# Package 'bruceR'

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Title Broadly Useful Convenient and Efficient R Functions

**Version** 0.8.10 **Date** 2023-03-03

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**Description** Broadly useful convenient and efficient R functions that bring users concise and elegant R data analyses.

This package includes easy-to-use functions for

(1) basic R programming

(e.g., set working directory to the path of currently opened file; import/export data from/to files in any format;

print tables to Microsoft Word);

(2) multivariate computation

(e.g., compute scale sums/means/... with reverse scoring);

- (3) reliability analyses and factor analyses;
- (4) descriptive statistics and correlation analyses;
- (5) t-test, multi-factor analysis of variance (ANOVA), simple-effect analysis, and post-hoc multiple comparison;
- (6) tidy report of statistical models
- (to R Console and Microsoft Word);
- (7) mediation and moderation analyses (PROCESS);

and (8) additional toolbox for statistics and graphics.

License GPL-3

**Encoding** UTF-8

LazyData true

LazyDataCompression xz

URL https://psychbruce.github.io/bruceR/

BugReports https://github.com/psychbruce/bruceR/issues

**Depends** R (>= 4.0.0)

Imports rstudioapi, data.table, rio, plyr, dplyr, tidyr, stringr, ggplot2, psych, afex, emmeans, effectsize, performance, mediation, interactions, lavaan, jtools, texreg, lmerTest, MuMIn

**Suggests** pacman, glue, crayon, tibble, forcats, haven, foreign, readxl, openxlsx, clipr, cowplot, ggtext, see, car, lmtest, lme4, nnet, vars, phia, MASS, BayesFactor, GGally, GPArotation

RoxygenNote 7.2.3 NeedsCompilation no Repository CRAN

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# $\mathsf{R}$ topics documented:

bruceR-package	
add	
Alpha	
cc	. 9
ecf_plot	. 10
CFA	. 12
Corr	. 13
cor_diff	. 15
Describe	. 15
ltime	
EFA	. 18
EMMEANS	. 21
export	
formatF	. 27
formatN	. 28
formula_expand	. 28
formula_paste	. 29
Freq	. 30
GLM_summary	. 31
grand_mean_center	. 32
granger_causality	. 33
granger_test	. 34
group_mean_center	. 35
HLM_ICC_rWG	. 36
HLM_summary	. 38
mport	. 39
avaan_summary	. 41
LOOKUP	
MANOVA	. 45
med_summary	
model_summary	
)	. 53
pkg_depend	
okg_install_suggested	
Print	
orint_table	. 56

bruceR-package	2
птисек-паскаде	<b>1</b>

PROCESS .					 		•						•	 •	•	•	•	 			
RECODE .					 													 			
regress					 													 			
rep_char					 													 			
RESCALE.					 													 			
RGB					 													 			
Run					 													 			
scaler					 													 			
set.wd					 													 			
$show\_colors$					 																
theme_bruce					 													 			
TTEST					 																
$% \operatorname{allin} % \left( {\operatorname{allin} % } \right) = \operatorname{allin} % \left( {\operatorname{allin} % } \right$					 													 			
$% \operatorname{anyin} % \left( {\operatorname{anyin} } \right) = 0$					 																
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%partin% .					 																
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bruceR-package

bruceR: BRoadly Useful Convenient and Efficient R functions

# Description

BRoadly Useful Convenient and Efficient R functions that BRing Users Concise and Elegant R data analyses.

Package homepage: https://psychbruce.github.io/bruceR/

Install the latest development version from GitHub: devtools::install\_github("psychbruce/bruceR") Report bugs at GitHub Issues.

#### **Details**

Loading bruceR by library(bruceR) will also load these R packages for you:

# [Data]:

- dplyr: Data manipulation and processing.
- tidyr: Data cleaning and reshaping.
- stringr: Toolbox for string operation (with regular expressions).
- forcats: Toolbox for factor manipulation (for categorical variables).
- data.table: Advanced data.frame with higher efficiency.

## [Stat]:

4 bruceR-package

- emmeans: Estimates of marginal means and multiple contrasts.
- effectsize: Estimates of effect sizes and standardized parameters.
- performance: Estimates of regression models performance.
- lmerTest: Tests of linear mixed effects models (LMM, also known as HLM and multilevel models).

# [Plot]:

- ggplot2: Data visualization.
- ggtext: Markdown/HTML rich text format for ggplot2 (geoms and themes).
- cowplot: Advanced toolbox for ggplot2 (arrange multiple plots and add labels).
- see: Advanced toolbox for ggplot2 (geoms, scales, themes, and color palettes).

#### Main Functions in bruceR

(1) Basic R Programming set.wd (alias: set\_wd)

```
import, export
cc
pkg_depend, pkg_install_suggested
formatF, formatN
print_table
Print, Glue, Run
%^%
%notin%
%allin%, %anyin%, %nonein%, %partin%
```

(2) Multivariate Computation add, added

```
. sum, .mean
SUM, MEAN, STD, MODE, COUNT, CONSEC
RECODE, RESCALE
LOOKUP
```

(3) Reliability and Factor Analyses Alpha

```
EFA / PCA
CFA
```

(4) Descriptive Statistics and Correlation Analyses Describe

```
Freq
Corr
cor_diff
```

(5) T-Test, Multi-Factor ANOVA, Simple-Effect Analysis, and Post-Hoc Multiple Comparison

```
TTEST
MANOVA
EMMEANS
```

add 5

### (6) Tidy Report of Regression Models model\_summary

```
lavaan_summary
GLM_summary
HLM_summary
HLM_ICC_rWG
regress
```

(7) Mediation and Moderation Analyses PROCESS

```
med_summary
```

(8) Additional Toolbox for Statistics and Graphics grand\_mean\_center

```
group_mean_center
ccf_plot
granger_test
granger_causality
theme_bruce
show_colors
```

#### Author(s)

Han-Wu-Shuang (Bruce) Bao

add

Create, modify, and delete variables.

#### **Description**

Enhanced functions to create, modify, and/or delete variables. The functions **combine** the advantages of within (base), mutate (dplyr), transmute (dplyr), and := (data.table). See examples below for the usage and convenience.

## Usage

```
add(data, expr, when, by, drop = FALSE)
added(data, expr, when, by, drop = FALSE)
```

# **Arguments**

data A data.table (preferred).

expr R expression(s) enclosed in {...} to compute variables.

Passing to data.table: DT[, `:=`(expr), ]

Execute each line of expression in {...} one by one, such that newly created variables are available immediately. This is an advantage of mutate and has been implemented here for data.table.

6 add

```
when [Optional] Compute for which rows or rows meeting what condition(s)?

Passing to data.table: DT[when, , ]

by [Optional] Compute by what group(s)?

Passing to data.table: DT[ , , by]

drop Drop existing variables and return only new variables? Default is FALSE, which returns all variables.
```

#### Value

```
add() returns a new data.table, with the raw data unchanged. added() returns nothing and has already changed the raw data.
```

#### **Functions**

• add(): Return the *new data*.

You need to assign the new data to an object:

```
data = add(data, {...})
```

• added(): Return nothing and *change the raw data immediately*. NO need to assign the new data:

```
added(data, {...})
```

```
## ===== Usage 1: add() ===== ##
d = as.data.table(within.1)
d$XYZ = 1:8
# add() does not change the raw data:
add(d, \{B = 1; C = 2\})
# new data should be assigned to an object:
d = d \% > % add({
  ID = str_extract(ID, "\\d") # modify a variable
  XYZ = NULL
                               # delete a variable
  A = .mean("A", 1:4)
                               # create a new variable
  B = A * 4 # new variable is immediately available
  C = 1
               # never need ,/; at the end of any line
})
d
## ===== Usage 2: added() ===== ##
d = as.data.table(within.1)
d$XYZ = 1:8
```

add 7

```
# added() has already changed the raw data:
added(d, \{B = 1; C = 2\})
# raw data has already become the new data:
 ID = str_extract(ID, "\\d")
 XYZ = NULL
 A = .mean("A", 1:4)
 B = A * 4
 C = 1
})
d
## ===== Using `when` and `by` ===== ##
d = as.data.table(between.2)
d
added(d, {SCORE2 = SCORE - mean(SCORE)},
      A == 1 \& B \%in\% 1:2, # `when`: for what conditions
                           # `by`: by what groups
      by=B)
na.omit(d)
## ===== Return Only New Variables ===== ##
newvars = add(within.1, {
 ID = str_extract(ID, "\\d")
 A = .mean("A", 1:4)
}, drop=TRUE)
newvars
## ===== Better Than `base::within()` ===== ##
d = as.data.table(within.1)
# wrong order: C B A
within(d, {
 A = 4
 B = A + 1
 C = 6
})
# correct order: A B C
add(d, {
 A = 4
 B = A + 1
```

8 Alpha

```
C = 6
```

Alpha

*Reliability analysis (Cronbach's*  $\alpha$  *and McDonald's*  $\omega$ ).

# Description

An extension of psych::alpha() and psych::omega(), reporting (1) scale statistics (Cronbach's  $\alpha$  and McDonald's  $\omega$ ) and (2) item statistics (item-rest correlation [i.e., corrected item-total correlation] and Cronbach's  $\alpha$  if item deleted).

Three options to specify variables:

- 1. var + items: common and unique parts of variable names (suggested).
- 2. vars: a character vector of variable names (suggested).
- 3. varrange: starting and stopping positions of variables (NOT suggested).

# Usage

```
Alpha(
  data,
  var,
  items,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  digits = 3,
  nsmall = digits
)
```

# Arguments

data	Data frame.
var	[Option 1] The common part across the variables. e.g., "RSES"
items	[Option 1] The unique part across the variables. e.g., 1:10
vars	[Option 2] A character vector specifying the variables. e.g., c("X1", "X2", "X3", "X4", "X5")
varrange	[Option 3] A character string specifying the positions ("starting:stopping") of variables. e.g., "A1:E5"
rev	[Optional] Variables that need to be reversed. It can be (1) a character vector specifying the reverse-scoring variables (recommended), or (2) a numeric vector specifying the item number of reverse-scoring variables (not recommended).
digits, nsmall	Number of decimal places of output. Default is 3.

cc 9

### Value

A list of results obtained from psych::alpha() and psych::omega().

#### See Also

```
MEAN, EFA, CFA
```

## **Examples**

```
# ?psych::bfi
data = psych::bfi
Alpha(data, "E", 1:5)  # "E1" & "E2" should be reversed
Alpha(data, "E", 1:5, rev=1:2)  # correct
Alpha(data, "E", 1:5, rev=cc("E1, E2"))  # also correct
Alpha(data, vars=cc("E1, E2, E3, E4, E5"), rev=cc("E1, E2"))
Alpha(data, varrange="E1:E5", rev=cc("E1, E2"))
# using dplyr::select()
data %>% select(E1, E2, E3, E4, E5) %>%
    Alpha(vars=names(.), rev=cc("E1, E2"))
```

CC

*Split up a string (with separators) into a character vector.* 

### **Description**

Split up a string (with separators) into a character vector (whitespace around separator is trimmed).

## Usage

```
cc(..., sep = "auto", trim = TRUE)
```

# Arguments

```
... Character string(s).
```

trim Remove whitespace from start and end of string(s)? Default is TRUE.

## Value

Character vector.

10 ccf\_plot

## **Examples**

ccf\_plot

Cross-correlation analysis.

# Description

Plot the results of cross-correlation analysis using ggplot2 (rather than R base plot) for more flexible modification of the plot.

## Usage

```
ccf_plot(
  formula,
  data,
  lag.max = 30,
  sig.level = 0.05,
  xbreaks = seq(-100, 100, 10),
 ybreaks = seq(-1, 1, 0.2),
  ylim = NULL,
  alpha.ns = 1,
  pos.color = "black",
  neg.color = "black",
  ci.color = "blue",
  title = NULL,
  subtitle = NULL,
 xlab = "Lag",
 ylab = "Cross-Correlation"
)
```

ccf\_plot

# Arguments

formula	Model formula like y ~ x.
data	Data frame.
lag.max	Maximum time lag. Default is 30.
sig.level	Significance level. Default is 0.05.
xbreaks	X-axis breaks.
ybreaks	Y-axis breaks.
ylim	Y-axis limits. Default is NULL to automatically estimate.
alpha.ns	Color transparency (opacity: 0~1) for non-significant values. Default is 1 for no transparency (i.e., opaque color).
pos.color	Color for positive values. Default is "black".
neg.color	Color for negative values. Default is "black".
ci.color	Color for upper and lower bounds of significant values. Default is "blue".
title	Plot title. Default is an illustration of the formula.
subtitle	Plot subtitle.
xlab	X-axis title. Default is "Lag".
ylab	Y-axis title. Default is "Cross-Correlation".

## **Details**

Significant correlations with *negative time lags* suggest shifts in a predictor *precede* shifts in an outcome.

## Value

A gg object, which you can further modify using ggplot2 syntax and save using ggsave().

#### See Also

```
granger_test
```

12 CFA

CFA

Confirmatory Factor Analysis (CFA).

# Description

```
An extension of lavaan::cfa().
```

# Usage

```
CFA(
   data,
   model = "A =~ a[1:5]; B =~ b[c(1,3,5)]; C =~ c1 + c2 + c3",
   estimator = "ML",
   highorder = "",
   orthogonal = FALSE,
   missing = "listwise",
   digits = 3,
   nsmall = digits,
   file = NULL
)
```

## **Arguments**

data	Data frame.
model	Model formula. See examples.
estimator	The estimator to be used (for details, see lavaan options). Default is "ML". Can be one of the following:
	"ML" Maximum Likelihood (can be extended to "MLM", "MLMV", "MLMVS", "MLF", or "MLR" for robust standard errors and robust test statistics)
	"GLS" Generalized Least Squares
	"WLS" Weighted Least Squares
	"ULS" Unweighted Least Squares
	"DWLS" Diagonally Weighted Least Squares
	"DLS" Distributionally-weighted Least Squares
highorder	High-order factor. Default is "".
orthogonal	Default is FALSE. If TRUE, all covariances among latent variables are set to zero.
missing	Default is "listwise". Alternative is "fiml" ("Full Information Maximum Likelihood").
digits, nsmall	Number of decimal places of output. Default is 3.
file	File name of MS Word (.doc).

# Value

A list of results returned by lavaan::cfa().

Corr 13

## See Also

```
Alpha, EFA, lavaan_summary
```

# **Examples**

```
data.cfa=lavaan::HolzingerSwineford1939
CFA(data.cfa, "Visual =~ x[1:3]; Textual =~ x[c(4,5,6)]; Speed =~ x7 + x8 + x9")
CFA(data.cfa, model="
    Visual =~ x[1:3]
    Textual =~ x[c(4,5,6)]
    Speed =~ x7 + x8 + x9
    ", highorder="Ability")

data.bfi = na.omit(psych::bfi)
CFA(data.bfi, "E =~ E[1:5]; A =~ A[1:5]; C =~ C[1:5]; N =~ N[1:5]; O =~ O[1:5]")
```

Corr

Correlation analysis.

# Description

Correlation analysis.

