

# Package ‘descsuppR’

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**Version** 1.2

**License** GPL-3

**Title** Support Functions for (Reproducible) Descriptive Statistics

**Description** Contains functions to help with generating tables with descriptive statistics. In addition, the package can display results of statistical tests for group comparisons. A wide range of test procedures is supported, and user-defined test functions can be incorporated.

**Depends** foreach

**Imports** plyr, descutils, tibble, dplyr, rlang, DescTools, nparcomp, rankFD, circular, glue, purrr

**Suggests** roxygen2, survival, stringr

**Encoding** UTF-8

**Collate** 'descsuppR-package.r' 'tod.r'

**RoxygenNote** 7.2.3

**NeedsCompilation** no

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---

as.tod	<i>Conversion to and from 'Time of Day'</i>
--------	---

---

### Description

Does not safeguard against "26:69".

### Usage

```
as.tod(x)
```

```
## S3 method for class 'character'
as.tod(x)
```

```
## S3 method for class 'circular'
as.tod(x)
```

```
is.tod(x)
```

```
## S3 method for class 'tod'
as.double(x, ...)
```

```
circular(x, ...)
```

```
## Default S3 method:
circular(
  x,
  type = c("angles", "directions"),
  units = c("radians", "degrees", "hours"),
  template = c("none", "geographics", "clock12", "clock24"),
  modulo = c("asis", "2pi", "pi"),
  zero = 0,
  rotation = c("counter", "clock"),
  names = NULL,
  ...
)

## S3 method for class 'tod'
circular(x, ...)
```

### Arguments

x	An object to be converted to or represented as a circular object.
...	Additional arguments passed to methods or lower-level functions.
type	Character string indicating if data represent "angles" or "directions".
units	Character string; the measurement units for the input data ("radians", "degrees", or "hours").
template	Character string specifying a specific template for circular data.
modulo	Character string indicating the modulo arithmetic to be used ("asis", "2pi", "pi").
zero	Numeric; direction assigned as zero (in the specified units).
rotation	Character string; direction of rotation for increasing values ("counter", "clock").
names	Optional character vector of names for the object.

### Value

x converted to/from 'tod'

### Author(s)

Dr. Andreas Leha

### Examples

```
times <- c("8:53", NA, "22:30")

## some conversions
as.tod(times)
as.numeric(as.tod(times))

is.tod(times)
is.tod(as.tod(times))
```

---

`buildDescrTbl`*buildDescrTbl*

---

**Description**

Calculate and Present Descriptive Values in Pritable

**Usage**

```
buildDescrTbl(  
  df,  
  tests,  
  prmnames,  
  prmunits,  
  addFactorLevelsToNames = TRUE,  
  excel_style = TRUE,  
  groupby,  
  addungrouped = FALSE,  
  dopvals = FALSE,  
  ignore_test_errors = FALSE,  
  p.adjust.method = "holm",  
  orderedAsUnordered = FALSE,  
  factorlevellimit = 14,  
  show.minmax = TRUE,  
  show.IQR = FALSE,  
  report_tests = FALSE,  
  report_testmessages = FALSE,  
  pvals_formatting = TRUE,  
  pvals_digits = 3,  
  pvals_signiflev = 0.05,  
  extraLevels = NULL,  
  missingName = "missing",  
  nonNAsName = "N",  
  removeZeroNAs = TRUE,  
  removeZeroExtraLevels = TRUE,  
  includeNAs = FALSE,  
  includeNonNAs = FALSE,  
  printOrgAlignment = FALSE,  
  useutf8 = "latex",  
  verbose = 0,  
  without_attrs = FALSE,  
  sd_digits = "by_mean",  
  descr_digits = 2,  
  significant_digits = TRUE,  
  percentages = "columnwise"  
)
```

