# Package 'sjPlot'

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Title Data Visualization for Statistics in Social Science

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Description Collection of plotting and table output functions for data visualization. Results of various statistical analyses (that are commonly used in social sciences) can be visualized using this package, including simple and cross tabulated frequencies, histograms, box plots, (generalized) linear models, mixed effects models, principal component analysis and correlation matrices, cluster analyses, scatter plots, stacked scales, effects plots of regression models (including interaction terms) and much more. This package supports labelled data.

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**Depends** R (>= 3.2), graphics, grDevices, stats, utils

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**Suggests** brms, car, cluster, cowplot, GPArotation, gridExtra, ggrepel, ggridges, pscl, rstanarm, survey, TMB, Zelig, testthat

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# **Description**

Collection of plotting and table output functions for data visualization. Results of various statistical analyses (that are commonly used in social sciences) can be visualized using this package, including simple and cross tabulated frequencies, histograms, box plots, (generalized) linear models, mixed effects models, PCA and correlation matrices, cluster analyses, scatter plots, Likert scales, effects plots of interaction terms in regression models, constructing index or score variables and much more.

The package supports labelled data, i.e. value and variable labels from labelled data (like vectors or data frames) are automatically used to label the output. Own labels can be specified as well.

What does this package do?

In short, the functions in this package mostly do two things:

- 1. compute basic or advanced statistical analyses
- 2. either plot the results as ggplot-figure or print them as html-table

How does this package help me?

One of the more challenging tasks when working with R is to get nicely formatted output of statistical analyses, either in graphical or table format. The sjPlot-package takes over these tasks and makes it easy to create beautiful figures or tables.

There are many examples for each function in the related help files and a comprehensive online documentation at <a href="http://www.strengejacke.de/sjPlot">http://www.strengejacke.de/sjPlot</a>.

A note on the package functions

The main functions follow specific naming conventions, hence starting with a specific prefix, which indicates what kind of task these functions perform.

- sjc cluster analysis functions
- sjp plotting functions
- sjt (HTML) table output functions

## Author(s)

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dist\_chisq

	dist_chisq	Plot chi-squared distributions	
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## **Description**

This function plots a simple chi-squared distribution or a chi-squared distribution with shaded areas that indicate at which chi-squared value a significant p-level is reached.

## Usage

```
dist_chisq(chi2 = NULL, deg.f = NULL, p = NULL, xmax = NULL,
  geom.colors = NULL, geom.alpha = 0.7)
```

# **Arguments**

chi2	Numeric, optional. If specified, a chi-squared distribution with deg.f degrees of freedom is plotted and a shaded area at chi2 value position is plotted that indicates whether or not the specified value is significant or not. If both chi2 and p are not specified, a distribution without shaded area is plotted.
deg.f	Numeric. The degrees of freedom for the chi-squared distribution. Needs to be specified.
p	Numeric, optional. If specified, a chi-squared distribution with deg.f degrees of freedom is plotted and a shaded area at the position where the specified plevel starts is plotted. If both chi2 and p are not specified, a distribution without shaded area is plotted.
xmax	Numeric, optional. Specifies the maximum x-axis-value. If not specified, the x-axis ranges to a value where a p-level of 0.00001 is reached.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
geom.alpha	Specifies the alpha-level of the shaded area. Default is 0.7, range between 0 to 1.

# **Examples**

```
# a simple chi-squared distribution
# for 6 degrees of freedom
dist_chisq(deg.f = 6)

# a chi-squared distribution for 6 degrees of freedom,
# and a shaded area starting at chi-squared value of ten.
# With a df of 6, a chi-squared value of 12.59 would be "significant",
# thus the shaded area from 10 to 12.58 is filled as "non-significant",
# while the area starting from chi-squared value 12.59 is filled as
# "significant"
dist_chisq(chi2 = 10, deg.f = 6)

# a chi-squared distribution for 6 degrees of freedom,
# and a shaded area starting at that chi-squared value, which has
```

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```
# a p-level of about 0.125 (which equals a chi-squared value of about 10). # With a df of 6, a chi-squared value of 12.59 would be "significant", # thus the shaded area from 10 to 12.58 (p-level 0.125 to p-level 0.05) # is filled as "non-significant", while the area starting from chi-squared # value 12.59 (p-level < 0.05) is filled as "significant". dist_chisq(p = 0.125, deg.f = 6)
```

 $dist_f$ 

Plot F distributions

# Description

This function plots a simple F distribution or an F distribution with shaded areas that indicate at which F value a significant p-level is reached.

# Usage

```
dist_f(f = NULL, deg.f1 = NULL, deg.f2 = NULL, p = NULL,
    xmax = NULL, geom.colors = NULL, geom.alpha = 0.7)
```

f	Numeric, optional. If specified, an F distribution with deg.f1 and deg.f2 degrees of freedom is plotted and a shaded area at f value position is plotted that indicates whether or not the specified value is significant or not. If both f and p are not specified, a distribution without shaded area is plotted.
deg.f1	Numeric. The first degrees of freedom for the F distribution. Needs to be specified.
deg.f2	Numeric. The second degrees of freedom for the F distribution. Needs to be specified.
p	Numeric, optional. If specified, a F distribution with deg.f1 and deg.f2 degrees of freedom is plotted and a shaded area at the position where the specified p-level starts is plotted. If both f and p are not specified, a distribution without shaded area is plotted.
xmax	Numeric, optional. Specifies the maximum x-axis-value. If not specified, the x-axis ranges to a value where a p-level of 0.00001 is reached.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
geom.alpha	Specifies the alpha-level of the shaded area. Default is $0.7$ , range between $0$ to $1$ .

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## **Examples**

```
# a simple F distribution for 6 and 45 degrees of freedom
dist_f(deg.f1 = 6, deg.f2 = 45)

# F distribution for 6 and 45 degrees of freedom,
# and a shaded area starting at F value of two.
# F-values equal or greater than 2.31 are "significant"
dist_f(f = 2, deg.f1 = 6, deg.f2 = 45)

# F distribution for 6 and 45 degrees of freedom,
# and a shaded area starting at a p-level of 0.2
# (F-Value about 1.5).
dist_f(p = 0.2, deg.f1 = 6, deg.f2 = 45)
```

 $dist\_norm$ 

Plot normal distributions

# **Description**

This function plots a simple normal distribution or a normal distribution with shaded areas that indicate at which value a significant p-level is reached.

# Usage

```
dist_norm(norm = NULL, mean = 0, sd = 1, p = NULL, xmax = NULL,
  geom.colors = NULL, geom.alpha = 0.7)
```

norm	Numeric, optional. If specified, a normal distribution with mean and sd is plotted and a shaded area at norm value position is plotted that indicates whether or not the specified value is significant or not. If both norm and p are not specified, a distribution without shaded area is plotted.
mean	Numeric. Mean value for normal distribution. By default 0.
sd	Numeric. Standard deviation for normal distribution. By default 1.
р	Numeric, optional. If specified, a normal distribution with mean and sd is plotted and a shaded area at the position where the specified p-level starts is plotted. If both norm and p are not specified, a distribution without shaded area is plotted.
xmax	Numeric, optional. Specifies the maximum x-axis-value. If not specified, the x-axis ranges to a value where a p-level of 0.00001 is reached.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
geom.alpha	Specifies the alpha-level of the shaded area. Default is 0.7, range between 0 to 1.

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# **Examples**