# Usage

```
Corr(
  data,
 method = "pearson",
  p.adjust = "none",
  all.as.numeric = TRUE,
  digits = 2,
  nsmall = digits,
  file = NULL,
  plot = TRUE,
  plot.range = c(-1, 1),
  plot.palette = NULL,
  plot.color.levels = 201,
  plot.file = NULL,
  plot.width = 8,
 plot.height = 6,
  plot.dpi = 500
)
```

14 Corr

#### Arguments

```
data
                  Data frame.
method
                  "pearson" (default), "spearman", or "kendall".
                  Adjustment of p values for multiple tests: "none", "fdr", "holm", "bonferroni",
p.adjust
                  ... For details, see stats::p.adjust().
all.as.numeric TRUE (default) or FALSE. Transform all variables into numeric (continuous).
digits, nsmall Number of decimal places of output. Default is 2.
file
                  File name of MS Word (.doc).
                  TRUE (default) or FALSE. Plot the correlation matrix.
plot
                  Range of correlation coefficients for plot. Default is c(-1, 1).
plot.range
plot.palette
                  Color gradient for plot. Default is c("#B52127", "white", "#2171B5"). You
                  may also set it to, e.g., c("red", "white", "blue").
plot.color.levels
                  Default is 201.
plot.file
                  NULL (default, plot in RStudio) or a file name ("xxx.png").
plot.width
                  Width (in "inch") of the saved plot. Default is 8.
plot.height
                  Height (in "inch") of the saved plot. Default is 6.
plot.dpi
                  DPI (dots per inch) of the saved plot. Default is 500.
```

#### Value

Invisibly return the correlation results obtained from psych::corr.test().

#### See Also

Describe

```
Corr(airquality)
Corr(airquality, p.adjust="bonferroni")

d = as.data.table(psych::bfi)
added(d, {
  gender = as.factor(gender)
  education = as.factor(education)
  E = .mean("E", 1:5, rev=c(1,2), range=1:6)
  A = .mean("A", 1:5, rev=1, range=1:6)
  C = .mean("C", 1:5, rev=c(4,5), range=1:6)
  N = .mean("N", 1:5, range=1:6)
  O = .mean("O", 1:5, rev=c(2,5), range=1:6)
})
Corr(d[, .(age, gender, education, E, A, C, N, O)])
```

cor\_diff

cor\_diff

Test the difference between two correlations.

# Description

Test the difference between two correlations.

# Usage

```
cor_diff(r1, n1, r2, n2, n = NULL, rcov = NULL)
```

# **Arguments**

we should also consider rcov: r(Y,Z)

#### Value

Invisibly return the *p* value.

## **Examples**

```
# two independent rs (X~Y vs. Z~W) cor_diff(r1=0.20, n1=100, r2=0.45, n2=100)  
# two nonindependent rs (X~Y vs. X~Z, with Y and Z also correlated [rcov]) cor_diff(r1=0.20, r2=0.45, n=100, rcov=0.80)
```

Describe

Descriptive statistics.

## **Description**

Descriptive statistics.

Describe Describe

## Usage

```
Describe(
  data,
  all.as.numeric = TRUE,
  digits = 2,
  nsmall = digits,
  file = NULL,
  plot = FALSE,
  upper.triangle = FALSE,
  upper.smooth = "none",
  plot.file = NULL,
  plot.width = 8,
  plot.height = 6,
  plot.dpi = 500
)
```

## **Arguments**

data	Data frame or numeric vector.
all.as.numeric	TRUE (default) or FALSE. Transform all variables into numeric (continuous).
digits, nsmall	Number of decimal places of output. Default is 2.
file	File name of MS Word (.doc).
plot	${\it TRUE or FALSE (default)}. \ Visualize the descriptive statistics using {\it GGally::ggpairs()}.$
upper.triangle	TRUE or FALSE (default). Add (scatter) plots to upper triangle (time consuming when sample size is large).
upper.smooth	"none" (default), "lm", or "loess". Add fitting lines to scatter plots (if any).
plot.file	NULL (default, plot in RStudio) or a file name ("xxx.png").
plot.width	Width (in "inch") of the saved plot. Default is 8.
plot.height	Height (in "inch") of the saved plot. Default is 6.
plot.dpi	DPI (dots per inch) of the saved plot. Default is 500.

## Value

Invisibly return a list consisting of (1) a data frame of descriptive statistics and (2) a ggplot2 object if users set plot=TRUE.

#### See Also

Corr

```
set.seed(1)
Describe(rnorm(1000000), plot=TRUE)
Describe(airquality)
```

dtime 17

```
Describe(airquality, plot=TRUE, upper.triangle=TRUE, upper.smooth="lm")
# ?psych::bfi
Describe(psych::bfi[c("age", "gender", "education")])

d = as.data.table(psych::bfi)
added(d, {
  gender = as.factor(gender)
  education = as.factor(education)
  E = .mean("E", 1:5, rev=c(1,2), range=1:6)
  A = .mean("A", 1:5, rev=1, range=1:6)
  C = .mean("C", 1:5, rev=c(4,5), range=1:6)
  N = .mean("N", 1:5, range=1:6)
  O = .mean("O", 1:5, rev=c(2,5), range=1:6)
})
Describe(d[, .(age, gender, education)], plot=TRUE, all.as.numeric=FALSE)
Describe(d[, .(age, gender, education, E, A, C, N, O)], plot=TRUE)
```

dtime

Timer (compute time difference).

## **Description**

Timer (compute time difference).

## Usage

```
dtime(t0, unit = "secs", digits = 0, nsmall = digits)
```

## Arguments

```
t0 Time at the beginning.

unit Options: "auto", "secs", "mins", "hours", "days", "weeks". Default is "secs".

digits, nsmall Number of decimal places of output. Default is 0.
```

#### Value

A character string of time difference.

```
## Not run:

t0 = Sys.time()
dtime(t0)

## End(Not run)
```

18 EFA

**EFA** 

Principal Component Analysis (PCA) and Exploratory Factor analysis (EFA).

## **Description**

An extension of psych::principal() and psych::fa(), performing either Principal Component Analysis (PCA) or Exploratory Factor Analysis (EFA).

Three options to specify variables:

- 1. var + items: use the common and unique parts of variable names.
- 2. vars: directly define a character vector of variables.
- 3. varrange: use the starting and stopping positions of variables.

## Usage

```
EFA(
  data,
  var,
  items,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  method = c("pca", "pa", "ml", "minres", "uls", "ols", "wls", "gls", "alpha"),
rotation = c("none", "varimax", "oblimin", "promax", "quartimax", "equamax"),
  nfactors = c("eigen", "parallel", "(any number >= 1)"),
  sort.loadings = TRUE,
  hide.loadings = 0,
  plot.scree = TRUE,
  kaiser = TRUE,
  max.iter = 25,
  min.eigen = 1,
  digits = 3,
  nsmall = digits,
  file = NULL
)
PCA(..., method = "pca")
```

#### **Arguments**

```
data
Data frame.

var

[Option 1] The common part across the variables. e.g., "RSES"

items

[Option 1] The unique part across the variables. e.g., 1:10

vars

[Option 2] A character vector specifying the variables. e.g., c("X1", "X2", "X3", "X4", "X5")
```

EFA 19

varrange [Option 3] A character string specifying the positions ("starting:stopping") of variables. e.g., "A1:E5"

rev [Optional] Variables that need to be reversed. It can be (1) a character vector specifying the reverse-scoring variables (recommended), or (2) a numeric vector

specifying the item number of reverse-scoring variables (not recommended).

method Extraction method.

• "pca" - Principal Component Analysis (default)

• "pa" - Principal Axis Factor Analysis

• "m1" - Maximum Likelihood Factor Analysis

• "minres" - Minimum Residual Factor Analysis

• "uls" - Unweighted Least Squares Factor Analysis

• "ols" - Ordinary Least Squares Factor Analysis

• "wls" - Weighted Least Squares Factor Analysis

• "gls" - Generalized Least Squares Factor Analysis

• "alpha" - Alpha Factor Analysis (Kaiser & Coffey, 1965)

rotation Rotation method.

• "none" - None (not suggested)

• "varimax" - Varimax (default)

• "oblimin" - Direct Oblimin

• "promax" - Promax

• "quartimax" - Quartimax

• "equamax" - Equamax

nfactors How to determine the number of factors/components?

• "eigen" - based on eigenvalue (> minimum eigenvalue) (default)

• "parallel" - based on parallel analysis

• (any number  $\geq$  1) - user-defined fixed number

sort.loadings Sort factor/component loadings by size? Default is TRUE.

hide.loadings A number  $(0\sim1)$  for hiding absolute factor/component loadings below this value.

Default is 0 (does not hide any loading).

plot.scree Display the scree plot? Default is TRUE.

kaiser Do the Kaiser normalization (as in SPSS)? Default is TRUE.

max.iter Maximum number of iterations for convergence. Default is 25 (the same as in

SPSS).

min.eigen Minimum eigenvalue (used if nfactors="eigen"). Default is 1.

digits, nsmall Number of decimal places of output. Default is 3.

file File name of MS Word (.doc).

... Arguments passed from PCA() to EFA().

20 EFA

#### Value

A list of results:

```
result The R object returned from psych::principal() or psych::fa()
result.kaiser The R object returned from psych::kaiser() (if any)
extraction.method Extraction method
rotation.method Rotation method
eigenvalues A data.frame of eigenvalues and sum of squared (SS) loadings
loadings A data.frame of factor/component loadings and communalities
scree.plot A ggplot2 object of the scree plot
```

#### **Functions**

- EFA(): Exploratory Factor Analysis
- PCA(): Principal Component Analysis a wrapper of EFA(..., method="pca")

#### Note

Results based on the varimax rotation method are identical to SPSS. The other rotation methods may produce results slightly different from SPSS.

#### See Also

```
MEAN, Alpha, CFA
```

```
data = psych::bfi
                       # var + items
EFA(data, "E", 1:5)
EFA(data, "E", 1:5, nfactors=2) # var + items
EFA(data, varrange="A1:05",
   nfactors="parallel",
   hide.loadings=0.45)
# the same as above:
# using dplyr::select() and dplyr::matches()
# to select variables whose names end with numbers
# (regexp: \d matches all numbers, $ matches the end of a string)
data %>% select(matches("\\d$")) %>%
 EFA(vars=names(.),
                         # all selected variables
     method="pca",
                          # default
     rotation="varimax", # default
     nfactors="parallel", # parallel analysis
     hide.loadings=0.45) # hide loadings < 0.45
```

**EMMEANS** 

Simple-effect analysis and post-hoc multiple comparison.

#### **Description**

Perform (1) simple-effect (and simple-simple-effect) analyses, including both simple main effects and simple interaction effects, and (2) post-hoc multiple comparisons (e.g., pairwise, sequential, polynomial), with p values adjusted for factors with >= 3 levels.

This function is based on and extends (1) emmeans::joint\_tests(), (2) emmeans::emmeans(), and (3) emmeans::contrast(). You only need to specify the model object, to-be-tested effect(s), and moderator(s). Almost all results you need will be displayed together, including effect sizes (partial  $\eta^2$  and Cohen's d) and their confidence intervals (CIs). 90% CIs for partial  $\eta^2$  and 95% CIs for Cohen's d are reported.

By default, the *root mean square error* (RMSE) is used to compute the pooled *SD* for Cohen's *d*. Specifically, it uses:

- 1. the square root of *mean square error* (MSE) for between-subjects designs;
- 2. the square root of *mean variance of all paired differences of the residuals of repeated measures* for within-subjects and mixed designs.

**Disclaimer:** There is substantial disagreement on the appropriate pooled *SD* to use in computing the effect size. For alternative methods, see emmeans::eff\_size() and effectsize::t\_to\_d(). Users should *not* take the default output as the only right results and are completely responsible for specifying sd.pooled.

#### Usage

```
EMMEANS(
  model,
  effect = NULL,
  by = NULL,
  contrast = "pairwise",
  reverse = TRUE,
  p.adjust = "bonferroni",
  sd.pooled = NULL,
  model.type = "multivariate",
  digits = 3,
  nsmall = digits
```

#### **Arguments**

model

The model object returned by MANOVA.

effect

Effect(s) you want to test. If set to a character string (e.g., "A"), it reports the results of omnibus test or simple main effect. If set to a character vector (e.g., c("A", "B")), it also reports the results of simple interaction effect.

by Moderator variable(s). Default is NULL. contrast Contrast method for multiple comparisons. Default is "pairwise". Alternatives can be "pairwise" ("revpairwise"), "seq" ("consec"), "poly", "eff". For details, see ?emmeans:: `contrast-methods`. The order of levels to be contrasted. Default is TRUE (higher level vs. lower reverse level). p.adjust Adjustment method of p values for multiple comparisons. Default is "bonferroni". For polynomial contrasts, default is "none". Alternatives can be "none", "fdr", "hochberg", "hommel", "holm", "tukey", "mvt", "dunnettx", "sidak", "scheffe", "bonferroni". For details, see stats::p.adjust() and emmeans::summary(). By default, it uses sqrt(MSE) (root mean square error, RMSE) as the pooled SDsd.pooled to compute Cohen's d. Users may specify this argument as the SD of a reference group, or use effectsize::sd\_pooled() to obtain a pooled SD. For an issue about the computation method of Cohen's d, see Disclaimer above. model.type "multivariate" returns the results of pairwise comparisons identical to SPSS, which uses the lm (rather than aov) object of the model for emmeans::joint\_tests() and emmeans::emmeans(). "univariate" requires also specifying aov.include=TRUE in MANOVA (not recommended by the afex package; for details, see afex::aov\_ez()). digits, nsmall Number of decimal places of output. Default is 3.

#### Value

The same model object as returned by MANOVA (for recursive use), along with a list of tables: sim (simple effects), emm (estimated marginal means), con (contrasts).

Each EMMEANS(...) appends one list to the returned object.

### **Interaction Plot (See Examples Below)**

You can save the returned object and use the emmeans::emmip() function to create an interaction plot (based on the fitted model and a formula). See examples below for the usage.

Note: emmeans::emmip() returns a ggplot object, which can be modified and saved with ggplot2 syntax.

#### **Statistical Details**

Some may confuse the statistical terms "simple effects", "post-hoc tests", and "multiple comparisons". Such a confusion is not uncommon. Here I explain what these terms actually refer to.

- 1. Simple Effect When we speak of "simple effect", we are referring to ...
  - simple main effect
  - simple interaction effect (only for designs with 3 or more factors)
  - simple simple effect (only for designs with 3 or more factors)

When the interaction effect in ANOVA is significant, we should then perform a "simple-effect analysis". In regression, we call this "simple-slope analysis". They are identical in statistical principles.

In a two-factors design, we only test "simple main effect". That is, at different levels of a factor "B", the main effects of "A" would be different. However, in a three-factors (or more) design, we may also test "simple interaction effect" and "simple simple effect". That is, at different combinations of levels of factors "B" and "C", the main effects of "A" would be different.

To note, simple effects  $per\ se$  never require p-value adjustment, because what we test in simple-effect analyses are still "omnibus F-tests".

2. Post-Hoc Test The term "post-hoc" means that the tests are performed after ANOVA. Given this, some may (wrongly) regard simple-effect analyses also as a kind of post-hoc tests. However, these two terms should be distinguished. In many situations, "post-hoc tests" only refer to "post-hoc comparisons" using t-tests and some p-value adjustment techniques. We need post-hoc comparisons only when there are factors with 3 or more levels.

Post-hoc tests are totally **independent of** whether there is a significant interaction effect. **It only deals with factors with multiple levels.** In most cases, we use pairwise comparisons to do post-hoc tests. See the next part for details.