**Arguments**

df	data.frame containing the variables of which to calc the descriptive values
tests	character vector or list of characters or list of functions or list of lists. In each case the i-th element gives the test to perform on the ith variable in the df (excluding stratification variables). The test can either be given as character (name of test function) or as function or as list where the first element is again either character or function and the following elements are *named* additional arguments to that test function. The individual function has to accept (at least) the arguments 'values' and 'grouping' which are vectors of equal length. For convenience, this package ships with some example functions; have a look at those if you want to supply your own. These convenience functions include w.chisq.test, w.cor.test, w.fisher.test, w.kruskal.test, w.wilcox.test. the whole list/vector is recycled if too short.
prmnames	names of the variables in df (if needed to be overwritten)
prmunits	units of the variables in df
addFactorLevelsToNames	logical. if TRUE expand 'sex' to 'sex [m/w]'. Defaults to TRUE.
excel_style	logical. if TRUE remove subsequent duplicates from the parameter column (as common in Excel). Default: TRUE
groupby	column of df. do more columns - one for each group. If the df\$column is an ordered factor, the order will be respected in the resulting table
addungrouped	logical. if TRUE add a column 'total' with the ungrouped summary statistics. Default: FALSE
dopvals	boolean. if TRUE an additional column containing the p-values comparing the two strata in groupby. Only implemented for a two-level stratum until now.
ignore_test_errors	logical. If TRUE returns an empty test results (as list).
p.adjust.method	character. if not NULL include an additional column with adjusted p values. see <a href="#">p.adjust.methods</a> for possible values and explanations. Defaults to "holm"
orderedAsUnordered	logical. treat ordered factors as unordered factors?
factorlevellimit	integer. for factors with more than factorlevellimit levels, not all levels are printed
show.minmax	logical. if TRUE show minimum and maximum for numeric variables. Defaults to TRUE.
show.IQR	logical. if TRUE show 25% and 75% quantiles for numeric variables. Defaults to FALSE.
report_tests	boolean. if TRUE one additional column in the result table will contain the test, that was performed to calculate the p value. Ignored if dopvals=FALSE
report_testmessages	boolean. if TRUE one additional column in the result table will contain any warnings that appeared while the test was performed. Ignored if dopvals=FALSE

pvals_formatting	boolean. If FALSE report numbers, else report formatted strings (via prettyP-value)
pvals_digits	integer. Number of digits for p value formatting. Ignored when pvals_formatting==FALSE. Defaults to 2
pvals_signiflev	double. The significance level for bold p value formatting. Ignored when pvals_formatting==FALSE. Defaults to 0.05
extraLevels	named list of lists. Names have to be variable names. Elements have to be named list of this form: `some label` = list(idxvec = idxvec, display = logical). Here idxvec needs to be a logical vector of length nrow(df) that specifies the affected rows. If display is TRUE the number of affected rows will be shown under some label.
missingName	character. name of missing values (default: missing)
nonNAsName	character. name of not missing values (default: N)
removeZeroNAs	boolean. if TRUE, rows for missing values containing only 0s are removed from the result.
removeZeroExtraLevels	boolean. if TRUE, rows for ExtraLevels containing only 0s are removed from the result.
includeNAs	boolean. Include number of NAs in the output? Currently only one of either includeNonNAs or includeNAs can be set to TRUE
includeNonNAs	boolean. Include number of not missing values (Non-NAs) in the output? Currently only one of either includeNonNAs or includeNAs can be set to TRUE
printOrgAlignment	boolean. If TRUE, than a row like "<l> <r> <r>" will be included in the result df
useutf8	character. one of c("latex", "utf8", "replace"). if 'latex' (the default) use \pm in the output; if 'replace' use +- in the output, if 'utf8' use the unicode character
verbose	numeric. level of verbosity (0 : silent)
without_attrs	logical. If TRUE return the descriptive table with attrs. Otherwise add df, groupby, and a 'full' (closer to tidy) version of the table as attributes. Defaults to TRUE.
sd_digits	character. one of c("by_mean", "fixed"). If 'by_mean', the number of decimal places of the standard deviation is limited by the number of decimal places of the mean.
descr_digits	integer. Number of digits for formatting of descriptive values. Defaults to 2.
significant_digits	boolean. if TRUE, the number of significant digits of is given by descr_digits. Otherwise the number of decimal places is fixed.
percentages	character. one of c("columnwise", "rowwise"). If 'rowwise', percentages are computed by row. Defaults to "columnwise"

