the data, which defaults to 2 decimal places.
nonnormal	When the data belongs to a non-normal distribution, this parameter is needed to indicate which is variable is non-normally distributed.
type	The type of encoding generally does not require input.
statistic	Statistical effect values. Usually, it is the default F, and selecting T will return a statistical effect value.
atotest	Check if the data is normally distributed. The default is T.
NormalTest	A method for detecting whether data is normally distributed. The default values are Kolmogorov Smirnov test and Kolmogorov Smirnov test. Other options are: "ad", "cvm", "pearson".
Overall	Generate summary data. The default is FALSE.
smd	The default is FALSE. If it is true, return the SMD value.

Details

Table 1 represents the relationship between the baseline values of the data. This function can be easily done.

Value

A data frame.

scitb1

Description

You can use it to draw a baseline table of data.

Usage

```
scitb1(vars,fvars=NULL,strata,data,dec,num,nonnormal=NULL,type=NULL,
statistic=F,atotest=T,NormalTest=NULL,fisher=FALSE,correct=FALSE,Overall=FALSE,smd=FALSE)
```

Arguments

vars	The full range of variables you don't want to compare.
fvars	Define the categorical variables in your data.
strata	Enter the variables to be layered. If you fill in consecutive variables, by default they will be split into 3 layers.
data	Enter your data.
dec	The precision of the data, which defaults to 2 decimal places.
num	When continuous variables are layered, use it to control the number of layers, which defaults to 3.
nonnormal	When the data belongs to a non-normal distribution, this parameter is needed to indicate which is variable is non-normally distributed.
type	The type of encoding generally does not require input.Contains three types: "A", "B", and "C".
statistic	Statistical effect values. Usually, it is the default F, and selecting T will return a statistical effect value.
atotest	Check if the data is normally distributed. The default is T.
NormalTest	A method for detecting whether data is normally distributed. The default values are Kolmogorov Smirnov test and Kolmogorov Smirnov test. Other options are: "ad", "cvm", "pearson".
fisher	Fisher's exact test. The default is FALSE.
correct	Chi square test for continuity correction. The default is FALSE.
Overall	Generate summary data. The default is FALSE.
smd	The default is FALSE. If it is true, return the SMD value.

Details

Table 1 represents the relationship between the baseline values of the data. This function can be easily done.Creates 'Table 1', i.e., description of baseline patient characteristics, which is essential in every medical research. Supports both continuous and categorical variables, as well as p-values and standardized mean differences.

scitb1

Value

A data frame.

Examples

```
## Import data
bc<-prematurity</pre>
## Hierarchical variables converted to factors.
bc$race<-as.factor(bc$race)</pre>
###Define all variables, categorical and stratified.
allVars <-c("age", "lwt", "smoke", "ptl", "ht", "ui", "ftv", "bwt")
fvars<-c("smoke","ht","ui")</pre>
strata<-"race"
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc)</pre>
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc,statistic=TRUE)</pre>
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc,statistic=TRUE,0verall=TRUE)</pre>
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc,statistic=TRUE,Overall=TRUE,smd=TRUE)
print(out)
###Stratified variables are continuous variables.
allVars <-c("race", "lwt", "smoke", "ptl", "ht", "ui", "ftv", "bwt")
fvars<-c("smoke","ht","ui","race")</pre>
strata<-"age"
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc)</pre>
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc,statistic=TRUE)</pre>
out<-scitb1(vars=allVars,fvars=fvars,strata=strata,data=bc,statistic=TRUE,Overall=TRUE,smd=TRUE)
```

```
print(out)
```

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