**3. Multiple Comparison** As mentioned above, multiple comparisons are indeed post-hoc tests but have no relationship with simple-effect analyses. Post-hoc multiple comparisons are **independent of** interaction effects and simple effects. Furthermore, if a simple main effect contains 3 or more levels, we also need to do multiple comparisons *within* the simple-effect analysis. In this situation, we also need *p*-value adjustment with methods such as Bonferroni, Tukey's HSD (honest significant difference), FDR (false discovery rate), and so forth.

Options for multiple comparison:

- "pairwise" Pairwise comparisons (default is "higher level lower level")
- "seq" or "consec" Consecutive (sequential) comparisons
- "poly" Polynomial contrasts (linear, quadratic, cubic, quartic, ...)
- "eff" Effect contrasts (vs. the grand mean)

#### See Also

TTEST, MANOVA, bruceR-demodata

```
#### Between-Subjects Design ####
between.1
MANOVA(between.1, dv="SCORE", between="A") %>%
    EMMEANS("A")
MANOVA(between.1, dv="SCORE", between="A") %>%
    EMMEANS("A", p.adjust="tukey")
MANOVA(between.1, dv="SCORE", between="A") %>%
    EMMEANS("A", contrast="seq")
MANOVA(between.1, dv="SCORE", between="A") %>%
    EMMEANS("A", contrast="poly")
```

```
between.2
MANOVA(between.2, dv="SCORE", between=c("A", "B")) %>%
 EMMEANS("A", by="B") %>%
 EMMEANS("B", by="A")
## How to create an interaction plot using `emmeans::emmip()`?
## See help page: ?emmeans::emmip()
m = MANOVA(between.2, dv="SCORE", between=c("A", "B"))
emmip(m, \sim A \mid B, CIs=TRUE)
emmip(m, ~ B | A, CIs=TRUE)
emmip(m, B ~ A, CIs=TRUE)
emmip(m, A ~ B, CIs=TRUE)
between.3
MANOVA(between.3, dv="SCORE", between=c("A", "B", "C")) %>%
 EMMEANS("A", by="B") %>%
 EMMEANS(c("A", "B"), by="C") %>%
 EMMEANS("A", by=c("B", "C"))
## Just to name a few...
## You may test other combinations...
#### Within-Subjects Design ####
within.1
MANOVA(within.1, dvs="A1:A4", dvs.pattern="A(.)",
      within="A") %>%
 EMMEANS("A")
within.2
MANOVA(within.2, dvs="A1B1:A2B3", dvs.pattern="A(.)B(.)",
      within=c("A", "B")) %>%
 EMMEANS("A", by="B") %>%
 EMMEANS("B", by="A") # singular error matrix
# This would produce a WARNING because of
# the linear dependence of A2B2 and A2B3.
# See: Corr(within.2[c("A2B2", "A2B3")])
within.3
MANOVA(within.3, dvs="A1B1C1:A2B2C2", dvs.pattern="A(.)B(.)C(.)",
      within=c("A", "B", "C")) %>%
 EMMEANS("A", by="B") %>%
 EMMEANS(c("A", "B"), by="C") %>%
 EMMEANS("A", by=c("B", "C"))
## Just to name a few...
## You may test other combinations...
#### Mixed Design ####
mixed.2_1b1w
MANOVA(mixed.2_1b1w, dvs="B1:B3", dvs.pattern="B(.)",
      between="A", within="B", sph.correction="GG") %>%
```

export 25

```
EMMEANS("A", by="B") %>%
 EMMEANS("B", by="A")
mixed.3_1b2w
MANOVA(mixed.3_1b2w, dvs="B1C1:B2C2", dvs.pattern="B(.)C(.)",
      between="A", within=c("B", "C")) %>%
 EMMEANS("A", by="B") %>%
 EMMEANS(c("A", "B"), by="C") %>%
 EMMEANS("A", by=c("B", "C"))
## Just to name a few...
## You may test other combinations...
mixed.3_2b1w
MANOVA(mixed.3_2b1w, dvs="B1:B2", dvs.pattern="B(.)",
      between=c("A", "C"), within="B") %>%
 EMMEANS("A", by="B") %>%
 EMMEANS("A", by="C") %>%
 EMMEANS(c("A", "B"), by="C") %>%
 EMMEANS("B", by=c("A", "C"))
## Just to name a few...
## You may test other combinations...
#### Other Examples ####
air = airquality
air$Day.1or2 = ifelse(air$Day %% 2 == 1, 1, 2) %>%
 factor(levels=1:2, labels=c("odd", "even"))
MANOVA(air, dv="Temp", between=c("Month", "Day.1or2"),
      covariate=c("Solar.R", "Wind")) %>%
 EMMEANS("Month", contrast="seq") %>%
 EMMEANS("Month", by="Day.1or2", contrast="poly")
```

export

Export data to a file (TXT, CSV, Excel, SPSS, Stata, ...) or clipboard.

# Description

Export data to a file, with format automatically judged from file extension. This function is inspired by rio::export() and has several modifications. Its purpose is to avoid using lots of write\_xxx() functions in your code and to provide one tidy function for data export.

It supports many file formats and uses corresponding R functions:

- Plain text (.txt, .csv, .csv2, .tsv, .psv), using data.table::fwrite(); if the encoding argument is specified, using utils::write.table() instead
- Excel (.xls, .xlsx), using openxlsx::write.xlsx()
- SPSS (.sav), using haven::write\_sav()

26 export

```
• Stata (.dta), using haven::write_dta()
```

- R objects (.rda, .rdata, .Rdata), using base::save()
- R serialized objects (.rds), using base::saveRDS()
- Clipboard (on Windows and Mac OS), using clipr::write\_clip()
- Other formats, using rio::export()

# Usage

```
export(
   x,
   file,
   sheet = NULL,
   encoding = NULL,
   header = "auto",
   overwrite = TRUE
)
```

## Arguments

X	Any R object	t, usually a data frame	(data.frame, data.	table, tbl_df). Mul-

tiple R objects should be included in a *named* list (see examples).

If you want to save R objects other than a data frame (e.g., model results), you'd better specify file with extensions .rda, .rdata, or .Rdata.

file File name (with extension). If unspecified, then data will be exported to clip-

board.

sheet [Only for Excel] Excel sheet name(s). Default is Sheet1, Sheet2, ... You may

specify multiple sheet names in a character vector c() with the  $same\ length$  as

x (see examples).

encoding File encoding. Default is NULL. Alternatives can be "UTF-8", "GBK", "CP936",

etc.

If you find messy code for Chinese text in the exported data (often in CSV when opened with Excel), it is usually effective to set encoding="GBK" or

encoding="CP936".

header Does the first row contain column names (TRUE or FALSE)? Default is "auto".

overwrite Overwrite the existing file (if any)? Default is TRUE.

#### Value

No return value.

#### See Also

```
import, print_table
```

formatF 27

#### **Examples**

```
## Not run:
 export(airquality) # paste to clipboard
 export(airquality, file="mydata.csv")
 export(airquality, file="mydata.sav")
 export(list(airquality, npk), file="mydata.xlsx") # Sheet1, Sheet2
 export(list(air=airquality, npk=npk), file="mydata.xlsx") # a named list
 export(list(airquality, npk), sheet=c("air", "npk"), file="mydata.xlsx")
 export(list(a=1, b=npk, c="character"), file="abc.Rdata") # .rda, .rdata
 d = import("abc.Rdata") # load only the first object and rename it to `d`
 load("abc.Rdata") # load all objects with original names to environment
 export(lm(yield ~ N*P*K, data=npk), file="lm_npk.Rdata")
 model = import("lm_npk.Rdata")
 load("lm_npk.Rdata") # because x is unnamed, the object has a name "List1"
 export(list(m1=lm(yield ~ N*P*K, data=npk)), file="lm_npk.Rdata")
 model = import("lm_npk.Rdata")
 load("lm_npk.Rdata") # because x is named, the object has a name "m1"
## End(Not run)
```

formatF

Format numeric values.

## **Description**

Format numeric values.

#### Usage

```
formatF(x, digits = 3, nsmall = digits)
```

### **Arguments**

x A number or numeric vector.digits, nsmall Number of decimal places of output. Default is 3.

#### Value

Formatted character string.

#### See Also

format, formatN

28 formula\_expand

## **Examples**

```
formatF(pi, 20)
```

formatN

Format "1234" to "1,234".

# Description

```
Format "1234" to "1,234".
```

# Usage

```
formatN(x, mark = ",")
```

# Arguments

x A number or numeric vector.

mark Usually ",".

## Value

Formatted character string.

## See Also

format, formatF

# **Examples**

formatN(1234)

formula\_expand

Expand all interaction terms in a formula.

# Description

Expand all interaction terms in a formula.

# Usage

```
formula_expand(formula, as.char = FALSE)
```

formula\_paste 29

# Arguments

formula R formula or a character string indicating the formula.

as.char Return character? Default is FALSE.

## Value

A formula/character object including all expanded terms.

# **Examples**

```
formula_expand(y ~ a*b*c)
formula_expand("y ~ a*b*c")
```

formula\_paste

Paste a formula into a string.

# Description

Paste a formula into a string.

# Usage

```
formula_paste(formula)
```

# Arguments

formula

R formula.

#### Value

A character string indicating the formula.

```
formula_paste(y \sim x)
formula_paste(y \sim x + (1 | g))
```

30 Freq

Freq	Frequency statistics.
------	-----------------------

# Description

Frequency statistics.

## Usage

```
Freq(x, varname, labels, sort = "", digits = 1, nsmall = digits, file = NULL)
```

# Arguments

X	A vector of values (or a data frame).
varname	[Optional] Variable name, if x is a data frame.
labels	[Optional] A vector re-defining the labels of values.
sort	"" (default, sorted by the order of variable values/labels), "-" (decreasing by N), or "+" (increasing by N).
digits, nsmall	Number of decimal places of output. Default is 1.
file	File name of MS Word (.doc).

## Value

A data frame of frequency statistics.

```
data = psych::bfi
## Input `data$variable`
Freq(data$education)
Freq(data$gender, labels=c("Male", "Female"))
Freq(data$age)
## Input one data frame and one variable name
Freq(data, "education")
Freq(data, "gender", labels=c("Male", "Female"))
Freq(data, "age")
```

GLM\_summary 31

**GLM\_summary** 

Tidy report of GLM (1m and g1m models).

### **Description**

NOTE: model\_summary is preferred.

# Usage

```
GLM_summary(
  model,
  robust = FALSE,
  cluster = NULL,
  digits = 3,
  nsmall = digits,
  ...
)
```

# **Arguments**

model A model fitted with 1m or glm function.

robust [Only for lm and glm] FALSE (default), TRUE (then the default is "HC1"), "HC0",

"HC1", "HC2", "HC3", "HC4", "HC4m", or "HC5". It will add a table with heteroskedasticity-

robust standard errors (aka. Huber-White standard errors). For details, see

?sandwich::vcovHC and ?jtools::summ.lm.

\*\*\* "HC1" is the default of Stata, whereas "HC3" is the default suggested by the

sandwich package.

cluster [Only for 1m and g1m] Cluster-robust standard errors are computed if cluster is

set to the name of the input data's cluster variable or is a vector of clusters.

digits, nsmall Number of decimal places of output. Default is 3.

.. Other arguments. You may re-define formula, data, or family.

## Value

No return value.

## See Also

```
print_table (print simple table)
model_summary (highly suggested)
HLM_summary
regress
```

32 grand\_mean\_center

#### **Examples**

grand\_mean\_center

Grand-mean centering.

# **Description**

Compute grand-mean centered variables. Usually used for GLM interaction-term predictors and HLM level-2 predictors.

## Usage

```
grand_mean_center(data, vars = names(data), std = FALSE, add.suffix = "")
```

## Arguments

data Data object.

vars Variable(s) to be centered.

std Standardized or not. Default is FALSE.

add.suffix The suffix of the centered variable(s). Default is "". You may set it to "\_c",

"\_center", etc.

#### Value

A new data object containing the centered variable(s).

#### See Also

```
group_mean_center
```

granger\_causality 33

#### **Examples**

```
d = data.table(a=1:5, b=6:10)
d.c = grand_mean_center(d, "a")
d.c
d.c = grand_mean_center(d, c("a", "b"), add.suffix="_center")
d.c
```

granger\_causality

Granger causality test (multivariate).

## **Description**

Granger test of predictive causality (between multivariate time series) based on vector autoregression (VAR) model. Its output resembles the output of the vargranger command in Stata (but here using an *F* test).

### Usage

```
granger_causality(
  varmodel,
  var.y = NULL,
  var.x = NULL,
  test = c("F", "Chisq"),
  file = NULL,
  check.dropped = FALSE
)
```

# **Arguments**

varmodel VAR model fitted using the vars::VAR() function.

var.y, var.x [Optional] Default is NULL (all variables). If specified, then perform tests for specific variables. Values can be a single variable (e.g., "X"), a vector of variables (e.g., c("X1", "X2")), or a string containing regular expression (e.g., "X1|X2").

test F test and/or Wald  $\chi^2$  test. Default is both: c("F", "Chisq").

file File name of MS Word (.doc).

check.dropped Check dropped variables. Default is FALSE.

#### **Details**

Granger causality test (based on VAR model) examines whether the lagged values of a predictor (or predictors) help to predict an outcome when controlling for the lagged values of the outcome itself. Granger causality does not necessarily constitute a true causal effect.

34 granger\_test

#### Value

A data frame of results.

#### See Also

```
ccf_plot, granger_test
```

#### **Examples**

```
## Not run:

# R package "vars" should be installed
library(vars)
data(Canada)
VARselect(Canada)
vm = VAR(Canada, p=3)
model_summary(vm)
granger_causality(vm)

## End(Not run)
```

granger\_test

Granger causality test (bivariate).

## **Description**

Granger test of predictive causality (between two time series) using the lmtest::grangertest() function.

#### Usage

```
granger_test(formula, data, lags = 1:5, test.reverse = TRUE, file = NULL)
```

## **Arguments**

formula Model formula like y ~ x.

data Data frame.

lags Time lags. Default is 1:5.

test.reverse Whether to test reverse causality. Default is TRUE.

file File name of MS Word (.doc).

## **Details**

Granger causality test examines whether the lagged values of a predictor have an incremental role in predicting (i.e., help to predict) an outcome when controlling for the lagged values of the outcome.

Granger causality does not necessarily constitute a true causal effect.

group\_mean\_center 35

## Value

A data frame of results.

#### See Also

```
ccf_plot, granger_causality
```

#### **Examples**

```
granger_test(chicken ~ egg, data=lmtest::ChickEgg)
granger_test(chicken ~ egg, data=lmtest::ChickEgg, lags=1:10, file="Granger.doc")
unlink("Granger.doc") # delete file for code check
```

group\_mean\_center

Group-mean centering.

## **Description**

Compute group-mean centered variables. Usually used for HLM level-1 predictors.

## Usage

```
group_mean_center(
  data,
  vars = setdiff(names(data), by),
  by,
  std = FALSE,
  add.suffix = "",
  add.group.mean = "_mean"
)
```

#### Arguments

data Data object.

vars Variable(s) to be centered.

by Grouping variable.

std Standardized or not. Default is FALSE.

add.suffix The suffix of the centered variable(s). Default is "". You may set it to "\_c", "\_center", etc.

add.group.mean The suffix of the variable name(s) of group means. Default is "\_mean" (see Examples).