## Details

Do a Table containing descriptive values

**Value**

formatted data.frame with descriptive values

**Author(s)**

Andreas Leha

**Examples**

```
ttt <- data.frame(data="training set",
  age=runif(100, 0, 100),
  sex=sample(c("m","f"), 100, replace=TRUE, prob=c(0.3, 0.7)),
  score=factor(sample(1:5, 100, replace=TRUE),
    ordered=TRUE,
    levels=1:5))
ttt2 <- data.frame(data="test set",
  age=runif(100, 0, 100),
  sex=sample(c("m","f"), 100, replace=TRUE, prob=c(0.5,0.5)),
  score=factor(sample(1:5, 100, replace=TRUE),
    ordered=TRUE,
    levels=1:5))

units <- c("years", "", "")
buildDescrTbl(rbind(ttt, ttt2),
  prmunits=units,
  groupby="data",
  includeNAs=TRUE)
```

---

buildDescrTbl.intern    *buildDescrTbl.intern*

---

**Description**

Internal wrapper around calc\_descr\_matrix

**Usage**

```
buildDescrTbl.intern(
  df,
  prmnames,
  prmunits,
  addFactorLevelsToNames = TRUE,
  extraLevels = NULL,
  includeNAs = FALSE,
  includeNonNAs = FALSE,
  orderedAsUnordered = FALSE,
  factorlevellimit = 14,
  show.minmax = TRUE,
```

```

show.IQR = FALSE,
sd_digits = "by_mean",
descr_digits = 2,
significant_digits = TRUE
)

```

### Arguments

<code>df</code>	data.frame containing the variables of which to calc the descriptive values
<code>prmnames</code>	names of the variables in df (if needed to be overwritten)
<code>prmunits</code>	units of the variables in df
<code>addFactorLevelsToNames</code>	logical. if TRUE expand 'sex' to 'sex [m/w]'. Defaults to TRUE.
<code>extraLevels</code>	named list of lists. Names have to be variable names. Elements have to have to be named list of this form: <code>`some label` = list(idxvec = idxvec, display = logical)</code> . Here <code>idxvec</code> needs to be a logical vector of length <code>nrow(df)</code> that specifies the affected rows. If <code>display</code> is TRUE the number of affected rows will be shown under some label.
<code>includeNAs</code>	boolean. Include number of NAs in the output? Currently only one of either <code>includeNonNAs</code> or <code>includeNAs</code> can be set to TRUE
<code>includeNonNAs</code>	boolean. Include number of not missing values (Non-NAs) in the output? Currently only one of either <code>includeNonNAs</code> or <code>includeNAs</code> can be set to TRUE
<code>orderedAsUnordered</code>	logical. treat ordered factors as unordered factors?
<code>factorlevellimit</code>	integer. for factors with more than <code>factorlevellimit</code> levels, not all levels are printed
<code>show.minmax</code>	logical. if TRUE show minimum and maximum for numeric variables. Defaults to TRUE.
<code>show.IQR</code>	logical. if TRUE show 25% and 75% quantiles for numeric variables. Defaults to FALSE.
<code>sd_digits</code>	character. one of <code>c("by_mean", "fixed")</code> . If 'by_mean', the number of decimal places of the standard deviation is limited by the number of decimal places of the mean.
<code>descr_digits</code>	integer. Number of digits for formatting of descriptive values. Defaults to 2.
<code>significant_digits</code>	boolean. if TRUE, the number of significant digits of is given by <code>descr_digits</code> . Otherwise the number of decimal places is fixed.