```
# a simple normal distribution
dist_norm()

# a simple normal distribution with different mean and sd.
# note that curve looks similar to above plot, but axis range
# has changed.
dist_norm(mean = 2, sd = 4)

# a simple normal distribution
dist_norm(norm = 1)

# a simple normal distribution
dist_norm(p = 0.2)
```

 $dist_t$ 

Plot t-distributions

# Description

This function plots a simple t-distribution or a t-distribution with shaded areas that indicate at which t-value a significant p-level is reached.

# Usage

```
dist_t(t = NULL, deg.f = NULL, p = NULL, xmax = NULL,
  geom.colors = NULL, geom.alpha = 0.7)
```

t	Numeric, optional. If specified, a t-distribution with deg.f degrees of freedom is plotted and a shaded area at t value position is plotted that indicates whether or not the specified value is significant or not. If both t and p are not specified, a distribution without shaded area is plotted.
deg.f	Numeric. The degrees of freedom for the t-distribution. Needs to be specified.
p	Numeric, optional. If specified, a t-distribution with deg.f degrees of freedom is plotted and a shaded area at the position where the specified p-level starts is plotted. If both t and p are not specified, a distribution without shaded area is plotted.
xmax	Numeric, optional. Specifies the maximum x-axis-value. If not specified, the x-axis ranges to a value where a p-level of 0.00001 is reached.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
geom.alpha	Specifies the alpha-level of the shaded area. Default is 0.7, range between 0 to 1.

## **Examples**

```
# a simple t-distribution
# for 6 degrees of freedom
dist_t(deg.f = 6)

# a t-distribution for 6 degrees of freedom,
# and a shaded area starting at t-value of one.
# With a df of 6, a t-value of 1.94 would be "significant".
dist_t(t = 1, deg.f = 6)

# a t-distribution for 6 degrees of freedom,
# and a shaded area starting at p-level of 0.4
# (t-value of about 0.26).
dist_t(p = 0.4, deg.f = 6)
```

efc

Sample dataset from the EUROFAMCARE project

#### **Description**

A SPSS sample data set, imported with the read\_spss function.

plot\_frq

Plot frequencies of variables

# **Description**

Plot frequencies of a variable as bar graph, histogram, box plot etc.

## Usage

```
plot_frq(data, ..., title = "", weight.by = NULL,
    title.wtd.suffix = NULL, sort.frq = c("none", "asc", "desc"),
    type = c("bar", "dot", "histogram", "line", "density", "boxplot",
    "violin"), geom.size = NULL, geom.colors = "#336699",
    errorbar.color = "darkred", axis.title = NULL, axis.labels = NULL,
    xlim = NULL, ylim = NULL, wrap.title = 50, wrap.labels = 20,
    grid.breaks = NULL, expand.grid = FALSE, show.values = TRUE,
    show.n = TRUE, show.prc = TRUE, show.axis.values = TRUE,
    show.ci = FALSE, show.na = FALSE, show.mean = FALSE,
    show.mean.val = TRUE, show.sd = TRUE, drop.empty = TRUE,
    mean.line.type = 2, mean.line.size = 0.5, inner.box.width = 0.15,
    inner.box.dotsize = 3, normal.curve = FALSE,
    normal.curve.color = "red", normal.curve.size = 0.8,
    normal.curve.alpha = 0.4, auto.group = NULL, coord.flip = FALSE,
    vjust = "bottom", hjust = "center", y.offset = NULL)
```

## **Arguments**

data A data frame, or a grouped data frame. Optional, unquoted names of variables that should be selected for further processing. Required, if data is a data frame (and no vector) and only selected variables from data should be processed. You may also use functions like: or tidyselect's select\_helpers. title Character vector, used as plot title. By default, get\_dv\_labels is called to retrieve the label of the dependent variable, which will be used as title. Use title = "" to remove title. Vector of weights that will be applied to weight all cases. Must be a vector of weight.by same length as the input vector. Default is NULL, so no weights are used. title.wtd.suffix Suffix (as string) for the title, if weight.by is specified, e.g. title.wtd.suffix=" (weighted)". Default is NULL, so title will not have a suffix when cases are weighted. sort.frq Determines whether categories should be sorted according to their frequencies or not. Default is "none", so categories are not sorted by frequency. Use "asc" or "desc" for sorting categories ascending or descending order. Specifies the plot type. May be abbreviated. type "bar" for simple bars (default) "dot" for a dot plot "histogram" for a histogram (does not apply to grouped frequencies) "line" for a line-styled histogram with filled area "density" for a density plot (does not apply to grouped frequencies) "boxplot" for box plot "violin" for violin plots geom.size size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes. geom.colors User defined color for geoms, e.g. geom. colors = "#0080ff". errorbar.color Color of confidence interval bars (error bars). Only applies to type = "bar". In case of dot plots, error bars will have same colors as dots (see geom. colors). axis.title Character vector of length one or two (depending on the plot function and type), used as title(s) for the x and y axis. If not specified, a default labelling is chosen. **Note:** Some plot types do not support this argument. In such cases, use the return value and add axis titles manually with labs, e.g.: plot.list[[1]] + labs(x = ...)axis.labels character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically. xlim Numeric vector of length two, defining lower and upper axis limits of the x scale. By default, this argument is set to NULL, i.e. the x-axis fits to the required range of the data. numeric vector of length two, defining lower and upper axis limits of the y scale. ylim By default, this argument is set to NULL, i.e. the y-axis fits to the required range of the data.

wrap.title Numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted. wrap.labels numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted. grid.breaks numeric; sets the distance between breaks for the axis, i.e. at every grid. breaks'th position a major grid is being printed. expand.grid logical, if TRUE, the plot grid is expanded, i.e. there is a small margin between axes and plotting region. Default is FALSE. show.values Logical, whether values should be plotted or not. logical, if TRUE, adds total number of cases for each group or category to the show.n labels. logical, if TRUE (default), percentage values are plotted to each bar If FALSE, show.prc percentage values are removed. show.axis.values logical, whether category, count or percentage values for the axis should be printed or not. show.ci Logical, if TRUE), adds notches to the box plot, which are used to compare groups; if the notches of two boxes do not overlap, medians are considered to be significantly different. logical, if TRUE, NA's (missing values) are added to the output. show.na show.mean Logical, if TRUE, a vertical line in histograms is drawn to indicate the mean value of the variables. Only applies to histogram-charts. Logical, if TRUE (default), the mean value is printed to the vertical line that show.mean.val indicates the variable's mean. Only applies to histogram-charts. Logical, if TRUE, the standard deviation is annotated as shaded rectangle around show.sd the mean intercept line. Only applies to histogram-charts. Logical, if TRUE and the variable's values are labelled, values that have no obserdrop.empty vations are still printed in the table (with frequency 0). If FALSE, values / factor levels with no occurence in the data are omitted from the output. mean.line.type Numeric value, indicating the linetype of the mean intercept line. Only applies to histogram-charts and when show.mean = TRUE. mean.line.size Numeric, size of the mean intercept line. Only applies to histogram-charts and when show.mean = TRUE. inner.box.width width of the inner box plot that is plotted inside of violin plots. Only applies if type = "violin". Default value is 0.15 inner.box.dotsize size of mean dot insie a violin or box plot. Applies only when type = "violin" or "boxplot". Logical, if TRUE, a normal curve, which is adjusted to the data, is plotted over normal.curve the histogram or density plot. Default is FALSE. Only applies when histograms or density plots are plotted (see type). normal.curve.color

Color of the normal curve line. Only applies if normal.curve = TRUE.

normal.curve.size

Numeric, size of the normal curve line. Only applies if normal.curve = TRUE.

normal.curve.alpha

Transparancy level (alpha value) of the normal curve. Only applies if normal.curve = TRUE.

auto.group

numeric value, indicating the minimum amount of unique values in the count variable, at which automatic grouping into smaller units is done (see group\_var). Default value for auto.group is NULL, i.e. auto-grouping is off. See group\_var

for examples on grouping.

coord.flip

logical, if TRUE, the x and y axis are swapped.

vjust

character vector, indicating the vertical position of value labels. Allowed are same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which

align text towards and away from the center of the plot respectively.

hjust

character vector, indicating the horizontal position of value labels. Allowed are same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which

align text towards and away from the center of the plot respectively.

y.offset

numeric, offset for text labels when their alignment is adjusted to the top/bottom

of the geom (see hjust and vjust).

## Value

A ggplot-object.

#### Note

This function only works with variables with integer values (or numeric factor levels), i.e. scales / centered variables with fractional part may result in unexpected behaviour.

# **Examples**