## Value

A new data object containing the centered variable(s).

36 HLM\_ICC\_rWG

### See Also

```
grand_mean_center
```

#### **Examples**

```
d = data.table(x=1:9, g=rep(1:3, each=3))
d.c = group_mean_center(d, "x", by="g")
d.c
d.c = group_mean_center(d, "x", by="g", add.suffix="_c")
d.c
```

HLM\_ICC\_rWG

Tidy report of HLM indices: ICC(1), ICC(2), and rWG/rWG(J).

## Description

Compute ICC(1) (non-independence of data), ICC(2) (reliability of group means), and  $r_{WG}/r_{WG(J)}$  (within-group agreement for single-item/multi-item measures) in multilevel analysis (HLM).

## Usage

```
HLM_ICC_rWG(
  data,
  group,
  icc.var,
  rwg.vars = icc.var,
  rwg.levels = 0,
  digits = 3,
  nsmall = digits
)
```

#### **Arguments**

data Data frame.

group Grouping variable.

icc.var Key variable for analysis (usually the dependent variable).

rwg.vars Default is icc.var. It can be:

• A single variable (single-item measure), then computing rWG.

• Multiple variables (multi-item measure), then computing rWG(J), where J = the number of items.

rwg.levels As  $r_{WG}/r_{WG(J)}$  compares the actual group variance to the expected random

As  $r_{WG}/r_{WG(J)}$  compares the actual group variance to the expected random variance (i.e., the variance of uniform distribution,  $\sigma_{EU}^2$ ), it is required to specify which type of uniform distribution is.

HLM\_ICC\_rWG 37

• For *continuous* uniform distribution,  $\sigma_{EU}^2=(max-min)^2/12$ . Then rwg.levels is not useful and will be set to 0 (the default).

• For *discrete* uniform distribution,  $\sigma_{EU}^2 = (A^2 - 1)/12$ , where A is the number of response options (levels). Then rwg.levels should be provided (= A in the above formula). For example, if the measure is a 5-point Likert scale, you should set rwg.levels=5.

digits, nsmall Number of decimal places of output. Default is 3.

#### **Details**

ICC(1) (intra-class correlation, or non-independence of data) ICC(1) = var.u0 / (var.u0 + var.e) =  $\sigma_{u0}^2/(\sigma_{u0}^2 + \sigma_e^2)$ 

ICC(1) is the ICC we often compute and report in multilevel analysis (usually in the Null Model, where only the random intercept of group is included). It can be interpreted as either "the proportion of variance explained by groups" (i.e., heterogeneity between groups) or "the expectation of correlation coefficient between any two observations within any group" (i.e., homogeneity within groups).

ICC(2) (reliability of group means) ICC(2) = mean(var.u0 / (var.u0 + var.e / n.k)) =  $\Sigma [\sigma_{u0}^2/(\sigma_{u0}^2 + \sigma_e^2/n_k)]/K$ 

ICC(2) is a measure of "the representativeness of group-level aggregated means for withingroup individual values" or "the degree to which an individual score can be considered a reliable assessment of a group-level construct".

 $r_{WG}/r_{WG(J)} \ \ (\text{within-group agreement for single-item/multi-item measures}) \ \ r_{WG} = 1 - \sigma^2/\sigma_{EU}^2 \\ r_{WG(J)} = 1 - (\sigma_{MJ}^2/\sigma_{EU}^2)/[J*(1-\sigma_{MJ}^2/\sigma_{EU}^2) + \sigma_{MJ}^2/\sigma_{EU}^2]$ 

 $r_{WG}/r_{WG(J)}$  is a measure of within-group agreement or consensus. Each group has an  $r_{WG}/r_{WG(J)}$ .

- \* Note for the above formulas  $\sigma_{u0}^2$ : between-group variance (i.e., tau00)
  - $\sigma_e^2$ : within-group variance (i.e., residual variance)
  - $n_k$ : group size of the k-th group
  - K: number of groups
  - $\sigma^2$ : actual group variance of the k-th group
  - $\sigma_{MJ}^2$ : mean value of actual group variance of the k-th group across all J items
  - $\sigma_{EU}^2$ : expected random variance (i.e., the variance of uniform distribution)
  - *J*: number of items

### Value

Invisibly return a list of results.

#### References

Bliese, P. D. (2000). Within-group agreement, non-independence, and reliability: Implications for data aggregation and Analysis. In K. J. Klein & S. W. Kozlowski (Eds.), *Multilevel theory, research, and methods in organizations* (pp. 349–381). San Francisco, CA: Jossey-Bass, Inc.

James, L.R., Demaree, R.G., & Wolf, G. (1984). Estimating within-group interrater reliability with and without response bias. *Journal of Applied Psychology*, 69, 85–98.

38 HLM\_summary

### See Also

R package "multilevel"

### **Examples**

**HLM\_summary** 

Tidy report of HLM (lmer and glmer models).

### Description

NOTE: model\_summary is preferred.

#### **Usage**

```
HLM_summary(model = NULL, test.rand = FALSE, digits = 3, nsmall = digits, ...)
```

## Arguments

model A model fitted with 1mer or glmer function using the 1merTest package.

test.rand [Only for lmer and glmer] TRUE or FALSE (default). Test random effects (i.e.,

variance components) by using the likelihood-ratio test (LRT), which is asymptotically chi-square distributed. For large datasets, it is much time-consuming.

digits, nsmall Number of decimal places of output. Default is 3.

... Other arguments. You may re-define formula, data, or family.

#### Value

No return value.

#### References

Hox, J. J. (2010). *Multilevel analysis: Techniques and applications* (2nd ed.). New York, NY: Routledge.

Nakagawa, S., & Schielzeth, H. (2013). A general and simple method for obtaining R<sup>2</sup> from generalized linear mixed-effects models. *Methods in Ecology and Evolution*, 4, 133–142.

Xu, R. (2003). Measuring explained variation in linear mixed effects models. *Statistics in Medicine*, 22, 3527–3541.

import 39

### See Also

```
print_table (print simple table)
model_summary (highly suggested)
GLM_summary
regress
```

#### **Examples**

```
library(lmerTest)
## Example 1: data from lme4::sleepstudy
# (1) 'Subject' is a grouping/clustering variable
# (2) 'Days' is a level-1 predictor nested within 'Subject'
# (3) No level-2 predictors
m1 = lmer(Reaction ~ (1 | Subject), data=sleepstudy)
m2 = lmer(Reaction ~ Days + (1 | Subject), data=sleepstudy)
m3 = lmer(Reaction ~ Days + (Days | Subject), data=sleepstudy)
HLM_summary(m1)
HLM_summary(m2)
HLM_summary(m3)
## Example 2: data from lmerTest::carrots
# (1) 'Consumer' is a grouping/clustering variable
# (2) 'Sweetness' is a level-1 predictor
# (3) 'Age' and 'Frequency' are level-2 predictors
hlm.1 = lmer(Preference ~ Sweetness + Age + Frequency +
               (1 | Consumer), data=carrots)
hlm.2 = lmer(Preference ~ Sweetness + Age + Frequency +
               (Sweetness | Consumer) + (1 | Product), data=carrots)
HLM_summary(hlm.1)
HLM_summary(hlm.2)
```

import

Import data from a file (TXT, CSV, Excel, SPSS, Stata, ...) or clipboard.

### **Description**

Import data from a file, with format automatically judged from file extension. This function is inspired by rio::import() and has several modifications. Its purpose is to avoid using lots of read\_xxx() functions in your code and to provide one tidy function for data import.

It supports many file formats (local or URL) and uses the corresponding R functions:

- Plain text (.txt, .csv, .csv2, .tsv, .psv), using data.table::fread()
- Excel (.xls, .xlsx), using readxl::read\_excel()
- SPSS (.sav), using foreign::read.spss(); if failed, using haven::read\_sav() instead

40 import

- Stata (.dta), using foreign::read.dta(); if failed, using haven::read\_dta() instead
- R objects (.rda, .rdata, .Rdata), using base::load()
- R serialized objects (.rds), using base::readRDS()
- Clipboard (on Windows and Mac OS), using clipr::read\_clip\_tbl()
- Other formats, using rio::import()

### Usage

```
import(
  file,
  sheet = NULL,
  range = NULL,
  encoding = NULL,
  header = "auto",
  setclass = as,
  as = "data.frame"
)
```

### **Arguments**

file	File name (with extension).	If unspecified, then data	will be imported from

clipboard.

sheet [Only for Excel] Excel sheet name (or sheet number). Default is the first sheet.

Ignored if the sheet is specified via range.

range [Only for Excel] Excel cell range. Default are all cells in a sheet. You may

specify it as range="A1:E100" or range="Sheet1!A1:E100".

encoding File encoding. Default is NULL. Alternatives can be "UTF-8", "GBK", "CP936",

etc.

If you find messy code for Chinese text in the imported data, it is usually effec-

tive to set encoding="UTF-8".

header Does the first row contain column names (TRUE or FALSE)? Default is "auto".

setclass, as Class of the imported data. Default is "data.frame". Ignored if the data file is

R object (.rds, .rda, .rdata, .Rdata).

Alternatives can be:

```
• data.frame: "data.frame", "df", "DF"
```

• data.table: "data.table", "dt", "DT"

• tbl\_df: "tibble", "tbl\_df", "tbl"

## Value

A data object (default class is data.frame).

### See Also

export

lavaan\_summary 41

## **Examples**

```
## Not run:

# Import data from system clipboard
data = import() # read from clipboard (on Windows and Mac OS)

# If you have an Excel file named "mydata.xlsx"
export(airquality, file="mydata.xlsx")

# Import data from a file
data = import("mydata.xlsx") # default: data.frame
data = import("mydata.xlsx", as="data.table")

## End(Not run)
```

lavaan\_summary

Tidy report of lavaan model.

# Description

Tidy report of lavaan model.

## Usage

```
lavaan_summary(
  lavaan,
  ci = c("raw", "boot", "bc.boot", "bca.boot"),
  nsim = 100,
  seed = NULL,
  digits = 3,
  nsmall = digits,
  print = TRUE,
  covariance = FALSE,
  file = NULL
)
```

## **Arguments**

lavaan	Model object fitted by lavaan.	
ci	Method for estimating standard error (SE) and 95% confidence interval (CI).	
	Default is "raw" (the standard approach of lavaan). Other options:	
	"boot" Percentile Bootstrap	
	"bc.boot" Bias-Corrected Percentile Bootstrap	
	"bca.boot" Bias-Corrected and Accelerated (BCa) Percentile Bootstrap	
nsim	Number of simulation samples (bootstrap resampling) for estimating SE and 95% CI. In formal analyses, nsim=1000 (or larger) is strongly suggested.	

42 lavaan\_summary

seed Random seed for obtaining reproducible results. Default is NULL.

digits, nsmall Number of decimal places of output. Default is 3.

print Print results. Default is TRUE.

covariance Print (co)variances. Default is FALSE.

file File name of MS Word (.doc).

### Value

Invisibly return a list of results:

fit Model fit indices.

measure Latent variable measures.

regression Regression paths.

covariance Variances and/or covariances.

effect Defined effect estimates.

#### See Also

PROCESS, CFA

```
## Simple Mediation:
## Solar.R (X) \Rightarrow Ozone (M) \Rightarrow Temp (Y)
# PROCESS(airquality, y="Temp", x="Solar.R",
          meds="Ozone", ci="boot", nsim=1000, seed=1)
model = "
Ozone ~ a*Solar.R
Temp ~ c.*Solar.R + b*Ozone
Indirect := a*b
Direct := c.
Total := c. + a*b
lv = lavaan::sem(model=model, data=airquality)
lavaan::summary(lv, fit.measure=TRUE, ci=TRUE, nd=3) # raw output
lavaan_summary(lv)
# lavaan_summary(lv, ci="boot", nsim=1000, seed=1)
## Serial Multiple Mediation:
## Solar.R (X) \Rightarrow Ozone (M1) \Rightarrow Wind(M2) \Rightarrow Temp (Y)
# PROCESS(airquality, y="Temp", x="Solar.R",
          meds=c("Ozone", "Wind"),
          med.type="serial", ci="boot", nsim=1000, seed=1)
model0 = "
```

LOOKUP 43

```
Ozone ~ a1*Solar.R
Wind ~ a2*Solar.R + d12*Ozone
Temp ~ c.*Solar.R + b1*Ozone + b2*Wind
Indirect_All := a1*b1 + a2*b2 + a1*d12*b2
Ind_X_M1_Y := a1*b1
Ind_X_M2_Y := a2*b2
Ind_X_M1_M2_Y := a1*d12*b2
Direct := c.
Total := c. + a1*b1 + a2*b2 + a1*d12*b2
lv0 = lavaan::sem(model=model0, data=airquality)
lavaan::summary(lv0, fit.measure=TRUE, ci=TRUE, nd=3) # raw output
lavaan_summary(1v0)
# lavaan_summary(lv0, ci="boot", nsim=1000, seed=1)
model1 = "
Ozone ~ a1*Solar.R
Wind ~ d12*Ozone
Temp ~ c.*Solar.R + b1*Ozone + b2*Wind
Indirect_All := a1*b1 + a1*d12*b2
Ind_X_M1_Y := a1*b1
Ind_X_M1_M2_Y := a1*d12*b2
Direct := c.
Total := c. + a1*b1 + a1*d12*b2
lv1 = lavaan::sem(model=model1, data=airquality)
lavaan::summary(lv1, fit.measure=TRUE, ci=TRUE, nd=3) # raw output
lavaan_summary(lv1)
# lavaan_summary(lv1, ci="boot", nsim=1000, seed=1)
```

L00KUP

Search, match, and look up values (like Excel's functions INDEX + MATCH).

### **Description**

In Excel, we can use VLOOKUP, HLOOKUP, XLOOKUP (a new function released in 2019), or the combination of INDEX and MATCH to search, match, and look up values. Here I provide a similar function.

## Usage

```
LOOKUP(
   data,
   vars,
   data.ref,
   vars.ref,
   vars.lookup,
   return = c("new.data", "new.var", "new.value")
)
```

44 LOOKUP

### **Arguments**

data	Main data.
vars	Character (vector), specifying the variable(s) to be searched in data.
data.ref	Reference data containing both the reference variable(s) and the lookup variable(s).
vars.ref	Character (vector), with the <b>same length and order</b> as vars, specifying the reference variable(s) to be matched in data.ref.
vars.lookup	Character (vector), specifying the $variable(s)$ to be looked up and returned from data.ref.
return	What to return. Default ("new.data") is to return a data frame with the lookup values added. You may also set it to "new.var" or "new.value".

### **Details**

If multiple values were simultaneously matched, a warning message would be printed.

### Value

New data object, new variable, or new value (see the argument return).

#### See Also

```
dplyr::left_join()
XLOOKUP: Excel University
```

MANOVA

Multi-factor ANOVA.

### **Description**

Multi-factor ANOVA (between-subjects, within-subjects, and mixed designs), with and without covariates (ANCOVA).

This function is based on and extends afex::aov\_ez(). You only need to specify the data, dependent variable(s), and factors (between-subjects and/or within-subjects). Almost all results you need will be displayed together, including effect sizes (partial  $\eta^2$ ) and their confidence intervals (CIs). 90% CIs for partial  $\eta^2$  (two-sided) are reported, following Steiger (2004). In addition to partial  $\eta^2$ , it also reports generalized  $\eta^2$ , following Olejnik & Algina (2003).