### Value

matrix with descriptive values

### Author(s)

Andreas Leha

---

calc\_descr\_matrix      *calc\_descr\_matrix*

---

## Description

Wrapper for single vectors

## Usage

```
calc_descr_matrix(
  ttt,
  format = "long",
  extraLevels = NULL,
  includeNAs = FALSE,
  includeNonNAs = FALSE,
  orderedAsUnordered = FALSE,
  factorlevellimit = 14,
  show.minmax = TRUE,
  show.IQR = FALSE,
  sd_digits = "by_mean",
  descr_digits = 2,
  significant_digits = TRUE
)
```

## Arguments

ttt	the data.frame
format	in c("long", "wide")
extraLevels	named list of lists. Names have to be variable names. Elements have to be named list of this form: <code>`some label` = list(idxvec = idxvec, display = logical)</code> . Here <code>idxvec</code> needs to be a logical vector of length <code>nrow(df)</code> that specifies the affected rows. If <code>display</code> is <code>TRUE</code> the number of affected rows will be shown under <code>some label</code> .
includeNAs	boolean. include number of NAs in the output?
includeNonNAs	boolean. include number of non-NAs in the output?
orderedAsUnordered	logical. treat ordered factors as unordered factors?
factorlevellimit	integer. for factors with more than <code>factorlevellimit</code> levels, not all levels are printed
show.minmax	logical. if <code>TRUE</code> show minimum and maximum for numeric variables. Defaults to <code>TRUE</code> .
show.IQR	logical. if <code>TRUE</code> show 25% and 75% quantiles for numeric variables. Defaults to <code>FALSE</code> .

<code>sd_digits</code>	character. one of <code>c("by_mean", "fixed")</code> . If <code>'by_mean'</code> , the number of decimal places of the standard deviation is limited by the number of decimal places of the mean.
<code>descr_digits</code>	integer. Number of digits for formatting of descriptive values. Defaults to 2.
<code>significant_digits</code>	boolean. if TRUE, the number of significant digits of is given by <code>descr_digits</code> . Otherwise the number of decimal places is fixed.

**Details**

calls the one dimensional functions

**Value**

matrix containing the descriptive values

**Author(s)**

Andreas Leha

---

`convertColumnHeading` *Convert Grouping Column Values into Headings with `ddply`*

---

**Description**

For each unique value in the specified column of a data frame, creates a "subtable" where that value is moved as a group heading (first row) and the column is blanked for the remaining rows. Optionally, can "move along" specified metadata columns, shifting their first element to the last row.

**Usage**

```
convertColumnHeading(df, col, movealong = NULL)
```

**Arguments**

<code>df</code>	A data frame to be processed.
<code>col</code>	Character. Name of the column in <code>df</code> to use for grouping and converting to headings.
<code>movealong</code>	Optional character vector of column names. For each group, these columns are rotated, placing their first value after the last row.

**Details**

The function uses `plyr::ddply` to split the data frame by the levels of `col`. For each group, - Moves the grouping value to a heading (top row) - Adds an empty row for formatting - Optionally rotates metadata columns specified in `movealong` This can be useful for preparing tables for presentation or reporting.

**Value**

A data frame formatted with expanded headings per group.

**Author(s)**

Andreas Leha

**Examples**

```
df <- data.frame(Group = c("A", "A", "B", "B"), Data = 1:4, Meta = c("x", "y", "z", "w"))
convertColumnHeading(df, "Group")
convertColumnHeading(df, "Group", movealong = "Meta")
```

---

descrSurvEstimate      *descrSurvEstimate*

---

**Description**

build a description table for survival estimates

**Usage**

```
descrSurvEstimate(
  S,
  strata,
  stratorder,
  survname = "survival time",
  evaltimes = c(3, 5),
  evaltimeunits = "years",
  digits = 2,
  includeNAs = TRUE,
  includeNonNAs = TRUE,
  missingName = "missing",
  nonNAsName = "N",
  stratheader = TRUE,
  pval = FALSE,
  pvals_formatting = TRUE,
  pvals_digits = 2,
  pvals_signiflev = 0.05,
  hr = FALSE
)
```

**Arguments**

**S** survival objects from [Surv](#)

**strata** a list of vectors containing strata. If the vectors are ordered factors the columns will be used in the given order.