```
library(sjlabelled)
library(dplyr)
data(efc)

# boxplot
plot_frq(efc$e17age, type = "box")

# histogram, pipe-workflow
efc %>%
    dplyr::select(e17age, c160age) %>%
    plot_frq(type = "hist", show.mean = TRUE)

# bar plot(s)
plot_frq(efc, e42dep, c172code)

# grouped data frame, all panels in one plot
efc %>%
    group_by(e42dep) %>%
```

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```
plot_frq(c161sex) %>%
  plot_grid()

library(sjmisc)
# grouped variable
ageGrp <- group_var(efc$e17age)
ageGrpLab <- group_labels(efc$e17age)
plot_frq(ageGrp, title = get_label(efc$e17age), axis.labels = ageGrpLab)

# plotting confidence intervals. expand grid and v/hjust for text labels
plot_frq(
  efc$e15relat, type = "dot", show.ci = TRUE, sort.frq = "desc",
      coord.flip = TRUE, expand.grid = TRUE, vjust = "bottom", hjust = "left"
)

# histogram with overlayed normal curve
plot_frq(efc$c160age, type = "h", show.mean = TRUE, show.mean.val = TRUE,
      normal.curve = TRUE, show.sd = TRUE, normal.curve.color = "blue",
      normal.curve.size = 3, ylim = c(0,50))</pre>
```

plot\_gpt

Plot grouped proportional tables

## Description

Plot grouped proportional crosstables, where the proportion of each level of x for the highest category in y is plotted, for each subgroup of grp.

#### Usage

```
plot_gpt(data, x, y, grp, colors = "metro ui", geom.size = 2.5,
    shape.fill.color = "#f0f0f0", shapes = c(15, 16, 17, 18, 21, 22, 23,
    24, 25, 7, 8, 9, 10, 12), title = NULL, axis.labels = NULL,
    axis.titles = NULL, legend.title = NULL, legend.labels = NULL,
    wrap.title = 50, wrap.labels = 15, wrap.legend.title = 20,
    wrap.legend.labels = 20, axis.lim = NULL, grid.breaks = NULL,
    show.total = TRUE, annotate.total = TRUE, show.p = TRUE,
    show.n = TRUE)
```

data	A data frame, or a grouped data frame.
Х	Categorical variable, where the proportion of each category in x for the highest category of y will be printed along the x-axis.
у	Categorical or numeric variable. If not a binary variable, y will be recoded into a binary variable, dichtomized at the highest category and all remaining categories.

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grp

Grouping variable, which will define the y-axis

colors

May be a character vector of color values in hex-format, valid color value names (see demo("colors")) or a name of a pre-defined color palette. Following options are valid for the colors argument:

- If not specified, a default color brewer palette will be used, which is suitable for the plot style.
- If "gs", a greyscale will be used.
- If "bw", and plot-type is a line-plot, the plot is black/white and uses different line types to distinguish groups (see this package-vignette).
- If colors is any valid color brewer palette name, the related palette will be used. Use display.brewer.all to view all available palette names.
- There are some pre-defined color palettes in this package, see sjPlot-themes for details.
- Else specify own color values or names as vector (e.g. colors = "#00ff00" or colors = c("firebrick", "blue")).

geom.size

size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.

shape.fill.color

Optional color vector, fill-color for non-filled shapes

shapes Numeric vector with shape styles, used to map the different categories of x.

Character vector, used as plot title. By default, get\_dv\_labels is called to title retrieve the label of the dependent variable, which will be used as title. Use

title = "" to remove title.

axis.labels character vector with labels used as axis labels. Optional argument, since in

most cases, axis labels are set automatically.

character vector of length one or two, defining the title(s) for the x-axis and axis.titles

y-axis.

legend.title Character vector, used as legend title for plots that have a legend.

legend.labels character vector with labels for the guide/legend.

wrap.title Numeric, determines how many chars of the plot title are displayed in one line

and when a line break is inserted.

wrap.labels numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.

wrap.legend.title

numeric, determines how many chars of the legend's title are displayed in one line and when a line break is inserted.

wrap.legend.labels

numeric, determines how many chars of the legend labels are displayed in one line and when a line break is inserted.

axis.lim Numeric vector of length 2, defining the range of the plot axis. Depending on plot type, may effect either x- or y-axis, or both. For multiple plot outputs (e.g., from type = "eff" or type = "slope" in plot\_model), axis.lim may also be a list of vectors of length 2, defining axis limits for each plot (only if nonfaceted).

plot\_grid

numeric; sets the distance between breaks for the axis, i.e. at every grid.breaks'th position a major grid is being printed.

show.total Logical, if TRUE, a total summary line for all aggregated grp is added.

Logical, if TRUE and show.total = TRUE, the total-row in the figure will be highlighted with a slightly shaded background.

show.p Logical, adds significance levels to values, or value and variable labels.

show.n logical, if TRUE, adds total number of cases for each group or category to the

labels.

## **Details**

The p-values are based on chisq.test of x and y for each grp.

#### Value

A ggplot-object.

# **Examples**

```
data(efc)

# the proportion of dependency levels in female
# elderly, for each family carer's relationship
# to elderly
plot_gpt(efc, e42dep, e16sex, e15relat)

# proportion of educational levels in highest
# dependency category of elderly, for different
# care levels
plot_gpt(efc, c172code, e42dep, n4pstu)
```

plot\_grid

Arrange list of plots as grid

# **Description**

Plot multiple ggplot-objects as a grid-arranged single plot.

# Usage

```
plot_grid(x, margin = c(1, 1, 1, 1))
```

# Arguments

x A list of ggplot-objects. See 'Details'.

margin A numeric vector of length 4, indicating the top, right, bottom and left margin

for each plot, in centimetres.

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## **Details**

This function takes a list of ggplot-objects as argument. Plotting functions of this package that produce multiple plot objects (e.g., when there is an argument facet.grid) usually return multiple plots as list (the return value is named plot.list). To arrange these plots as grid as a single plot, use plot\_grid.

#### Value

An object of class gtable.