How to prepare your data and specify the arguments of MANOVA?

• Wide-format data (one person in one row, and repeated measures in multiple columns):

```
Betweem-subjects design MANOVA(data=, dv=, between=, ...)
Within-subjects design MANOVA(data=, dvs=, dvs.pattern=, within=, ...)
Mixed design MANOVA(data=, dvs=, dvs.pattern=, between=, within=, ...)
```

• Long-format data (one person in multiple rows, and repeated measures in one column):

```
Betweem-subjects design (not applicable)
Within-subjects design MANOVA(data=, subID=, dv=, within=, ...)
Mixed design MANOVA(data=, subID=, dv=, between=, within=, ...)
```

### Usage

```
MANOVA(
  data,
  subID = NULL,
  dv = NULL,
  dvs = NULL,
  dvs.pattern = NULL,
  between = NULL,
  within = NULL,
  covariate = NULL,
  ss.type = "III",
  sph.correction = "none",
  aov.include = FALSE,
  digits = 3,
  nsmall = digits,
  file = NULL
)
```

#### **Arguments**

Data frame. Both **wide-format** and **long-format** are supported.

subID Subject ID (the column name). Only necessary for **long-format** data.

dv Dependent variable.

• For wide-format data, dv only can be used for between-subjects designs. For within-subjects and mixed designs, please use dvs and dvs.pattern.

• For **long-format** data, dv is the outcome variable.

Repeated measures. Only for **wide-format** data (within-subjects or mixed designs).

#### Can be:

- "start:stop" to specify the range of variables (sensitive to the order of variables):
  - e.g., "A1B1: A2B3" is matched to all variables in the data between "A1B1" and "A2B3"
- a character vector to directly specify variables (insensitive to the order of variables):

e.g., c("Cond1", "Cond2", "Cond3") or cc("Cond1, Cond2, Cond3") See cc for its usage.

dvs.pattern

If you use dvs, you should also specify the pattern of variable names using regular expression.

#### Examples:

- "Cond(.)" extracts levels from "Cond1", "Cond2", "Cond3", ... You may rename the factor using the within argument (e.g., within="Condition")
- "X(..)Y(..)" extracts levels from "X01Y01", "X02Y02", "XaaYbc", ...
- "X(.+)Y(.+)" extracts levels from "X1Y1", "XaYb", "XaY002", ...

Tips on regular expression:

- "(.)" extracts any single character (number, letter, and other symbols)
- "(.+)" extracts >= 1 character(s)
- "(.\*)" extracts >= 0 character(s)
- "([0-9])" extracts any single number
- "([a-z])" extracts any single letter
- More information: Link 1 (in English) and Link 2 (in Chinese)

between

Between-subjects factor(s). Multiple variables should be included in a character vector c().

within

Within-subjects factor(s). Multiple variables should be included in a character vector c().

covariate

Covariates. Multiple variables should be included in a character vector c().

ss.type

Type of sums of squares (SS) for ANOVA. Default is "III". Possible values are "II", "III", 2, or 3.

sph.correction

[Only for repeated measures with >= 3 levels]

Sphericity correction method for adjusting the degrees of freedom (df) when the sphericity assumption is violated. Default is "none". If Mauchly's test of sphericity is significant, you may set it to "GG" (Greenhouse-Geisser) or "HF" (Huynh-Feldt).

dvs

aov.include Include the aov object in the returned object? Default is FALSE, as suggested by afex::aov\_ez() (please see the include\_aov argument in this help page, which provides a detailed explanation). If TRUE, you should also specify model.type="univariate" in EMMEANS.

digits, nsmall Number of decimal places of output. Default is 3.

file File name of MS Word (.doc).

### **Details**

If observations are not uniquely identified in user-defined long-format data, the function takes averages across those multiple observations for each case. In technical details, it specifies fun\_aggregate=mean in afex::aov\_ez() and values\_fn=mean in tidyr::pivot\_wider().

#### Value

A result object (list) returned by afex::aov\_ez(), along with several other elements: between, within, data.wide, data.long.

#### **Interaction Plot**

You can save the returned object and use the emmeans::emmip() function to create an interaction plot (based on the fitted model and a formula specification). For usage, please see the help page of emmeans::emmip(). It returns an object of class ggplot, which can be easily modified and saved using ggplot2 syntax.

### References

Olejnik, S., & Algina, J. (2003). Generalized eta and omega squared statistics: Measures of effect size for some common research designs. *Psychological Methods*, 8(4), 434–447.

Steiger, J. H. (2004). Beyond the F test: Effect size confidence intervals and tests of close fit in the analysis of variance and contrast analysis. *Psychological Methods*, 9(2), 164–182.

#### See Also

```
TTEST, EMMEANS, bruceR-demodata
```

```
#### Between-Subjects Design ####
between.1
MANOVA(between.1, dv="SCORE", between="A")
between.2
MANOVA(between.2, dv="SCORE", between=c("A", "B"))
between.3
MANOVA(between.3, dv="SCORE", between=c("A", "B", "C"))
## How to create an interaction plot using `emmeans::emmip()`?
```

```
## See help page for its usage: ?emmeans::emmip()
m = MANOVA(between.2, dv="SCORE", between=c("A", "B"))
emmip(m, ~ A | B, CIs=TRUE)
emmip(m, ~ B | A, CIs=TRUE)
emmip(m, B \sim A, CIs=TRUE)
emmip(m, A \sim B, CIs=TRUE)
#### Within-Subjects Design ####
within.1
MANOVA(within.1, dvs="A1:A4", dvs.pattern="A(.)",
       within="A")
## the same:
MANOVA(within.1, dvs=c("A1", "A2", "A3", "A4"), dvs.pattern="A(.)",
       within="MyFactor") # renamed the within-subjects factor
within.2
MANOVA(within.2, dvs="A1B1:A2B3", dvs.pattern="A(.)B(.)",
       within=c("A", "B"))
within.3
MANOVA(within.3, dvs="A1B1C1:A2B2C2", dvs.pattern="A(.)B(.)C(.)",
       within=c("A", "B", "C"))
#### Mixed Design ####
mixed.2_1b1w
MANOVA(mixed.2_1b1w, dvs="B1:B3", dvs.pattern="B(.)",
       between="A", within="B")
MANOVA(mixed.2_1b1w, dvs="B1:B3", dvs.pattern="B(.)",
       between="A", within="B", sph.correction="GG")
mixed.3_1b2w
MANOVA(mixed.3_1b2w, dvs="B1C1:B2C2", dvs.pattern="B(.)C(.)",
       between="A", within=c("B", "C"))
mixed.3_2b1w
MANOVA(mixed.3_2b1w, dvs="B1:B2", dvs.pattern="B(.)",
       between=c("A", "C"), within="B")
#### Other Examples ####
data.new = mixed.3_1b2w
names(data.new) = c("Group", "Cond_01", "Cond_02", "Cond_03", "Cond_04")
MANOVA(data.new,
       dvs="Cond_01:Cond_04",
       dvs.pattern="Cond_(..)",
       between="Group",
       within="Condition") # rename the factor
```

med\_summary 49

med\_summary

Tidy report of mediation analysis.

## Description

Tidy report of mediation analysis, which is performed using the mediation package.

### Usage

```
med_summary(model, digits = 3, nsmall = digits, file = NULL)
```

#### **Arguments**

```
model Mediation model built using mediation::mediate().

digits, nsmall Number of decimal places of output. Default is 3.

file File name of MS Word (.doc).
```

#### Value

Invisibly return a data frame containing the results.

#### See Also

**PROCESS** 

```
## Not run:
library(mediation)
# ?mediation::mediate

## Example 1: OLS Regression
## Bias-corrected and accelerated (BCa) bootstrap confidence intervals

## Hypothesis: Solar radiation -> Ozone -> Daily temperature

lm.m = lm(Ozone ~ Solar.R + Month + Wind, data=airquality)

lm.y = lm(Temp ~ Ozone + Solar.R + Month + Wind, data=airquality)

set.seed(123) # set a random seed for reproduction
```

50 model\_summary

```
med = mediate(lm.m, lm.y,
            treat="Solar.R", mediator="Ozone",
            sims=1000, boot=TRUE, boot.ci.type="bca")
med_summary(med)
## Example 2: Multilevel Linear Model (Linear Mixed Model)
## (models must be fit using "lme4::lmer" rather than "lmerTest::lmer")
## Monte Carlo simulation (quasi-Bayesian approximation)
## (bootstrap method is not applicable to "lmer" models)
## Hypothesis: Crips -> Sweetness -> Preference (for carrots)
data = lmerTest::carrots # long-format data
data = na.omit(data) # omit missing values
lmm.m = lme4::lmer(Sweetness ~ Crisp + Gender + Age + (1 | Consumer), data=data)
lmm.y = lme4::lmer(Preference \sim Sweetness + Crisp + Gender + Age + (1 | Consumer), data=data)
set.seed(123) # set a random seed for reproduction
med.lmm = mediate(lmm.m, lmm.y,
                  treat="Crisp", mediator="Sweetness",
                  sims=1000)
med_summary(med.lmm)
## End(Not run)
```

model\_summary

Tidy report of regression models.

## **Description**

Tidy report of regression models (most model types are supported). This function uses:

```
texreg::screenreg()
texreg::htmlreg()
MuMIn::std.coef()
MuMIn::r.squaredGLMM()
performance::r2_mcfadden()
performance::r2_nagelkerke()
```

## Usage

```
model_summary(
  model.list,
  std = FALSE,
  digits = 3,
  nsmall = digits,
  file = NULL,
  check = TRUE,
  zero = ifelse(std, FALSE, TRUE),
```

model\_summary 51

```
modify.se = NULL,
modify.head = NULL,
line = TRUE,
bold = 0,
...
)
```

### **Arguments**

model.list A single model or a list of (various types of) models. Most types of regression

models are supported!

std Standardized coefficients? Default is FALSE. Only applicable to linear models

and linear mixed models. Not applicable to generalized linear (mixed) models.

digits, nsmall Number of decimal places of output. Default is 3.

file File name of MS Word (.doc).

check If there is only one model in model.list, it checks for multicollinearity us-

ing performance::check\_collinearity(). You may turn it off by setting

check=FALSE.

zero Display "0" before "."? Default is TRUE.

modify.se Replace standard errors. Useful if you need to replace raw SEs with robust

SEs. New SEs should be provided as a list of numeric vectors. See usage in

texreg::screenreg().

modify.head Replace model names.

line Lines look like true line (TRUE) or === --- === (FALSE). Only relevant to R

Console output.

bold The p-value threshold below which the coefficients will be formatted in bold.

... Other arguments passed to texreg::screenreg() or texreg::htmlreg().

### Value

Invisibly return the output (character string).

### See Also

```
print_table (print simple table)
GLM_summary
HLM_summary
med_summary
lavaan_summary
PROCESS
```

52 model\_summary

```
#### Example 1: Linear Model ####
lm1 = lm(Temp ~ Month + Day, data=airquality)
lm2 = lm(Temp ~ Month + Day + Wind + Solar.R, data=airquality)
model_summary(lm1)
model_summary(1m2)
model_summary(list(lm1, lm2))
model_summary(list(lm1, lm2), std=TRUE, digits=2)
model_summary(list(lm1, lm2), file="OLS Models.doc")
unlink("OLS Models.doc") # delete file for code check
#### Example 2: Generalized Linear Model ####
glm1 = glm(case ~ age + parity,
          data=infert, family=binomial)
glm2 = glm(case ~ age + parity + education + spontaneous + induced,
          data=infert, family=binomial)
model_summary(list(glm1, glm2)) # "std" is not applicable to glm
model_summary(list(glm1, glm2), file="GLM Models.doc")
unlink("GLM Models.doc") # delete file for code check
#### Example 3: Linear Mixed Model ####
library(lmerTest)
hlm1 = lmer(Reaction ~ (1 | Subject), data=sleepstudy)
hlm2 = lmer(Reaction ~ Days + (1 | Subject), data=sleepstudy)
hlm3 = lmer(Reaction ~ Days + (Days | Subject), data=sleepstudy)
model_summary(list(hlm1, hlm2, hlm3))
model_summary(list(hlm1, hlm2, hlm3), std=TRUE)
model_summary(list(hlm1, hlm2, hlm3), file="HLM Models.doc")
unlink("HLM Models.doc") # delete file for code check
#### Example 4: Generalized Linear Mixed Model ####
library(lmerTest)
data.glmm = MASS::bacteria
glmm1 = glmer(y ~ trt + week + (1 | ID), data=data.glmm, family=binomial)
glmm2 = glmer(y ~ trt + week + hilo + (1 | ID), data=data.glmm, family=binomial)
model_summary(list(glmm1, glmm2)) # "std" is not applicable to glmm
model_summary(list(glmm1, glmm2), file="GLMM Models.doc")
unlink("GLMM Models.doc") # delete file for code check
#### Example 5: Multinomial Logistic Model ####
library(nnet)
d = airquality
d$Month = as.factor(d$Month) # Factor levels: 5, 6, 7, 8, 9
mn1 = multinom(Month ~ Temp, data=d, Hess=TRUE)
mn2 = multinom(Month ~ Temp + Wind + Ozone, data=d, Hess=TRUE)
model_summary(mn1)
model summarv(mn2)
model_summary(mn2, file="Multinomial Logistic Model.doc")
unlink("Multinomial Logistic Model.doc") # delete file for code check
```

p 53

p

Compute p value.

# Description

Compute *p* value.

# Usage

```
p(
  z = NULL,
 t = NULL,
  f = NULL
  r = NULL
  chi2 = NULL,
 n = NULL,
  df = NULL,
  df1 = NULL,
 df2 = NULL
 digits = 2,
  nsmall = digits
)
p.z(z)
p.t(t, df)
p.f(f, df1, df2)
p.r(r, n)
p.chi2(chi2, df)
```

# Arguments

```
z, t, f, r, chi2 z, t, F, r, \chi^2 value.
n, df, df1, df2 Sample size or degree of freedom.
digits, nsmall Number of decimal places of output. Default is 2.
```

# Value

p value statistics.

## **Functions**

• p.z(): Two-tailed *p* value of *z*.

54 pkg\_depend

```
• p.t(): Two-tailed p value of t.
```

- p.f(): One-tailed *p* value of *F*. (Note: *F* test is one-tailed only.)
- p.r(): Two-tailed p value of r.
- p.chi2(): One-tailed p value of  $\chi^2$ . (Note:  $\chi^2$  test is one-tailed only.)

# **Examples**

```
p.z(1.96)
p.t(2, 100)
p.f(4, 1, 100)
p.r(0.2, 100)
p.chi2(3.84, 1)
p(z=1.96)
p(t=2, df=100)
p(f=4, df1=1, df2=100)
p(r=0.2, n=100)
p(chi2=3.84, df=1)
```

pkg\_depend

Check dependencies of R packages.

## **Description**

Check dependencies of R packages.

## Usage

```
pkg_depend(pkgs, excludes = NULL)
```

### **Arguments**

pkgs Package(s).

excludes [Optional] Package(s) and their dependencies excluded from the dependencies

of pkgs. Useful if you want to see the unique dependencies of pkgs.

## Value

A character vector of package names.