stratorder	(list of) character vector for the order of the reported columns. Overrides any order of strata
survname	the name of the survival time, e.g. 'DFS'
evaltimes	numeric vector. for which times to calculate the survival estimate
evaltimeunits	the unit of the survival times (years, months, ...)
digits	round to
includeNAs	boolean. Include number of NAs in the output?
includeNonNAs	Logical; if TRUE, include a column reporting the number of non-missing (non-NA) observations in the output table. Default is TRUE.
missingName	character. name of the rows with missing numbers. Defaults to "missing".
nonNAsName	Character string; the name to use as the column heading for the number of non-missing (non-NA) values. Default is "N".
stratheader	boolean. print the stratheader? Turn off for inclusion into a bigger table
pval	boolean. if TRUE, the p value from a cox model is printed in a separate column
pvals_formatting	boolean. If FALSE report numbers, else report formatted strings (via prettyP-value)
pvals_digits	integer. Number of digits for p value formatting. Ignored when pvals_formatting==FALSE. Defaults to 2
pvals_signiflev	double. The significance level for bold p value formatting. Ignored when pvals_formatting==FALSE. Defaults to 0.05
hr	boolean. if TRUE, the hazard ratio (with confidence interval) is printed as well. (only has an effect if pval==TRUE)

### Details

calculate the survival estimates at the specified times and return a nicely formatted table

### Value

a character matrix

### Author(s)

Andreas Leha

### Examples

```
if (require("survival")) {
  S <- Surv(aml$time, aml$status)
  descrSurvEstimate(S,
    evaltimes=c(19, 24),
    evaltimeunits="months")
}
```

---

hours.tod	<i>Extract the hours/minutes Component of time-of-day</i>
-----------	---

---

**Description**

Extract the hours/minutes Component of time-of-day

**Usage**

```
hours.tod(x)
```

```
minutes.tod(x)
```

**Arguments**

x                    vector of class tod

**Value**

numeric vector of length length(x) with hours/minutes

**Author(s)**

Dr. Andreas Leha

**Examples**

```
times <- as.tod(c("8:53", NA, "22:30"))
hours.tod(times)
minutes.tod(times)
```

---

length.tod	<i>Basic Functions on time-of-day Vectors</i>
------------	---

---

**Description**

These should just do whatever the same function does for character vectors

**Usage**

```
## S3 method for class 'tod'
length(x)

## S3 method for class 'tod'
x[i, ...]

## S3 replacement method for class 'tod'
x[i, ...] <- value

## S3 method for class 'tod'
x[[i, ...]]

## S3 replacement method for class 'tod'
x[[i, ...]] <- value

## S3 method for class 'tod'
c(...)
```

**Arguments**

x	vector of class tod
i	integer of indices
...	to match the generics' arguments
value	of type tod. to be inserted

**Value**

as expected

**Author(s)**

Dr. Andreas Leha

**Examples**

```
times <- c("8:53", NA, "22:30")
times <- as.tod(times)

length(times)
times[1]
times[1] <- "07:00"
c(times, times)
```

---

`mean.tod`*Descriptive Statistics for time-of-day Vectors*

---

**Description**

Functions to calculate descriptive values for time-of-day vectors. The heavy lifting is done by the circular package.

**Usage**

```
## S3 method for class 'tod'
mean(x, ...)

sd(x, ...)

## Default S3 method:
sd(x, na.rm = FALSE, ...)

## S3 method for class 'tod'
sd(x, na.rm = FALSE, ...)

## S3 method for class 'tod'
median(x, na.rm = FALSE, ...)

## S3 method for class 'tod'
min(..., na.rm = FALSE)

## S3 method for class 'tod'
max(..., na.rm = FALSE)

## S3 method for class 'tod'
quantile(x, ...)
```

**Arguments**

<code>x</code>	vector of class <code>tod</code>
<code>...</code>	additional args passed on to base stats or circular stats functions
<code>na.rm</code>	often passed through to corresponding functions in the circular packages. Otherwise logical. Defaults to <code>FALSE</code> .

**Details**

These functions are meant to provide the least surprising descriptive values.