## **Examples**

```
library(ggeffects)
data(efc)
# fit model
fit <- glm(
  tot_sc_e \sim c12hour + e17age + e42dep + neg_c_7,
  family = poisson
)
# plot marginal effects for each predictor, each as single plot
p1 <- ggpredict(fit, "c12hour") %>%
  plot(show.y.title = FALSE, show.title = FALSE)
p2 <- ggpredict(fit, "e17age") %>%
  plot(show.y.title = FALSE, show.title = FALSE)
p3 <- ggpredict(fit, "e42dep") %>%
  plot(show.y.title = FALSE, show.title = FALSE)
p4 <- ggpredict(fit, "neg_c_7") %>%
  plot(show.y.title = FALSE, show.title = FALSE)
# plot grid
plot_grid(list(p1, p2, p3, p4))
```

plot\_kfold\_cv

Plot model fit from k-fold cross-validation

# Description

This function plots the aggregated residuals of k-fold cross-validated models against the outcome. This allows to evaluate how the model performs according over- or underestimation of the outcome.

## Usage

```
plot_kfold_cv(data, formula, k = 5, fit)
```

plot\_kfold\_cv

#### **Arguments**

data A data frame, used to split the data into k trainig-test-pairs.

formula A model formula, used to fit linear models (1m) over all k training data sets.

Use fit to specify a fitted model (also other models than linear models), which will be used to compute cross validation. If fit is not missing, formula will be

ignored.

k Number of folds.

fit Model object, which will be used to compute cross validation. If fit is not

missing, formula will be ignored. Currently, only linear, poisson and negative

binomial regression models are supported.

#### **Details**

This function, first, generates k cross-validated test-training pairs (using the crossv\_kfold-function) and fits the same model, specified in the formula- or fit- argument, over all training data sets.

Then, the test data is used to predict the outcome from all models that have been fit on the training data, and the residuals from all test data is plotted against the observed values (outcome) from the test data (note: for poisson or negative binomial models, the deviance residuals are calculated). This plot can be used to validate the model and see, whether it over- (residuals > 0) or underestimates (residuals < 0) the model's outcome.

#### Note

Currently, only linear, poisson and negative binomial regression models are supported.

## **Examples**

```
data(efc)
plot_kfold_cv(efc, neg_c_7 ~ e42dep + c172code + c12hour)
plot_kfold_cv(mtcars, mpg ~.)

# for poisson models. need to fit a model and use 'fit'-argument
fit <- glm(tot_sc_e ~ neg_c_7 + c172code, data = efc, family = poisson)
plot_kfold_cv(efc, fit = fit)

# and for negative binomial models
fit <- MASS::glm.nb(tot_sc_e ~ neg_c_7 + c172code, data = efc)
plot_kfold_cv(efc, fit = fit)</pre>
```

prot_riner c row interest as centered statement burs	plot_likert	Plot likert scales as centered stacked bars
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## **Description**

Plot likert scales as centered stacked bars.

## Usage

```
plot_likert(items, groups = NULL, groups.titles = "auto",
   title = NULL, legend.title = NULL, legend.labels = NULL,
   axis.titles = NULL, axis.labels = NULL, catcount = NULL,
   cat.neutral = NULL, sort.frq = NULL, weight.by = NULL,
   title.wtd.suffix = NULL, wrap.title = 50, wrap.labels = 30,
   wrap.legend.title = 30, wrap.legend.labels = 28, geom.size = 0.6,
   geom.colors = "BrBG", cat.neutral.color = "grey70",
   intercept.line.color = "grey50", reverse.colors = FALSE,
   values = "show", show.n = TRUE, show.legend = TRUE,
   show.prc.sign = FALSE, grid.range = 1, grid.breaks = 0.2,
   expand.grid = TRUE, digits = 1, reverse.scale = FALSE,
   coord.flip = TRUE, sort.groups = TRUE, legend.pos = "bottom",
   rel_heights = 1, cowplot.options = list(label_x = 0.01, hjust = 0,
   align = "v"))
```

items	Data frame, with each column representing one item.
groups	(optional) Must be a vector of same length as ncol(items), where each item in this vector represents the group number of the related columns of items. See 'Examples'.
groups.titles	(optional, only used if groups are supplied) Titles for each factor group that will be used as table caption for each component-table. Must be a character vector of same length as length(unique(groups)). Default is "auto", which means that each table has a standard caption <i>Component x</i> . Use NULL to use names as supplied to groups and use FALSE to suppress table captions.
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If $title = ""$ , no title is printed. For effect-plots, may also be a character vector of length $> 1$ , to define titles for each sub-plot or facet.
legend.title	character vector, used as title for the plot legend.
legend.labels	character vector with labels for the guide/legend.
axis.titles	character vector of length one or two, defining the title(s) for the x-axis and y-axis.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.

catcount optional, amount of categories of items (e.g. "strongly disagree", "disagree", "disagree", "agree" and "strongly agree" would be categories of items (e.g. "strongly disagree", "like the this argu-

"agree" and "strongly agree" would be catcount = 4). Note that this argument only applies to "valid" answers, i.e. if you have an additional neutral category (see cat.neutral) like "don't know", this won't count for catcount (e.g. "strongly disagree", "disagree", "agree", "strongly agree" and neutral category

"don't know" would still mean that catcount = 4). See 'Note'.

cat.neutral If there's a neutral category (like "don't know" etc.), specify the index number

(value) for this category. Else, set cat.neutral = NULL (default). The proportions of neutral category answers are plotted as grey bars on the left side of the

figure.

sort.frq Indicates whether the items of items should be ordered by total sum of positive

or negative answers.

"pos.asc" to order ascending by sum of positive answers

"pos.desc" to order descending by sum of positive answers

"neg.asc" for sorting ascending negative answers

"neg.desc" for sorting descending negative answers

NULL (default) for no sorting

weight.by Vector of weights that will be applied to weight all cases. Must be a vector of

same length as the input vector. Default is NULL, so no weights are used.

title.wtd.suffix

Suffix (as string) for the title, if weight.by is specified, e.g. title.wtd.suffix=" (weighted)".

Default is NULL, so title will not have a suffix when cases are weighted.

wrap.title numeric, determines how many chars of the plot title are displayed in one line

and when a line break is inserted.

wrap.labels numeric, determines how many chars of the value, variable or axis labels are

displayed in one line and when a line break is inserted.

wrap.legend.title

numeric, determines how many chars of the legend's title are displayed in one

line and when a line break is inserted.

wrap.legend.labels

numeric, determines how many chars of the legend labels are displayed in one

line and when a line break is inserted.

geom. size size resp. width of the geoms (bar width, line thickness or point size, depending

on plot type and function). Note that bar and bin widths mostly need smaller

values than dot sizes.

geom. colors user defined color for geoms. See 'Details' in sjp.grpfrq.

cat.neutral.color

Color of the neutral category, if plotted (see cat.neutral).

intercept.line.color

Color of the vertical intercept line that divides positive and negative values.

reverse.colors logical, if TRUE, the color scale from geom.colors will be reversed, so positive

and negative values switch colors.

values Determines style and position of percentage value labels on the bars:

> "show" (default) shows percentage value labels in the middle of each category bar

"hide" hides the value labels, so no percentage values on the bars are printed

"sum.inside" shows the sums of percentage values for both negative and positive values and prints them inside the end of each bar

"sum.outside" shows the sums of percentage values for both negative and positive values and prints them outside the end of each bar

show.n logical, if TRUE, adds total number of cases for each group or category to the labels.