## See Also

```
pkg_install_suggested
```

pkg\_install\_suggested 55

```
pkg_install_suggested Install suggested R packages.
```

# Description

Install suggested R packages.

### Usage

```
pkg_install_suggested(by)
```

## **Arguments**

by

Suggested by which package?

## Value

No return value.

### See Also

```
pkg_depend
```

## **Examples**

```
## Not run:
pkg_install_suggested() # install all packages suggested by me
## End(Not run)
```

Print

Print strings with rich formats and colors.

# Description

Be frustrated with print() and cat()? Try Print()! Run examples to see what it can do.

# Usage

```
Print(...)
Glue(...)
```

56 print\_table

# Arguments

... Character strings enclosed by "{ }" will be evaluated as R code.

Character strings enclosed by "<< >>" will be printed as formatted and colored text.

Long strings are broken by line and concatenated together.

Leading whitespace and blank lines from the first and last lines are automatically trimmed.

#### **Details**

Possible formats/colors that can be used in "<< >>" include:

- (1) bold, italic, underline, reset, blurred, inverse, hidden, strikethrough;
- (2) black, white, silver, red, green, blue, yellow, cyan, magenta;
- (3) bgBlack, bgWhite, bgRed, bgGreen, bgBlue, bgYellow, bgCyan, bgMagenta.

See more details in glue::glue() and glue::glue\_col().

### Value

Formatted text.

### **Functions**

- Print(): Paste and print strings.
- Glue(): Paste strings.

## **Examples**

print\_table

*Print a three-line table (to R Console and Microsoft Word).* 

## Description

This basic function prints any data frame as a three-line table to either R Console or Microsoft Word (.doc). It has been used in many other functions of bruceR (see below).

print\_table 57

### Usage

```
print_table(
    x,
    digits = 3,
    nsmalls = digits,
    nspaces = 1,
    row.names = TRUE,
    col.names = TRUE,
    title = "",
    note = "",
    append = "",
    line = TRUE,
    file = NULL,
    file.align.head = "auto",
    file.align.text = "auto")
```

### Arguments

x Matrix, data.frame (or data.table), or any model object (e.g., lm, glm, lmer, glmer, ...).

digits, nsmalls

Numeric vector specifying the number of decimal places of output. Default is 3.

nspaces Number of whitespaces between columns. Default is 1.

row.names, col.names

Print row/column names. Default is TRUE (column names are always printed). To modify the names, you can use a character vector with the same length as the

raw names.

title Title text, which will be inserted in (HTML code).

note Note text, which will be inserted in (HTML code).

append Other contents, which will be appended in the end (HTML code).

line Lines looks like true line (TRUE) or === --- === (FALSE).

file File name of MS Word (.doc).

file.align.head, file.align.text

Alignment of table head or table text: "left", "right", "center". Either one value of them OR a character vector of mixed values with the same length as the table columns. Default alignment (if set as "auto"): left, right, right, ..., right.

#### Value

Invisibly return a list of data frame and HTML code.

### See Also

These functions have implemented MS Word file output using this function:

• Describe

- Freq
- Corr
- EFA / PCA
- CFA
- TTEST
- MANOVA
- model\_summary
- med\_summary
- lavaan\_summary
- PROCESS
- granger\_test
- granger\_causality

## **Examples**

```
print_table(data.frame(x=1))
print_table(airquality, file="airquality.doc")
unlink("airquality.doc")  # delete file for code check

model = lm(Temp ~ Month + Day + Wind + Solar.R, data=airquality)
print_table(model)
print_table(model, file="model.doc")
unlink("model.doc")  # delete file for code check
```

**PROCESS** 

PROCESS for mediation and/or moderation analyses.

# Description

To perform mediation, moderation, and conditional process (moderated mediation) analyses, people may use software like Mplus, SPSS "PROCESS" macro, and SPSS "MLmed" macro. Some R packages can also perform such analyses separately and in a complex way, including R package "mediation", R package "interactions", and R package "lavaan". Some other R packages or scripts/modules have been further developed to improve the convenience, including jamovi module "jAMM" (by *Marcello Gallucci*, based on the lavaan package), R package "processR" (by *Keon-Woong Moon*, not official, also based on the lavaan package), and R script file "process.R" (the official PROCESS R code by *Andrew F. Hayes*, but it is not yet an R package and has some bugs and limitations).

Here, the bruceR::PROCESS() function provides an alternative to performing mediation/moderation analyses in R. This function supports a total of **24** kinds of SPSS PROCESS models (Hayes, 2018)

and also supports multilevel mediation/moderation analyses. Overall, it supports the most frequently used types of mediation, moderated moderated moderation (3-way interaction), and moderated mediation (conditional indirect effect) analyses for (generalized) linear or linear mixed models.

Specifically, the bruceR::PROCESS() function fits regression models based on the data, variable names, and a few other arguments that users input (with **no need to** specify the PROCESS model number and **no need to** manually mean-center the variables). The function can automatically judge the model number/type and also conduct grand-mean centering before model building (using the bruceR::grand\_mean\_center() function).

This automatic grand-mean centering can be turned off by setting center=FALSE.

Note that this automatic grand-mean centering (1) makes the results of main effects accurate for interpretation; (2) does not change any results of model fit (it only affects the interpretation of main effects); (3) is only conducted in "PART 1" (for an accurate estimate of main effects) but not in "PART 2" because it is more intuitive and interpretable to use the raw values of variables for the simple-slope tests in "PART 2"; (4) is not optional to users because mean-centering should always be done when there is an interaction; (5) is not conflicted with group-mean centering because after group-mean centering the grand mean of a variable will also be 0, such that the automatic grand-mean centering (with mean = 0) will not change any values of the variable.

If you need to do group-mean centering, please do this before using PROCESS. bruceR::group\_mean\_center() is a useful function of group-mean centering. Remember that the automatic grand-mean centering in PROCESS never affects the values of a group-mean centered variable, which already has a grand mean of 0.

The bruceR::PROCESS() function uses:

- 1. the interactions::sim\_slopes() function to estimate simple slopes (and conditional direct effects) in moderation, moderated moderation, and moderated mediation models (PROCESS Models 1, 2, 3, 5, 7, 8, 9, 10, 11, 12, 14, 15, 16, 17, 18, 19, 58, 59, 72, 73, 75, 76).
- 2. the mediation::mediate() function to estimate (conditional) indirect effects in (moderated) mediation models (PROCESS Models 4, 5, 7, 8, 9, 10, 11, 12, 14, 15, 16, 17, 18, 19, 58, 59, 72, 73, 75, 76).
- 3. the lavaan::sem() function to perform serial multiple mediation analysis (PROCESS Model 6).

If you use this function in your research and report its results in your paper, please cite not only bruceR but also the other R packages it uses internally (mediation, interactions, and/or lavaan).

Two parts of results are printed:

PART 1. Regression model summary (using bruceR::model\_summary() to summarize the models)

PART 2. Mediation/moderation effect estimates (using one or a combination of the above packages and functions to estimate the effects)

To organize the PART 2 output, the results of **Simple Slopes** are titled in **green**, whereas the results of **Indirect Path** are titled in **blue**.

**Disclaimer:** Although this function is named after PROCESS, Andrew F. Hayes has no role in its design, and its development is independent from the official SPSS PROCESS macro and "process.R" script. Any error or limitation should be attributed to the three R packages/functions that bruceR::PROCESS() uses internally. Moreover, as mediation analyses include *random processes* 

(i.e., bootstrap resampling or Monte Carlo simulation), the results of mediation analyses are *unlikely* to be exactly the same across different software (even if you set the same random seed in different software).

## Usage

```
PROCESS(
  data,
  y = ""
  x = "",
  meds = c(),
  mods = c(),
  covs = c(),
  clusters = c(),
  hlm.re.m = "",
  hlm.re.y = "",
  hlm.type = c("1-1-1", "2-1-1", "2-2-1"),
  med.type = c("parallel", "serial"),
  mod.type = c("2-way", "3-way"),
  mod.path = c("x-y", "x-m", "m-y", "all"),
cov.path = c("y", "m", "both"),
  mod1.val = NULL,
  mod2.val = NULL,
  ci = c("boot", "bc.boot", "bca.boot", "mcmc"),
  nsim = 100,
  seed = NULL
  center = TRUE,
  std = FALSE,
  digits = 3,
  nsmall = digits,
  file = NULL
```

## **Arguments**

data	Data frame.
y, x	Variable name of outcome (Y) and predictor (X).
	It supports both continuous (numeric) and dichotomous (factor) variables.
meds	Variable name(s) of mediator(s) (M). Use c() to combine multiple mediators.
	It supports both continuous (numeric) and dichotomous (factor) variables.
	It allows an infinite number of mediators in parallel or 2~4 mediators in serial.
	* Order matters when med.type="serial" (PROCESS Model 6: serial mediation).
mods	Variable name(s) of 0~2 moderator(s) (W). Use c() to combine multiple moderators.
	It supports all types of variables: continuous (numeric), dichotomous (factor),

and multicategorical (factor).

\* Order matters when mod.type="3-way" (PROCESS Models 3, 5.3, 11, 12, 18, 19, 72, and 73).

\*\* Do not set this argument when med.type="serial" (PROCESS Model 6).

covs

Variable name(s) of covariate(s) (i.e., control variables). Use c() to combine multiple covariates. It supports all types of (and an infinite number of) variables.

clusters

HLM (multilevel) cluster(s): e.g., "School", c("Prov", "City"), c("Sub", "Item").

hlm.re.m, hlm.re.y

HLM (multilevel) random effect term of M model and Y model. By default, it converts clusters to lme4 syntax of random intercepts: e.g., "(1 | School)" or "(1 | Sub) + (1 | Item)".

You may specify these arguments to include more complex terms: e.g., random slopes "(X | School)", or 3-level random effects "(1 | Prov/City)".

hlm.type

HLM (multilevel) mediation type (levels of "X-M-Y"): "1-1-1" (default), "2-1-1" (indeed the same as "1-1-1" in a mixed model), or "2-2-1" (currently *not fully supported*, as limited by the mediation package). In most cases, no need to set this argument.

med.type

Type of mediator: "parallel" (default) or "serial" (only relevant to PRO-CESS Model 6). Partial matches of "p" or "s" also work. In most cases, no need to set this argument.

mod.type

Type of moderator: "2-way" (default) or "3-way" (relevant to PROCESS Models 3, 5.3, 11, 12, 18, 19, 72, and 73). Partial matches of "2" or "3" also work.

mod.path

Which path(s) do the moderator(s) influence? "x-y", "x-m", "m-y", or any combination of them (use c() to combine), or "all" (i.e., all of them). No default value.

cov.path

Which path(s) do the control variable(s) influence? "y", "m", or "both" (default).

mod1.val, mod2.val

By default (NULL), it uses **Mean +/- SD** of a continuous moderator (numeric) or **all levels** of a dichotomous/multicategorical moderator (factor) to perform simple slope analyses and/or conditional mediation analyses. You may manually specify a vector of certain values: e.g., mod1.val=c(1, 3, 5) or mod1.val=c("A", "B", "C").

ci

Method for estimating the standard error (SE) and 95% confidence interval (CI) of indirect effect(s). Default is "boot" for (generalized) linear models or "mcmc" for (generalized) linear mixed models (i.e., multilevel models).

"boot" Percentile Bootstrap

"bc.boot" Bias-Corrected Percentile Bootstrap

"bca.boot" Bias-Corrected and Accelerated (BCa) Percentile Bootstrap

"mcmc" Markov Chain Monte Carlo (Quasi-Bayesian)

\* Note that these methods *never* apply to the estimates of simple slopes. You *should not* report the 95% CIs of simple slopes as Bootstrap or Monte Carlo CIs, because they are just standard CIs without any resampling method.

Number of simulation samples (bootstrap resampling or Monte Carlo simulation) for estimating SE and 95% CI. Default is 100 for running examples faster. In formal analyses, however, nsim=1000 (or larger) is strongly suggested!

Random seed for obtaining reproducible results. Default is NULL. You may set

to any number you prefer (e.g., seed=1234, just an uncountable number).

\* Note that all mediation models include random processes (i.e., bootstrap resampling or Monte Carlo simulation). To get exactly the same results between runs, you need to set a random seed. However, even if you set the same seed number, it is unlikely to get exactly the same results across different R packages (e.g., lavaan vs. mediation) and software (e.g., SPSS, Mplus, R, jamovi).

center Centering numeric (continuous) predictors? Default is TRUE (suggested).

std Standardizing variables to get standardized coefficients? Default is FALSE. If

TRUE, it will standardize all numeric (continuous) variables before building regression models. However, it is *not suggested* to set std=TRUE for *generalized* 

linear (mixed) models.

digits, nsmall Number of decimal places of output. Default is 3.

file File name of MS Word (.doc). Currently, only regression model summary can

be saved.

#### **Details**

seed

For more details and illustrations, see PROCESS-bruceR-SPSS (PDF and Markdown files).

## Value

Invisibly return a list of results:

process.id PROCESS model number.

process.type PROCESS model type.

model.m "Mediator" (M) models (a list of multiple models).

model.y "Outcome" (Y) model.

results Effect estimates and other results (unnamed list object).

#### References

Hayes, A. F. (2018). *Introduction to mediation, moderation, and conditional process analysis (second edition): A regression-based approach.* Guilford Press.

Yzerbyt, V., Muller, D., Batailler, C., & Judd, C. M. (2018). New recommendations for testing indirect effects in mediational models: The need to report and test component paths. *Journal of Personality and Social Psychology*, 115(6), 929–943.

#### See Also

```
lavaan_summary
model_summary
med_summary
```

```
#### NOTE ####
## In the following examples, I set nsim=100 to save time.
## In formal analyses, nsim=1000 (or larger) is suggested!
#### Demo Data ####
# ?mediation::student
data = mediation::student %>%
  dplyr::select(SCH_ID, free, smorale, pared, income,
                gender, work, attachment, fight, late, score)
names(data)[2:3] = c("SCH_free", "SCH_morale")
names(data)[4:7] = c("parent_edu", "family_inc", "gender", "partjob")
datagender01 = 1 - data gender # 0 = female, 1 = male
# dichotomous X: as.factor()
data$gender = factor(data$gender01, levels=0:1, labels=c("Female", "Male"))
# dichotomous Y: as.factor()
data$pass = as.factor(ifelse(data$score>=50, 1, 0))
#### Descriptive Statistics and Correlation Analyses ####
Freq(data$gender)
Freq(data$pass)
Describe(data)
                  # file="xxx.doc"
Corr(data[,4:11]) # file="xxx.doc"
#### PROCESS Analyses ####
## Model 1 ##
PROCESS(data, y="score", x="late", mods="gender") # continuous Y
PROCESS(data, y="pass", x="late", mods="gender") # dichotomous Y
# (multilevel moderation)
PROCESS(data, y="score", x="late", mods="gender", # continuous Y (LMM)
        clusters="SCH_ID")
PROCESS(data, y="pass", x="late", mods="gender", # dichotomous Y (GLMM)
        clusters="SCH_ID")
# (Johnson-Neyman (J-N) interval and plot)
PROCESS(data, y="score", x="gender", mods="late") -> P
P$results[[1]]$jn[[1]]  # Johnson-Neyman interval
P$results[[1]]$jn[[1]]$plot # Johnson-Neyman plot (ggplot object)
                        # detailed results of regression
GLM_summary(P$model.y)
# (allows multicategorical moderator)
d = airquality
d$Month = as.factor(d$Month) # moderator: factor with levels "5"~"9"
PROCESS(d, y="Temp", x="Solar.R", mods="Month")
## Model 2 ##
PROCESS(data, y="score", x="late",
        mods=c("gender", "family_inc"),
        mod.type="2-way") # or omit "mod.type", default is "2-way"
```

64 RECODE

```
## Model 3 ##
PROCESS(data, y="score", x="late",
        mods=c("gender", "family_inc"),
        mod.type="3-way")
PROCESS(data, y="pass", x="gender", mods=c("late", "family_inc"),
        mod1.val=c(1, 3, 5),  # moderator 1: late
        mod2.val=seq(1, 15, 2), # moderator 2: family_inc
        mod.type="3-way")
## Model 4 ##
PROCESS(data, y="score", x="parent_edu",
        meds="family_inc", covs="gender",
        ci="boot", nsim=100, seed=1)
# (allows an infinite number of multiple mediators in parallel)
PROCESS(data, y="score", x="parent_edu",
        meds=c("family_inc", "late"),
        covs=c("gender", "partjob"),
        ci="boot", nsim=100, seed=1)
# (multilevel mediation)
PROCESS(data, y="score", x="SCH_free",
        meds="late", clusters="SCH_ID",
        ci="mcmc", nsim=100, seed=1)
## Model 6 ##
PROCESS(data, y="score", x="parent_edu",
        meds=c("family_inc", "late"),
        covs=c("gender", "partjob"),
        med.type="serial",
        ci="boot", nsim=100, seed=1)
## Model 8 ##
PROCESS(data, y="score", x="fight",
        meds="late",
        mods="gender",
        mod.path=c("x-m", "x-y"),
        ci="boot", nsim=100, seed=1)
## For more examples and details, see the "note" subfolder at:
## https://github.com/psychbruce/bruceR/tree/main/note
```

RECODE

Recode a variable.