**Value**

descriptive values as would be returned for non-circular numeric vectors

**Author(s)**

Dr. Andreas Leha

**Examples**

```
times <- as.tod(c("21:53", NA, "22:30", "23:10", "23:58", "00:15", "01:01"))  
  
mean(times)  
mean(times, na.rm = TRUE)
```

---

pred.survfit

*pred.survfit*

---

**Description**

X-years prediction

**Usage**

```
pred.survfit(Sfit, time)
```

**Arguments**

Sfit	a survfit object
time	the time to calculate the estimate

**Details**

Calculate the X-years estimate of a survfit

**Value**

the estimate

**Author(s)**

Andreas Leha

---

replaceGermanUmlauts *Replace German Umlaute*

---

**Description**

Replace German Umlaute

**Usage**

```
replaceGermanUmlauts(txt)
```

**Arguments**

txt                    character. within this txt the German umlauts will be replaced

**Value**

character. version of txt with all 'Umlaute' and 'scharfes s' replaced.

**Author(s)**

Andreas Leha

**Examples**

```
replaceGermanUmlauts("gefräßig")
```

---

testWrapper *Collect Warnings From Running testfun*

---

**Description**

wrap this around 'correlation' tests to get output formatted for buildDescrTbl

**Usage**

```
testWrapper(testfun, values, grouping, ignore_test_errors = FALSE, ...)
```

**Arguments**

testfun                character or function. Which function to call.  
 values                vector. The values to compare (age, toxicity score, gene expression, ...)  
 grouping              vector of the same length as values. treated as factor giving the group membership  
 ignore\_test\_errors    logical. If TRUE returns an empty test results (as list).  
 ...                    additional parameters. are passed on to the testfun

**Details**

This function is called by buildDescrTbl in order to generate the comparison p values. Basically it just calls the provided testfun. Main purpose is, that it collects warnings and returns them as well.

**Value**

list. the results from testfun plus the element 'warnings' containing all warnings collected from the run of testfun. the results from testfun are assumed to be of type list and are additionally assumed to contain at least the elements 'p.value' and 'method'.

**Author(s)**

Andreas Leha

---

w.anova.test

*ANOVA with unified interface*

---

**Description**

One-way ANOVA that unifies the api to other tests

**Usage**

```
w.anova.test(values, grouping, na.rm = TRUE, ...)
```

**Arguments**

values	vector. The values to compare (age, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
na.rm	logical. if TRUE (default) the values are subset to only non-missing and finite values
...	additional parameters. are passed on to lm()

**Value**

the value of anova augmented by 'p.value' and 'method'

**Author(s)**

Dr. Andreas Leha

---

w.chisq.test	<i>chisq.test with unified interface</i>
--------------	--

---

**Description**

chisq.test with unified interface

**Usage**

```
w.chisq.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the chisq.test

**Details**

just a call to chisq.test that unifies the api to other tests

**Value**

the value chisq.test

**Author(s)**

Andreas Leha

---

w.CochraneArmitageTrend.test	<i>Cochran-Armitage Test for Trend</i>
------------------------------	--

---

**Description**

Wrapper for [CochranArmitageTest](#) which is a test for trend in binomial proportions across the levels of a single variable

**Usage**

```
w.CochraneArmitageTrend.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare. Will be passed to <a href="#">circular</a> to be converted to circular format
grouping	of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the <a href="#">CochranArmitageTest</a>

**Value**

the value of [CochranArmitageTest](#)

**Author(s)**

Dr. Andreas Leha

---

w.cor.test

*cor.test with unified interface*

---

**Description**

cor.test with unified interface

**Usage**

```
w.cor.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the cor.test

**Details**

just a call to cor.test that unifies the api to other tests

**Value**

the value cor.test

**Author(s)**

Andreas Leha

---

w.fisher.test	<i>fisher.test with unified interface</i>
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---

**Description**

fisher.test with unified interface

**Usage**

```
w.fisher.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the fisher.test

**Details**

just a call to fisher.test that unifies the api to other tests computes the exact p-value if possible and simulates the p-value otherwise

**Value**

the value fisher.test

**Author(s)**

Andreas Leha

---

w.JonckheereTerpstraTest	<i>JonckheereTerpstraTest with unified interface</i>
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---