> logical, if TRUE, and depending on plot type and function, a legend is added to the plot.

logical, if TRUE, %-signs for value labels are shown. show.prc.sign

> Numeric, limits of the x-axis-range, as proportion of 100. Default is 1, so the xscale ranges from zero to 100% on both sides from the center. Can alternatively be supplied as a vector of 2 positive numbers (e.g. grid.range = c(1, .8)) to set the left and right limit separately. You can use values beyond 1 (100%) in case bar labels are not printed because they exceed the axis range. E.g. grid.range = 1.4 will set the axis from -140 to +140%, however, only (valid) axis labels from -100 to +100% are printed. Neutral categories are adjusted to the most left limit.

numeric; sets the distance between breaks for the axis, i.e. at every grid. breaks'th grid.breaks position a major grid is being printed.

logical, if TRUE, the plot grid is expanded, i.e. there is a small margin between axes and plotting region. Default is FALSE.

Numeric, amount of digits after decimal point when rounding estimates or values.

logical, if TRUE, the ordering of the categories is reversed, so positive and negative values switch position.

coord.flip logical, if TRUE, the x and y axis are swapped.

sort.groups (optional, only used if groups are supplied) logical, if groups should be sorted according to the values supplied to groups. Defaults to TRUE.

> (optional, only used if groups are supplied) Defines the legend position. Possible values are c("bottom", "top", "both", "all", "none"). If the is only one group or this option is set to "all" legends will be printed as defined with set\_theme.

> (optional, only used if groups are supplied) This option can be used to adjust the height of the subplots. The bars in subplots can have different heights due to a differing number of items or due to legend placement. This can be adjusted here. Takes a vector of numbers, one for each plot. Values are evaluated relative to each other.

cowplot.options

(optional, only used if groups are supplied) List of label options to be passed to plot\_grid.

show.legend

grid.range

expand.grid

digits

reverse.scale

legend.pos

rel\_heights

#### Value

A ggplot-object.

## Note

Note that only even numbers of categories are possible to plot, so the "positive" and "negative" values can be splitted into two halfs. A neutral category (like "don't know") can be used, but must be indicated by cat.neutral.

The catcount-argument indicates how many item categories are in the Likert scale. Normally, this argument can be ignored because the amount of valid categories is retrieved automatically. However, sometimes (for instance, if a certain category is missing in all items), auto-detection of the amount of categories fails. In such cases, specify the amount of categories with the catcountargument.

## **Examples**

```
library(sjmisc)
data(efc)
# find all variables from COPE-Index, which all have a "cop" in their
# variable name, and then plot that subset as likert-plot
mydf <- find_var(efc, pattern = "cop", out = "df")</pre>
plot_likert(mydf)
plot_likert(
  mydf,
  grid.range = c(1.2, 1.4),
  expand.grid = FALSE,
  values = "sum.outside",
  show.prc.sign = TRUE
)
# Plot in groups
plot_likert(mydf, c(2,1,1,1,1,2,2,2,1))
groups <- sjt.pca(mydf)$factor.index</pre>
plot_likert(mydf, groups = groups)
plot_likert(mydf,
            c(rep("B", 4), rep("A", 5)),
            sort.groups = FALSE,
            grid.range = c(0.9, 1.1),
            geom.colors = "RdBu",
            rel_heights = c(6, 8),
            wrap.labels = 40,
            reverse.scale = TRUE)
```

plot\_model

Plot regression models

#### **Description**

plot\_model() creates plots from regression models, either estimates (as so-called forest or dot whisker plots) or marginal effects.

#### Usage

```
plot_model(model, type = c("est", "re", "eff", "pred", "int", "std",
  "std2", "slope", "resid", "diag"), transform, terms = NULL,
  sort.est = NULL, rm.terms = NULL, group.terms = NULL,
 order.terms = NULL, pred.type = c("fe", "re"),
 mdrt.values = c("minmax", "meansd", "zeromax", "quart", "all"),
  ri.nr = NULL, title = NULL, axis.title = NULL,
  axis.labels = NULL, legend.title = NULL, wrap.title = 50,
 wrap.labels = 25, axis.lim = NULL, grid.breaks = NULL,
  ci.lvl = NULL, se = NULL, vcov.fun = NULL, vcov.type = c("HC3",
  "const", "HC", "HC0", "HC1", "HC2", "HC4", "HC4m", "HC5"),
 vcov.args = NULL, colors = "Set1", show.intercept = FALSE,
  show.values = FALSE, show.p = TRUE, show.data = FALSE,
  show.legend = TRUE, show.zeroinf = TRUE, value.offset = NULL,
  value.size, jitter = NULL, digits = 2, dot.size = NULL,
  line.size = NULL, vline.color = NULL, p.threshold = c(0.05, 0.01,
  0.001), grid, case, auto.label = TRUE, prefix.labels = c("none",
  "varname", "label"), bpe = "median", bpe.style = "line",
 bpe.color = "white", ci.style = c("whisker", "bar"), ...)
get_model_data(model, type = c("est", "re", "eff", "pred", "int", "std",
  "std2", "slope", "resid", "diag"), transform, terms = NULL,
  sort.est = NULL, rm.terms = NULL, group.terms = NULL,
 order.terms = NULL, pred.type = c("fe", "re"), ri.nr = NULL,
  ci.lvl = NULL, colors = "Set1", grid, case = "parsed",
 digits = 2, \ldots)
```

## **Arguments**

model

A regression model object. Depending on the type, many kinds of models are supported, e.g. from packages like stats, lme4, nlme, rstanarm, survey, glmmTMB, MASS, brms etc.

type

Type of plot. There are three groups of plot-types:

Coefficients (related vignette)

type = "est" Forest-plot of estimates. If the fitted model only contains one predictor, slope-line is plotted.

type = "re" For mixed effects models, plots the random effects.

type = "std" Forest-plot of standardized beta values.

type = "std2" Forest-plot of standardized beta values, however, standardization is done by dividing by two sd (see 'Details').

Marginal Effects (related vignette)

type = "pred" Predicted values (marginal effects) for specific model terms. See ggpredict for details.

type = "eff" Similar to type = "pred", however, discrete predictors are held constant at their proportions (not reference level). See ggeffect for details.

type = "int" Marginal effects of interaction terms in model.

Model diagnostics

type = "slope" Slope of coefficients for each single predictor, against the response (linear relationship between each model term and response). See 'Details'

type = "resid" Slope of coefficients for each single predictor, against the residuals (linear relationship between each model term and residuals). See 'Details'.

type = "diag" Check model assumptions. See 'Details'.

**Note:** For mixed models, the diagnostic plots like linear relationship or check for Homoscedasticity, do **not** take the uncertainty of random effects into account, but is only based on the fixed effects part of the model.

transform

A character vector, naming a function that will be applied on estimates and confidence intervals. By default, transform will automatically use "exp" as transformation for applicable classes of model (e.g. logistic or poisson regression). Estimates of linear models remain untransformed. Use NULL if you want the raw, non-transformed estimates.

terms

Character vector with the names of those terms from model that should be plotted. This argument depends on the plot-type:

Coefficients Select terms that should be plotted. All other term are removed from the output. Note that the term names must match the names of the model's coefficients. For factors, this means that the variable name is suffixed with the related factor level, and each category counts as one term.
E.g. rm.terms = "t\_name [2,3]" would remove the terms "t\_name2" and "t\_name3" (assuming that the variable t\_name is categorical and has at least the factor levels 2 and 3). Another example for the iris-dataset: terms = "Species" would not work, instead you would write terms = "Species [versicolor, vi to remove these two levels, or terms = "Speciesversicolor" if you just want to remove the level versicolor from the plot.