## **Description**

A wrapper of car::recode().

regress 65

## Usage

```
RECODE(var, recodes)
```

# Arguments

var Variable (numeric, character, or factor).

recodes A character string definine the rule of recoding. e.g., "lo:1=0; c(2,3)=1; 4=2; 5:hi=3; else=999"

## Value

A vector of recoded variable.

# **Examples**

```
d = data.table(var=c(NA, 0, 1, 2, 3, 4, 5, 6))
added(d, {
  var.new = RECODE(var, "lo:1=0; c(2,3)=1; 4=2; 5:hi=3; else=999")
})
d
```

regress

Regression analysis.

# Description

NOTE: model\_summary is preferred.

# Usage

```
regress(
  formula,
  data,
  family = NULL,
  digits = 3,
  nsmall = digits,
  robust = FALSE,
  cluster = NULL,
  test.rand = FALSE
)
```

66 regress

### **Arguments**

formula Model formula. data Data frame.

family [Optional] The same as in glm and glmer (e.g., family=binomial fits a logistic

regression model).

digits, nsmall Number of decimal places of output. Default is 3.

robust [Only for lm and glm] FALSE (default), TRUE (then the default is "HC1"), "HC0",

"HC1", "HC2", "HC3", "HC4", "HC4m", or "HC5". It will add a table with heteroskedasticity-

robust standard errors (aka. Huber-White standard errors). For details, see

?sandwich::vcovHC and ?jtools::summ.lm.

\*\*\* "HC1" is the default of Stata, whereas "HC3" is the default suggested by the

sandwich package.

cluster [Only for 1m and g1m] Cluster-robust standard errors are computed if cluster is

set to the name of the input data's cluster variable or is a vector of clusters.

test.rand [Only for lmer and glmer] TRUE or FALSE (default). Test random effects (i.e.,

variance components) by using the likelihood-ratio test (LRT), which is asymptotically chi-square distributed. For large datasets, it is much time-consuming.

#### Value

No return value.

## See Also

```
print_table (print simple table)
model_summary (highly suggested)
GLM_summary
HLM_summary
```

rep\_char 67

```
library(lmerTest)
data.glmm = MASS::bacteria
regress(y ~ trt + week + (1 | ID), data=data.glmm, family=binomial)
regress(y ~ trt + week + hilo + (1 | ID), data=data.glmm, family=binomial)
## End(Not run)
```

rep\_char

Repeat a character string for many times and paste them up.

## **Description**

Repeat a character string for many times and paste them up.

# Usage

```
rep_char(char, rep.times)
```

# Arguments

char Character string.
rep.times Times for repeat.

## Value

Character string.

## **Examples**

```
rep_char("a", 5)
```

**RESCALE** 

Rescale a variable (e.g., from 5-point to 7-point).

## **Description**

Rescale a variable (e.g., from 5-point to 7-point).

## Usage

```
RESCALE(var, from = range(var, na.rm = T), to)
```

68 *RGB* 

## **Arguments**

var Variable (numeric).

from Numeric vector, the range of old scale (e.g., 1:5). If not defined, it will compute

the range of var.

to Numeric vector, the range of new scale (e.g., 1:7).

### Value

A vector of rescaled variable.

### **Examples**

```
d = data.table(var=rep(1:5, 2))
added(d, {
  var1 = RESCALE(var, to=1:7)
  var2 = RESCALE(var, from=1:5, to=1:7)
})
d # var1 is equal to var2
```

RGB

A simple extension of rgb().

## Description

A simple extension of rgb().

### Usage

```
RGB(r, g, b, alpha)
```

### **Arguments**

r, g, b Red, Green, Blue: 0~255.

alpha Color transparency (opacity): 0~1. If not specified, an opaque color will be

generated.

## Value

```
"#rrggbb" or "#rrggbbaa".
```

```
RGB(255, 0, 0) # red: "#FF0000"
RGB(255, 0, 0, 0.8) # red with 80\% opacity: "#FF0000CC"
```

Run 69

Run

Run code parsed from text.

# Description

Run code parsed from text.

# Usage

```
Run(..., silent = FALSE)
```

### **Arguments**

... Character string(s) to run. You can use "{ }" to insert any R object in the environment.

silent Suppress error/warning messages. Default is FALSE.

### Value

Invisibly return the running expression(s).

## **Examples**

```
Run("a=1", "b=2")
Run("print({a+b})")
```

scaler

Min-max scaling (min-max normalization).

## **Description**

This function resembles RESCALE() and it is just equivalent to RESCALE(var, to=0:1).

# Usage

```
scaler(v, min = 0, max = 1)
```

#### **Arguments**

```
Variable (numeric vector).min Minimum value (default is 0).max Maximum value (default is 1).
```

70 set.wd

### Value

A vector of rescaled variable.

# **Examples**

```
scaler(1:5)
# the same: RESCALE(1:5, to=0:1)
```

set.wd

Set working directory to the path of currently opened file.

## **Description**

Set working directory to the path of currently opened file (usually an R script). You can use this function in both **.R/.Rmd files and R Console**. **RStudio** (version >= 1.2) is required for running this function.

# Usage

```
set.wd(path = NULL, ask = FALSE)
set_wd(path = NULL, ask = FALSE)
```

# Arguments

path NULL (default) or a specific path. Default is to extract the path of the currently

 $opened\ file\ (usually\ .R\ or\ .Rmd)\ using\ the\ \verb"rstudioapi::getSourceEditorContext"$ 

function.

ask TRUE or FALSE (default). If TRUE, you can select a folder with the prompt of a

dialog.

#### Value

Invisibly return the path.

### **Functions**

```
• set.wd(): Main function
```

• set\_wd(): The alias of set.wd (the same)

#### See Also

setwd

show\_colors 71

### **Examples**

```
## Not run:

# RStudio (version >= 1.2) is required for running this function.
set.wd() # set working directory to the path of the currently opened file
set.wd("~/") # set working directory to the home path
set.wd("../") # set working directory to the parent path
set.wd(ask=TRUE) # select a folder with the prompt of a dialog

## End(Not run)
```

show\_colors

Show colors.

### **Description**

Show colors.

### Usage

```
show_colors(colors)
```

## Arguments

colors

Color names.

e.g.,

- "red" (R base color names)
- "#FF0000" (hex color names)
- see::social\_colors()
- viridis::viridis\_pal()(10)
- RColorBrewer::brewer.pal(name="Set1", n=9)
- RColorBrewer::brewer.pal(name="Set2", n=8)
- RColorBrewer::brewer.pal(name="Spectral", n=11)

### Value

A gg object.

```
show_colors("blue")
show_colors("#0000FF") # blue (hex name)
show_colors(RGB(0, 0, 255)) # blue (RGB)
show_colors(see::social_colors())
show_colors(see::pizza_colors())
```

72 theme\_bruce

theme\_bruce

A nice ggplot2 theme that enables Markdown/HTML rich text.

### **Description**

A nice ggplot2 theme for scientific publication. It uses ggtext::element\_markdown() to render Markdown/HTML formatted rich text. You can use a combination of Markdown and/or HTML syntax (e.g., "\*y\* = \*x\*<sup>2</sup>") in plot text or title, and this function draws text elements with rich text format.

For more usage, see:

```
ggtext::geom_richtext()ggtext::geom_textbox()ggtext::element_markdown()ggtext::element_textbox()
```

### Usage

```
theme_bruce(
 markdown = FALSE,
 base.size = 12,
  line.size = 0.5,
  border = "black",
  bg = "white",
  panel.bg = "white",
  tag = "bold",
  plot.title = "bold",
  axis.title = "plain",
  title.pos = 0.5,
  subtitle.pos = 0.5,
  caption.pos = 1,
  font = NULL,
  grid.x = "",
 grid.y = "",
  line.x = TRUE,
  line.y = TRUE,
  tick.x = TRUE,
  tick.y = TRUE
)
```

## Arguments

markdown Use element\_markdown() instead of element\_text(). Default is FALSE. If set to TRUE, then you should also use element\_markdown() in theme() (if any).

base.size Basic font size. Default is 12.

theme\_bruce 73

```
line.size
                  Line width. Default is 0.5.
                  TRUE, FALSE, or "black" (default).
border
                  Background color of whole plot. Default is "white". You can use any colors or
bg
                  choose from some pre-set color palettes: "stata", "stata.grey", "solar",
                  "wsj", "light", "dust".
                  To see these colors, you can type:
                  ggthemr::colour_plot(c(stata="#EAF2F3", stata.grey="#E8E8E8",solar="#FDF6E3",
                  wsj="#F8F2E4", light="#F6F1EB", dust="#FAF7F2"))
panel.bg
                  Background color of panel. Default is "white".
                  Font face of tag. Choose from "plain", "italic", "bold", "bold.italic".
tag
                  Font face of title. Choose from "plain", "italic", "bold", "bold.italic".
plot.title
                  Font face of axis text. Choose from "plain", "italic", "bold", "bold.italic".
axis.title
title.pos
                  Title position (0\sim1).
subtitle.pos
                  Subtitle position (0\sim1).
caption.pos
                  Caption position (0\sim1).
font
                  Text font. Only applicable to Windows system.
                  FALSE, "" (default), or a color (e.g., "grey90") to set the color of panel grid (x).
grid.x
grid.y
                  FALSE, "" (default), or a color (e.g., "grey90") to set the color of panel grid (y).
                  Draw the x-axis line. Default is TRUE.
line.x
                  Draw the y-axis line. Default is TRUE.
line.y
tick.x
                  Draw the x-axis ticks. Default is TRUE.
tick.y
                  Draw the y-axis ticks. Default is TRUE.
```

#### Value

A theme object that should be used for ggplot2.

74 TTEST

**TTEST** 

*One-sample, independent-samples, and paired-samples t-test.* 

#### **Description**

One-sample, independent-samples, and paired-samples t-test, with both Frequentist and Bayesian approaches. The output includes descriptives, t statistics, mean difference with 95% CI, Cohen's d with 95% CI, and Bayes factor (BF10; BayesFactor package needs to be installed). It also tests the assumption of homogeneity of variance and allows users to determine whether variances are equal or not.

Users can simultaneously test multiple dependent and/or independent variables. The results of one pair of Y-X would be summarized in one row in the output. Key results can be saved in APA format to MS Word.

#### Usage

```
TTEST(
  data,
  у,
  x = NULL
  paired = FALSE,
  paired.d.type = "dz",
  var.equal = TRUE,
  mean.diff = TRUE,
  test.value = 0,
  test.sided = c("=", "<", ">"),
  factor.rev = TRUE,
  bayes.prior = "medium",
  digits = 2,
  nsmall = digits,
  file = NULL
)
```

TTEST 75

#### **Arguments**

data	Data frame (wide-format only, i.e., one case in one row).
У	Dependent variable(s). Multiple variables should be included in a character vector c().
	For paired-samples <i>t</i> -test, the number of variables should be 2, 4, 6, etc.
х	Independent variable(s). Multiple variables should be included in a character vector c().
	Only necessary for independent-samples <i>t</i> -test.
paired	For paired-samples <i>t</i> -test, set it as TRUE. Default is FALSE.
paired.d.type	Type of Cohen's <i>d</i> for paired-samples <i>t</i> -test (see Lakens, 2013).
	Default is "dz". Options include:
	"dz" ( <i>d</i> for standardized difference) Cohen's $d_z = \frac{M_{diff}}{SD_{diff}}$
	"dav" ( <i>d</i> for average standard deviation) Cohen's $d_{av} = \frac{M_{diff}}{\frac{SD_1 + SD_2}{2}}$
	"drm" (d for repeated measures, corrected for correlation) Cohen's $d_{rm}=$
	$\frac{{M_{diff}} \times \sqrt{2(1 - r_{1,2})}}{\sqrt{SD_1^2 + SD_2^2 - 2 \times r_{1,2} \times SD_1 \times SD_2}}$
var.equal	If Levene's test indicates a violation of the homogeneity of variance, then you should better set this argument as FALSE. Default is TRUE.
mean.diff	Whether to display results of mean difference and its 95% CI. Default is TRUE.
test.value	The true value of the mean (or difference in means for a two-samples test). Default is $\emptyset$ .
test.sided	Any of "=" (two-sided, the default), "<" (one-sided), or ">" (one-sided).
factor.rev	Whether to reverse the levels of factor (X) such that the test compares higher vs. lower level. Default is TRUE.
bayes.prior	Prior scale in Bayesian <i>t</i> -test. Default is 0.707. See details in BayesFactor::ttestBF().
digits, nsmall	Number of decimal places of output. Default is 2.
file	File name of MS Word (.doc).

#### **Details**

Note that the point estimate of Cohen's d is computed using the common method "Cohen's d = mean difference / (pooled) standard deviation", which is consistent with results from other R packages (e.g., effectsize) and software (e.g., jamovi). The 95% CI of Cohen's d is estimated based on the 95% CI of mean difference (i.e., also divided by the pooled standard deviation).

However, different packages and software diverge greatly on the estimate of the 95% CI of Cohen's d. R packages such as psych and effectsize, R software jamovi, and several online statistical tools for estimating effect sizes indeed produce surprisingly inconsistent results on the 95% CI of Cohen's d.

See an illustration of this issue in the section "Examples".

#### References

Lakens, D. (2013). Calculating and reporting effect sizes to facilitate cumulative science: A practical primer for *t*-tests and ANOVAs. *Frontiers in Psychology, 4*, Article 863.