**Description**

JonckheereTerpstraTest with unified interface

**Usage**

```
w.JonckheereTerpstraTest(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the JonckheereTerpstraTest

**Details**

just a call to JonckheereTerpstraTest that unifies the api to other tests

**Value**

the value JonckheereTerpstraTest

**Author(s)**

Fabian Kück

---

w.kruskal.test	<i>kruskal.test with unified interface</i>
----------------	--

---

**Description**

kruskal.test with unified interface

**Usage**

```
w.kruskal.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the kruskal.test

**Details**

just a call to kruskal.test that unifies the api to other tests

**Value**

the value kruskal.test

**Author(s)**

Andreas Leha

---

w.no.test	<i>no test</i>
-----------	----------------

---

**Description**

no test but unified interface

**Usage**

```
w.no.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. unused

**Details**

just returns NA. Included as pass through for non-testable variables

**Value**

NA

**Author(s)**

Andreas Leha

---

w.npar.t.test	<i>npar.t.test with unified interface</i>
---------------	---

---

**Description**

npar.t.test with unified interface

**Usage**

```
w.npar.t.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the npar.t.test

**Details**

just a call to npar.t.test that unifies the api to other tests computes the ordinary Brunner-Munzel test for group sizes > 9 and the studentized permutation test version otherwise

**Value**

the value npar.t.test

**Author(s)**

Fabian Kück

---

w.npar.t.test.permu    *npar.t.test with unified interface*

---

**Description**

npar.t.test with unified interface

**Usage**

```
w.npar.t.test.permu(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the npar.t.test

**Details**

just a call to npar.t.test that unifies the api to other tests computes the the studentized permutation test version of the Brunner-Munzel test

**Value**

the value npar.t.test

**Author(s)**

Fabian Kück

---

w.prop.trend.test      *prop.trend.test with unified interface*

---

**Description**

prop.trend.test with unified interface

**Usage**

```
w.prop.trend.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the prop.trend.test

**Details**

just a call to prop.trend.test that unifies the api to other tests

**Value**

the value prop.trend.test

**Author(s)**

Fabian Kück

---

w.rankFD.mid.ranks      *rankFD with unified interface*

---

**Description**

rankFD with unified interface

**Usage**

```
w.rankFD.mid.ranks(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the rankFD

**Details**

just a call to rankFD with effect="weighted" that unifies the api to other tests

**Value**

the value rankFD

**Author(s)**

Fabian Kück

---

w.rankFD.pseudo.ranks *rankFD with unified interface*

---

**Description**

rankFD with unified interface

**Usage**

```
w.rankFD.pseudo.ranks(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the rankFD

**Details**

just a call to rankFD that unifies the api to other tests

**Value**

the value rankFD

**Author(s)**

Fabian Kück

---

w.t.test	<i>t.test with unified interface</i>
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---

**Description**

just a call to t.test that unifies the api to other tests

**Usage**

```
w.t.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the t.test

**Value**

the value of t.test

**Author(s)**

Andreas Leha

---

w.watson.williams.test	<i>Watson-Williams Test of Homogeneity of Means</i>
------------------------	---

---

**Description**

just a call to [watson.williams.test](#) that unifies the api to other tests

**Usage**

```
w.watson.williams.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare. Will be passed to <a href="#">circular</a> to be converted to circular format
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the <a href="#">watson.williams.test</a>

**Value**

the value of `watson.williams.test`

**Author(s)**

Dr. Andreas Leha

---

w.wilcox.test

*wilcox.test with unified interface*

---

**Description**

wilcox.test with unified interface

**Usage**

```
w.wilcox.test(values, grouping, ...)
```

**Arguments**

values	vector. The values to compare (age, toxicity score, gene expression, ...)
grouping	vector of the same length as values. treated as factor giving the group membership
...	additional parameters. are passed on to the wilcox.test

**Details**

just a call to wilcox.test that unifies the api to other tests

**Value**

the value wilcox.test

**Author(s)**

Andreas Leha

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