Marginal Effects Here terms indicates for which terms marginal effects should be displayed. At least one term is required to calculate effects, maximum length is three terms, where the second and third term indicate the groups, i.e. predictions of first term are grouped by the levels of the second (and third) term. terms may also indicate higher order terms (e.g. interaction terms). Indicating levels in square brackets allows for selecting only specific groups. Term name and levels in brackets must be separated by a

whitespace character, e.g. terms = c("age", "education [1,3]"). It is also possible to specify a range of numeric values for the predictions with a colon, for instance terms = c("education [1,3]", "age [30:50]"). Furthermore, it is possible to specify a function name. Values for predictions will then be transformed, e.g. terms = "income [exp]". This is useful when model predictors were transformed for fitting the model and should be back-transformed to the original scale for predictions. Finally, numeric vectors for which no specific values are given, a "pretty range" is calculated, to avoid memory allocation problems for vectors with many unique values. If a numeric vector is specified as second or third term (i.e. if this vector represents a grouping structure), representative values (see rprs\_values) are chosen. If all values for a numeric vector should be used to compute predictions, you may use e.g. terms = "age [all]". For more details, see ggpredict.

sort.est

Determines in which way estimates are sorted in the plot:

- If NULL (default), no sorting is done and estimates are sorted in the same order as they appear in the model formula.
- If TRUE, estimates are sorted in descending order, with highest estimate at the top.
- If sort.est = "sort.all", estimates are re-sorted for each coefficient (only applies if type = "re" and grid = FALSE), i.e. the estimates of the random effects for each predictor are sorted and plotted to an own plot.
- If type = "re", specify a predictor's / coefficient's name to sort estimates according to this random effect.

rm.terms

Character vector with names that indicate which terms should be removed from the plot. Counterpart to terms. rm.terms = "t\_name" would remove the term t\_name. Default is NULL, i.e. all terms are used. For factors, levels that should be removed from the plot need to be explicitly indicated in square brackets, and match the model's coefficient names, e.g. rm.terms = "t\_name [2,3]" would remove the terms "t\_name2" and "t\_name3" (assuming that the variable t\_name was categorical and has at least the factor levels 2 and 3). Another example for the *iris* dataset would be rm.terms = "Species [versicolor, virginica]". Note that the rm.terms-argument does not apply to *Marginal Effects* plots.

group.terms

Numeric vector with group indices, to group coefficients. Each group of coefficients gets its own color (see 'Examples').

order.terms

Numeric vector, indicating in which order the coefficients should be plotted. See examples in this package-vignette.

pred.type

Character, only applies for *Marginal Effects* plots with mixed effects models. Indicates whether predicted values should be conditioned on random effects (pred.type = "re") or fixed effects only (pred.type = "fe", the default). For details, see documentation of the type-argument in ggpredict.

mdrt.values

Indicates which values of the moderator variable should be used when plotting interaction terms (i.e. type = "int").

"minmax" (default) minimum and maximum values (lower and upper bounds) of the moderator are used to plot the interaction between independent variable and moderator(s).

> "meansd" uses the mean value of the moderator as well as one standard deviation below and above mean value to plot the effect of the moderator on the independent variable (following the convention suggested by Cohen and Cohen and popularized by Aiken and West (1991), i.e. using the mean, the value one standard deviation above, and the value one standard deviation below the mean as values of the moderator, see Grace-Martin K: 3 Tips to Make Interpreting Moderation Effects Easier).

> "zeromax" is similar to the "minmax" option, however, 0 is always used as minimum value for the moderator. This may be useful for predictors that don't have an empirical zero-value, but absence of moderation should be simulated by using 0 as minimum.

> "quart" calculates and uses the quartiles (lower, median and upper) of the moderator value.

"all" uses all values of the moderator variable.

Numeric vector. If type = "re" and fitted model has more than one random intercept, ri.nr indicates which random effects of which random intercept (or: which list elements of ranef) will be plotted. Default is NULL, so all random effects will be plotted.

> Character vector, used as plot title. By default, get\_dv\_labels is called to retrieve the label of the dependent variable, which will be used as title. Use title = "" to remove title.

> Character vector of length one or two (depending on the plot function and type), used as title(s) for the x and y axis. If not specified, a default labelling is chosen. Note: Some plot types may not support this argument sufficiently. In such cases, use the returned ggplot-object and add axis titles manually with labs. Use axis.title = "" to remove axis titles.

> Character vector with labels for the model terms, used as axis labels. By default, get\_term\_labels is called to retrieve the labels of the coefficients, which will be used as axis labels. Use axis.labels = "" or auto.label = FALSE to use the variable names as labels instead. If axis.labels is a named vector, axis labels (by default, the names of the model's coefficients) will be matched with the names of axis.label. This ensures that labels always match the related axis value, no matter in which way axis labels are sorted.

Character vector, used as legend title for plots that have a legend.

wrap.title Numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.

wrap.labels Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.

> Numeric vector of length 2, defining the range of the plot axis. Depending on plot-type, may effect either x- or y-axis. For Marginal Effects plots, axis.lim may also be a list of two vectors of length 2, defining axis limits for both the x and y axis.

> Numeric value or vector; if grid.breaks is a single value, sets the distance between breaks for the axis at every grid.breaks'th position, where a major grid line is plotted. If grid.breaks is a vector, values will be used to define the axis positions of the major grid lines.

ri.nr

title

axis.title

axis.labels

legend.title

axis.lim

grid.breaks

ci.lvl

Numeric, the level of the confidence intervals (error bars). Use ci.lvl = NA to remove error bars. For stanneg-models, ci.lvl defines the (outer) probability for the *credible interval* that is plotted (see ci). By default, stanreg-models are printed with two intervals: the "inner" interval, which defaults to the 50%-CI; and the "outer" interval, which defaults to the 89%-CI. ci.lvl affects only the outer interval in such cases. See prob.inner and prob.outer under the ...-argument for more details.

se

Logical, if TRUE, the standard errors are also printed. If robust standard errors are required, use arguments vcov. fun, vcov. type and vcov. args (see robust for details). se overrides ci.lvl: if not NULL, arguments ci.lvl and transform will be ignored. Currently, se only applies to *Coefficients* plots.

vcov.fun

Character vector, indicating the name of the vcov\*()-function from the sandwichpackage, e.g. vcov.fun = "vcovCL", if robust standard errors are required.

vcov.type

Character vector, specifying the estimation type for the robust covariance matrix estimation (see vcovHC for details).

vcov.args

List of named vectors, used as additional arguments that are passed down to vcov.fun.

colors

May be a character vector of color values in hex-format, valid color value names (see demo("colors")) or a name of a pre-defined color palette. Following options are valid for the colors argument:

- If not specified, a default color brewer palette will be used, which is suitable for the plot style.
- If "gs", a greyscale will be used.
- If "bw", and plot-type is a line-plot, the plot is black/white and uses different line types to distinguish groups (see this package-vignette).
- If colors is any valid color brewer palette name, the related palette will be used. Use display.brewer.all to view all available palette names.
- There are some pre-defined color palettes in this package, see sjPlot-themes for details.
- Else specify own color values or names as vector (e.g. colors = "#00ff00" or colors = c("firebrick", "blue")).

show intercept Logical, if TRUE, the intercept of the fitted model is also plotted. Default is FALSE. If transform = "exp", please note that due to exponential transformation of estimates, the intercept in some cases is non-finite and the plot can not be created.

show.values

Logical, whether values should be plotted or not.

show.p

Logical, adds asterisks that indicate the significance level of estimates to the value labels.

show.data

Logical, for *Marginal Effects* plots, also plots the raw data points.

show.legend

For Marginal Effects plots, shows or hides the legend.

show.zeroinf

Logical, if TRUE, shows the zero-inflation part of hurdle- or zero-inflated models.

value.offset

Numeric, offset for text labels to adjust their position relative to the dots or lines.

value.size

Numeric, indicates the size of value labels. Can be used for all plot types where the argument show. values is applicable, e.g. value.size = 4.

jitter Numeric, between 0 and 1. If show.data = TRUE, you can add a small amount of random variation to the location of each data point. jitter then indicates the width, i.e. how much of a bin's width will be occupied by the jittered values. digits Numeric, amount of digits after decimal point when rounding estimates or values. dot.size Numeric, size of the dots that indicate the point estimates. line.size Numeric, size of the lines that indicate the error bars. vline.color Color of the vertical "zero effect" line. Default color is inherited from the current theme. p.threshold Numeric vector of length 3, indicating the treshold for annotating p-values with asterisks. Only applies if p. style = "asterisk". Logical, if TRUE, multiple plots are plotted as grid layout. grid Desired target case. Labels will automatically converted into the specified charcase acter case. See to\_any\_case for more details on this argument. By default, if case is not specified, it will be set to "parsed", unless prefix.labels is not "none". If prefix.labels is either "label" (or "l") or "varname" (or "v") and case is not specified, it will be set to NULL - this is a more convenient default when prefixing labels. auto.label Logical, if TRUE (the default), plot-labels are based on value and variable labels, if the data is labelled. See get\_label and get\_term\_labels for details. If FALSE, original variable names and value labels (factor levels) are used. Indicates whether the value labels of categorical variables should be prefixed, prefix.labels e.g. with the variable name or variable label. See argument prefix in get\_term\_labels for details. bpe For **Stan**-models (fitted with the **rstanarm**- or **brms**-package), the Bayesian point estimate is, by default, the median of the posterior distribution. Use bpe to define other functions to calculate the Bayesian point estimate. bpe needs to be a character naming the specific function, which is passed to the fun-argument in typical\_value. So, bpe = "mean" would calculate the mean value of the posterior distribution. bpe.style For **Stan**-models (fitted with the **rstanarm**- or **brms**-package), the Bayesian point estimate is indicated as a small, vertical line by default. Use bpe.style = "dot" to plot a dot instead of a line for the point estimate. bpe.color Character vector, indicating the color of the Bayesian point estimate. Setting bpe.color = NULL will inherit the color from the mapped aesthetic to match it with the geom's color. ci.style Character vector, defining whether inner and outer intervals for Bayesion models are shown in boxplot-style ("whisker") or in bars with different alpha-levels ("bar"). Other arguments, passed down to various functions. Here is a list of supported arguments and their description in detail. prob.inner and prob.outer For Stan-models (fitted with the rstanarm- or brms-package) and coefficients plot-types, you can specify numeric values

between 0 and 1 for prob. inner and prob. outer, which will then be used

- as inner and outer probabilities for the uncertainty intervals (HDI). By default, the inner probability is 0.5 and the outer probability is 0.89 (unless ci.lvl is specified in this case, ci.lvl is used as outer probability).
- size.inner For **Stan**-models and *Coefficients* plot-types, you can specify the width of the bar for the inner probabilities. Default is 0.1. Setting size.inner = 0 removes the inner probability regions.
- width, alpha, and scale Passed down to geom\_errorbar() or geom\_density\_ridges(), for forest or diagnostic plots.
- width, alpha, dot.alpha, dodge **and** log.y Passed down to plot.ggeffects for *Marginal Effects* plots.
- show.loess Logical, for diagnostic plot-types "slope" and "resid", adds (or hides) a loess-smoothed line to the plot.
- *Marginal Effects* **plot-types** When plotting marginal effects, arguments are also passed down to ggpredict, ggeffect or plot.ggeffects.
- **Case conversion of labels** For case conversion of labels (see argument case), arguments sep\_in and sep\_out will be passed down to to\_any\_case. This only applies to automatically retrieved term labels, *not* if term labels are provided by the axis.labels-argument.

#### **Details**

get\_model\_data() simply calls plot\_model() and returns the data from the ggplot-object. Hence, it is rather inefficient and should be used as alternative to **brooms** tidy()-function only in specific situations.

Some details on the different plot-types:

- type = "std2" Plots standardized beta values, however, standardization follows Gelman's (2008) suggestion, rescaling the estimates by dividing them by two standard deviations instead of just one. Resulting coefficients are then directly comparable for untransformed binary predictors.
- type = "pred" Plots marginal effects. Simply wraps ggpredict. See also this package-vignette.
- type = "eff" Plots marginal effects. Simply wraps ggeffect. See also this package-vignette.
- type = "int" A shortcut for marginal effects plots, where interaction terms are automatically detected and used as terms-argument. Furthermore, if the moderator variable (the second and third term in an interaction) is continuous, type = "int" automatically chooses useful values based on the mdrt.values-argument, which are passed to terms. Then, ggpredict is called. type = "int" plots the interaction term that appears first in the formula along the x-axis, while the second (and possibly third) variable in an interaction is used as grouping factor(s) (moderating variable). Use type = "pred" or type = "eff" and specify a certain order in the terms-argument to indicate which variable(s) should be used as moderator. See also this package-vignette.
- type = "slope" and type = "resid" Simple diagnostic-plots, where a linear model for each single predictor is plotted against the response variable, or the model's residuals. Additionally, a loess-smoothed line is added to the plot. The main purpose of these plots is to check whether the relationship between outcome (or residuals) and a predictor is roughly linear or not. Since the plots are based on a simple linear regression with only one model predictor at the moment, the slopes (i.e. coefficients) may differ from the coefficients of the complete model.

type = "diag" For **Stan-models**, plots the prior versus posterior samples. For **linear (mixed) models**, plots for multicollinearity-check (Variance Inflation Factors), QQ-plots, checks for normal distribution of residuals and homoscedasticity (constant variance of residuals) are shown. For **generalized lineare mixed models**, returns the QQ-plot for random effects.

#### Value

Depending on the plot-type, plot\_model() returns a ggplot-object or a list of such objects. get\_model\_data returns the associated data with the plot-object as tidy data frame, or (depending on the plot-type) a list of such data frames.

#### References

Gelman A (2008) "Scaling regression inputs by dividing by two standard deviations." *Statistics in Medicine* 27: 2865-2873. http://www.stat.columbia.edu/~gelman/research/published/standardizing7.pdf

Aiken and West (1991). Multiple Regression: Testing and Interpreting Interactions.