76 TTEST

#### See Also

MANOVA, EMMEANS

```
## Demo data ##
d1 = between.3
d1$Y1 = d1$SCORE # shorter name for convenience
d1$Y2 = rnorm(32) # random variable
d1$B = factor(d1$B, levels=1:2, labels=c("Low", "High"))
d1$C = factor(d1$C, levels=1:2, labels=c("M", "F"))
d2 = within.1
## One-sample t-test ##
TTEST(d1, "SCORE")
TTEST(d1, "SCORE", test.value=5)
## Independent-samples t-test ##
TTEST(d1, "SCORE", x="A")
TTEST(d1, "SCORE", x="A", var.equal=FALSE)
TTEST(d1, y="Y1", x=c("A", "B", "C"))
TTEST(d1, y=c("Y1", "Y2"), x=c("A", "B", "C"),
      mean.diff=FALSE, # remove to save space
      file="t-result.doc")
unlink("t-result.doc") # delete file for code check
## Paired-samples t-test ##
TTEST(d2, y=c("A1", "A2"), paired=TRUE)
TTEST(d2, y=c("A1", "A2", "A3", "A4"), paired=TRUE)
## Not run:
  ## Illustration for the issue stated in "Details"
  # Inconsistency in the 95% CI of Cohen's d between R packages:
  # In this example, the true point estimate of Cohen's d = 3.00
  # and its 95% CI should be equal to 95% CI of mean difference.
  data = data.frame(X=rep(1:2, each=3), Y=1:6)
  data # simple demo data
  TTEST(data, y="Y", x="X")
  \# d = 3.00 [0.73, 5.27] (estimated based on 95% CI of mean difference)
  MANOVA(data, dv="Y", between="X") %>%
   EMMEANS("X")
  \# d = 3.00 [0.73, 5.27] (the same as TTEST)
  psych::cohen.d(x=data, group="X")
  \# d = 3.67 [0.04, 7.35] (strange)
```

%allin% 77

```
psych::d.ci(d=3.00, n1=3, n2=3)
 \# d = 3.00 [-0.15, 6.12] (significance inconsistent with t-test)
 # jamovi uses psych::d.ci() to compute 95% CI
 # so its results are also: 3.00 [-0.15, 6.12]
 effectsize::cohens_d(Y ~ rev(X), data=data)
 \# d = 3.00 [0.38, 5.50] (using the noncentrality parameter method)
 effectsize::t_to_d(t=t.test(Y ~ rev(X), data=data, var.equal=TRUE)$statistic,
                     df_error=4)
 # d = 3.67 [0.47, 6.74] (merely an approximate estimate, often overestimated)
 # see ?effectsize::t_to_d
 # https://www.psychometrica.de/effect_size.html
 # d = 3.00 [0.67, 5.33] (slightly different from TTEST)
 # https://www.campbellcollaboration.org/escalc/
 \# d = 3.00 [0.67, 5.33] (slightly different from TTEST)
 # Conclusion:
 # TTEST() provides a reasonable estimate of Cohen's d and its 95% CI,
 \# and effectsize::cohens_d() offers another method to compute the CI.
## End(Not run)
```

%allin%

A simple extension of %in%.

## **Description**

A simple extension of %in%.

#### Usage

x %allin% vector

#### **Arguments**

x Numeric or character vector.

vector Numeric or character vector.

#### Value

TRUE or FALSE.

#### See Also

%in%, %anyin%, %nonein%, %partin%

78 %anyin%

# Examples

```
1:2 %allin% 1:3 # TRUE
3:4 %allin% 1:3 # FALSE
```

%anyin%

A simple extension of %in%.

# Description

A simple extension of %in%.

## Usage

x %anyin% vector

# Arguments

x Numeric or character vector.

vector Numeric or character vector.

## Value

TRUE or FALSE.

## See Also

```
%in%, %allin%, %nonein%, %partin%
```

```
3:4 %anyin% 1:3 # TRUE
4:5 %anyin% 1:3 # FALSE
```

%%COMPUTE%% 79

%%COMPUTE%%

Multivariate computation.

#### **Description**

Easily compute multivariate sum, mean, and other scores. Reverse scoring can also be easily implemented without saving extra variables. Alpha function uses a similar method to deal with reverse scoring.

Three options to specify variables:

- 1. var + items: common and unique parts of variable names (suggested).
- 2. vars: a character vector of variable names (suggested).
- 3. varrange: starting and stopping positions of variables (NOT suggested).

#### Usage

```
COUNT(data, var = NULL, items = NULL, vars = NULL, varrange = NULL, value = NA)
MODE(data, var = NULL, items = NULL, vars = NULL, varrange = NULL)
SUM(
  data,
  var = NULL,
  items = NULL,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  range = likert,
  likert = NULL,
  na.rm = TRUE
)
.sum(
  var = NULL,
  items = NULL,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  range = likert,
  likert = NULL,
  na.rm = TRUE
)
MEAN(
  data,
  var = NULL,
```

80 %%COMPUTE%%

```
items = NULL,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  range = likert,
  likert = NULL,
  na.rm = TRUE
)
.mean(
  var = NULL,
  items = NULL,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  range = likert,
  likert = NULL,
  na.rm = TRUE
)
STD(
  data,
  var = NULL,
  items = NULL,
  vars = NULL,
  varrange = NULL,
  rev = NULL,
  range = likert,
  likert = NULL,
  na.rm = TRUE
)
CONSEC(
  data,
  var = NULL,
  items = NULL,
  vars = NULL,
  varrange = NULL,
  values = 0:9
)
```

# Arguments

data	Data frame.
var	[Option 1] The common part across the variables. e.g., "RSES"
items	[Option 1] The unique part across the variables. e.g., 1:10
vars	[Option 2] A character vector specifying the variables. e.g., $c("X1", "X2", "X3", "X4", "X5")$

%%COMPUTE%% 81

varrange	[Option 3] A character string specifying the positions ("starting:stopping") of variables. e.g., "A1:E5"
value	[Only for COUNT] The value to be counted.
rev	[Optional] Variables that need to be reversed. It can be (1) a character vector specifying the reverse-scoring variables (recommended), or (2) a numeric vector specifying the item number of reverse-scoring variables (not recommended).
range, likert	[Optional] Range of likert scale (e.g., 1:5, c(1, 5)). If not provided, it will be automatically estimated from the given data (BUT you should use this carefully).
na.rm	Ignore missing values. Default is TRUE.
values	[Only for CONSEC] Values to be counted as consecutive identical values. Default is all numbers $(0:9)$ .

#### Value

A vector of computed values.

#### **Functions**

- COUNT(): Count a certain value across variables.
- MODE(): Compute mode across variables.
- SUM(): Compute sum across variables.
- .sum(): Tidy version of SUM, only can be used in add()/added()
- MEAN(): Compute **mean** across variables.
- .mean(): Tidy version of MEAN, only can be used in add()/added()
- STD(): Compute **standard deviation** across variables.
- CONSEC(): Compute **consecutive identical digits** across variables (especially useful in detecting careless responding).

82 %nonein%

```
cons1 = CONSEC(d, "x", 1:5),
  cons2 = CONSEC(d, varrange="x1:x5")
)]
d

## ====== Usage 2: `add()` & `added()` ===== ##
data = as.data.table(psych::bfi)
added(data, {
  gender = as.factor(gender)
  education = as.factor(education)
  E = .mean("E", 1:5, rev=c(1,2), range=1:6)
  A = .mean("A", 1:5, rev=1, range=1:6)
  C = .mean("C", 1:5, rev=c(4,5), range=1:6)
  N = .mean("N", 1:5, range=1:6)
  O = .mean("O", 1:5, rev=c(2,5), range=1:6)
}, drop=TRUE)
data
```

%nonein%

A simple extension of %in%.

## Description

A simple extension of %in%.

## Usage

x %nonein% vector

#### **Arguments**

x Numeric or character vector.vector Numeric or character vector.

# Value

TRUE or FALSE.

## See Also

%in%, %allin%, %anyin%, %partin%

```
3:4 %nonein% 1:3 # FALSE
4:5 %nonein% 1:3 # TRUE
```

%notin% 83

%notin%

The opposite of %in%.

## Description

The opposite of %in%.

#### Usage

```
x %notin% vector
```

#### **Arguments**

x Numeric or character vector.vector Numeric or character vector.

#### Value

A vector of TRUE or FALSE.

## See Also

%in%

# **Examples**

```
data = data.table(ID=1:10, X=sample(1:10, 10))
data
data[ID %notin% c(1, 3, 5, 7, 9)]
```

%partin%

A simple extension of %in%.

# Description

A simple extension of %in%.

## Usage

```
pattern %partin% vector
```

## Arguments

pattern Character string containing **regular expressions** to be matched.

vector Character vector.

84

#### Value

TRUE or FALSE.

#### See Also

```
%in%, %allin%, %anyin%, %nonein%
```

## **Examples**

```
"Bei" %partin% c("Beijing", "Shanghai") # TRUE
"bei" %partin% c("Beijing", "Shanghai") # FALSE
"[aeiou]ng" %partin% c("Beijing", "Shanghai") # TRUE
```

%^%

Paste strings together.

## Description

Paste strings together. A wrapper of paste0(). Why %^%? Because typing % and ^ is pretty easy by pressing **Shift** + 5 + 6 + 5.

#### Usage

```
x %^% y
```

# Arguments

x,y

Any objects, usually a numeric or character string or vector.

#### Value

A character string/vector of the pasted values.

```
"He" %^% "llo"
"X" %^% 1:10
"Q" %^% 1:5 %^% letters[1:5]
```

# **Index**

.mean, 4	CONSEC (%%COMPUTE%%), 79
.mean (%%COMPUTE%%), 79	cor_diff, <i>4</i> , 15
$.  sum,   extit{4}$	Corr, 4, 13, 16, 58
.sum (%%COMPUTE%%), 79	COUNT, 4
:=, <i>5</i>	COUNT (%%COMPUTE%%), 79
%%COMPUTE%%, 79	
% <b>^</b> %, <i>4</i> , 84	data.table, 5, 6
%allin%, 4, 77, 78, 82, 84	data.table::fread(),39
%anyin%, 4, 77, 78, 82, 84	data.table::fwrite(),25
%in%, 77, 78, 82–84	Describe, 4, 14, 15, 57
%nonein%, 4, 77, 78, 82, 84	dplyr::left_join(),44
%notin%, 4, 83	dtime, 17
%partin%, 4, 77, 78, 82, 83	
	EFA, 4, 9, 13, 18, 58
add, 4, 5	effectsize::sd_pooled(), 22
add()/added(), <i>81</i>	effectsize::t_to_d(), 21
added, 4	EMMEANS, 4, 21, 47, 76
added (add), 5	emmeans::contrast(), 21
afex::aov_ez(), 22, 45, 47	emmeans::eff_size(), 21
Alpha, 4, 8, 13, 20, 79	emmeans::emmeans(), 21, 22
	emmeans::emmip(), 22, 47
base::load(), <i>40</i>	<pre>emmeans::joint_tests(), 21, 22</pre>
base::readRDS(), 40	emmeans::summary(), 22
base::save(), 26	export, 4, 25, 40
base::saveRDS(), 26	
BayesFactor::ttestBF(), 75	foreign::read.dta(),40
bruceR (bruceR-package), 3	foreign::read.spss(), 39
bruceR-package, 3	format, 27, 28
bruceR::grand_mean_center(), 59	formatF, 4, 27, 28
<pre>bruceR::group_mean_center(), 59</pre>	formatN, 4, 27, 28
bruceR::model_summary(), 59	formula_expand, 28
bruceR::PROCESS(), 58, 59	formula_paste, 29
<b>V</b> / /	Freq, 4, 30, 58
car::recode(), 64	, ,
cc, 4, 9, 46	GGally::ggpairs(), 16
ccf_plot, 5, 10, 34, 35	<pre>ggtext::element_markdown(), 72</pre>
CFA, 4, 9, 12, 20, 42, 58	<pre>ggtext::element_textbox(), 72</pre>
clipr::read_clip_tbl(), 40	<pre>ggtext::geom_richtext(), 72</pre>
clipr::write_clip(), 26	<pre>ggtext::geom_textbox(), 72</pre>
CONSEC, 4	GLM_summary, 5, 31, 39, 51, 66
<i>'</i>	,.,,,

86 INDEX

01 4	
Glue, 4	pkg_depend, 4, 54, 55
Glue (Print), 55	pkg_install_suggested, 4, 54, 55
glue::glue(), 56	Print, 4, 55
glue::glue_col(), 56	print_table, 4, 26, 31, 39, 51, 56, 66
$grand_mean_center, 5, 32, 36$	PROCESS, 5, 42, 49, 51, 58, 58
granger_causality, 5, 33, 35, 58	psych::alpha(), 8, 9
granger_test, 5, 11, 34, 34, 58	psych::corr.test(), 14
group_mean_center, <i>5</i> , <i>32</i> , 35	psych::fa(), 18, 20
	psych::kaiser(), 20
haven::read_dta(),40	psych::omega(), 8, 9
haven::read_sav(), 39	psych::principal(), 18, 20
haven::write_dta(), 26	
haven::write_sav(), 25	readxl::read_excel(), 39
HLM_ICC_rWG, 5, 36	RECODE, 4, 64
HLM_summary, 5, 31, 38, 51, 66	regress, 5, 31, 39, 65
_ , , , , ,	rep_char, 67
import, 4, 26, 39	RESCALE, 4, 67
<pre>interactions::sim_slopes(), 59</pre>	RESCALE(), 69
	RGB, 68
lavaan, <i>41</i> , <i>62</i>	rio::export(), 25, 26
lavaan options, 12	rio::import(), 39, 40
lavaan::cfa(), <i>12</i>	
lavaan::sem(), 59	Run, 4, 69
lavaan_summary, 5, 13, 41, 51, 58, 62	scaler, 69
lme4, <i>61</i>	
<pre>lmtest::grangertest(), 34</pre>	set.wd, 4, 70
LOOKUP, 4, 43	set_wd, 4
200101, 7, 13	set_wd (set.wd), 70
MANOVA, 4, 21–23, 45, 58, 76	setwd, 70
MEAN, 4, 9, 20	show_colors, 5, 71
MEAN (%%COMPUTE%%), 79	stats::p.adjust(), <i>14</i> , <i>22</i>
med_summary, 5, 49, 51, 58, 62	STD, 4
mediation, 49, 61, 62	STD (%%COMPUTE%%), 79
mediation::mediate(), 49, 59	SUM, 4
MODE, 4	SUM (%%COMPUTE%%), 79
MODE (%COMPUTE%), 79	texreg::htmlreg(), <i>50</i> , <i>51</i>
model_summary, 5, 31, 38, 39, 50, 58, 62, 65,	texreg::screenreg(), <i>50</i> , <i>51</i>
66	theme_bruce, 5, 72
MuMIn::r.squaredGLMM(), 50	tidyr::pivot_wider(),47
MuMIn::std.coef(), 50	transmute, 5
mutate, 5	TTEST, 4, 23, 47, 58, 74
openxlsx::write.xlsx(), 25	utils::write.table(), 25
p, 53	VAR, <i>33</i>
PCA, 4, 58	vars::VAR(), 33
PCA (EFA), 18	ναι 3 ΥΛΙΝ( <i>)</i> , 33
<pre>performance::check_collinearity(), 51</pre>	within, 5
performance::r2_mcfadden(), 50	
performance::r2_nagelkerke(), 50	