## **Examples**

```
# prepare data
library(sjmisc)
data(efc)
efc <- to_factor(efc, c161sex, e42dep, c172code)
m < -lm(neg_c_7 \sim pos_v_4 + c12hour + e42dep + c172code, data = efc)
# simple forest plot
plot_model(m)
# grouped coefficients
plot_{model}(m, group.terms = c(1, 2, 3, 3, 4, 4))
# keep only selected terms in the model: pos_v_4, the
# levels 3 and 4 of factor e42dep and levels 2 and 3 for c172code
plot_model(m, terms = c("pos_v_4", "e42dep [3,4]", "c172code [2,3]"))
# multiple plots, as returned from "diagnostic"-plot type,
# can be arranged with 'plot_grid()'
## Not run:
p <- plot_model(m, type = "diag")</pre>
plot_grid(p)
## End(Not run)
# plot random effects
library(lme4)
m <- lmer(Reaction ~ Days + (Days | Subject), sleepstudy)</pre>
plot_model(m, type = "re")
# plot marginal effects
plot_model(m, type = "pred", terms = "Days")
```

```
# plot interactions
## Not run:
m \leftarrow glm(
  tot\_sc\_e \sim c161sex + c172code * neg\_c\_7,
  data = efc,
  family = poisson()
# type = "int" automatically selects groups for continuous moderator
# variables - see argument 'mdrt.values'. The following function call is
# identical to:
# plot_model(m, type = "pred", terms = c("c172code", "neg_c_7 [7,28]"))
plot_model(m, type = "int")
# switch moderator
plot_model(m, type = "pred", terms = c("neg_c_7", "c172code"))
# ggeffects::ggpredict(m, terms = c("neg_c_7", "c172code"))
## End(Not run)
# plot Stan-model
## Not run:
if (require("rstanarm")) {
  data(mtcars)
  m <- stan_glm(mpg ~ wt + am + cyl + gear, data = mtcars, chains = 1)</pre>
  plot_model(m, bpe.style = "dot")
## End(Not run)
```

plot\_models

Forest plot of multiple regression models

## **Description**

Plot and compare regression coefficients with confidence intervals of multiple regression models in one plot.

# Usage

```
plot_models(..., transform, std.est = NULL, rm.terms = NULL,
  title = NULL, m.labels = NULL,
  legend.title = "Dependent Variables", legend.pval.title = "p-level",
  axis.labels = NULL, axis.title = NULL, axis.lim = NULL,
  wrap.title = 50, wrap.labels = 25, wrap.legend.title = 20,
  grid.breaks = NULL, dot.size = 3, spacing = 0.4, colors = "Set1",
  show.values = FALSE, show.legend = TRUE, show.intercept = FALSE,
  show.p = TRUE, p.shape = FALSE, p.threshold = c(0.05, 0.01, 0.001),
  ci.lvl = 0.95, vline.color = NULL, digits = 2, grid = FALSE,
  auto.label = TRUE, prefix.labels = c("none", "varname", "label"))
```

#### **Arguments**

One or more regression models, including glm's or mixed models. May also be a list with fitted models. See 'Examples'.

transform A character vector, naming a function that will be applied on estimates and confidence intervals. By default, transform will automatically use "exp" as transformation for applicable classes of model (e.g. logistic or poisson regression). Estimates of linear models remain untransformed. Use NULL if you want

the raw, non-transformed estimates.

std.est For linear models, choose whether standardized coefficients should be used for plotting. Default is no standardization.

NULL (default) no standardization, returns original estimates.

"std" standardized beta values.

"std2" standardized beta values, however, standardization is done by rescaling estimates by dividing them by two sd (see std\_beta).

Character vector with names that indicate which terms should be removed from rm.terms the plot. Counterpart to terms. rm. terms = "t\_name" would remove the term

t\_name. Default is NULL, i.e. all terms are used. For factors, levels that should be removed from the plot need to be explicitly indicated in square brackets, and match the model's coefficient names, e.g. rm. terms = "t\_name [2,3]" would remove the terms "t\_name2" and "t\_name3" (assuming that the variable t\_name was categorical and has at least the factor levels 2 and 3). Another example for the *iris* dataset would be rm.terms = "Species [versicolor, virginica]".

Note that the rm. terms-argument does not apply to *Marginal Effects* plots.

Character vector, used as plot title. By default, get\_dv\_labels is called to retrieve the label of the dependent variable, which will be used as title. Use title = "" to remove title.

Character vector, used to indicate the different models in the plot's legend. If not specified, the labels of the dependent variables for each model are used.

Character vector, used as legend title for plots that have a legend. legend.title

legend.pval.title

Character vector, used as title of the plot legend that indicates the p-values. Default is "p-level". Only applies if p. shape = TRUE.

Character vector with labels for the model terms, used as axis labels. By default, get\_term\_labels is called to retrieve the labels of the coefficients, which will be used as axis labels. Use axis.labels = "" or auto.label = FALSE to use the variable names as labels instead. If axis.labels is a named vector, axis labels (by default, the names of the model's coefficients) will be matched with the names of axis.label. This ensures that labels always match the related

axis value, no matter in which way axis labels are sorted.

Character vector of length one or two (depending on the plot function and type), used as title(s) for the x and y axis. If not specified, a default labelling is chosen. Note: Some plot types may not support this argument sufficiently. In such cases, use the returned ggplot-object and add axis titles manually with labs. Use axis.title = "" to remove axis titles.

title

m.labels

axis.labels

axis.title

axis.lim Numeric vector of length 2, defining the range of the plot axis. Depending on plot-type, may effect either x- or y-axis. For Marginal Effects plots, axis.lim may also be a list of two vectors of length 2, defining axis limits for both the x and y axis. wrap.title Numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted. wrap.labels Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted. wrap.legend.title numeric, determines how many chars of the legend's title are displayed in one line and when a line break is inserted. Numeric value or vector; if grid.breaks is a single value, sets the distance grid.breaks between breaks for the axis at every grid.breaks'th position, where a major grid line is plotted. If grid.breaks is a vector, values will be used to define the axis positions of the major grid lines. dot.size Numeric, size of the dots that indicate the point estimates. spacing Numeric, spacing between the dots and error bars of the plotted fitted models. Default is 0.3. colors May be a character vector of color values in hex-format, valid color value names (see demo("colors")) or a name of a pre-defined color palette. Following options are valid for the colors argument: • If not specified, a default color brewer palette will be used, which is suitable for the plot style. • If "gs", a greyscale will be used. • If "bw", and plot-type is a line-plot, the plot is black/white and uses different line types to distinguish groups (see this package-vignette). • If colors is any valid color brewer palette name, the related palette will be used. Use display.brewer.all to view all available palette names. • There are some pre-defined color palettes in this package, see sjPlot-themes for details. • Else specify own color values or names as vector (e.g. colors = "#00ff00" or colors = c("firebrick", "blue")). show.values Logical, whether values should be plotted or not. show.legend For Marginal Effects plots, shows or hides the legend. show.intercept Logical, if TRUE, the intercept of the fitted model is also plotted. Default is FALSE. If transform = "exp", please note that due to exponential transformation of estimates, the intercept in some cases is non-finite and the plot can not be created. show.p Logical, adds asterisks that indicate the significance level of estimates to the value labels. Logical, if TRUE, significant levels are distinguished by different point shapes p.shape

and a related legend is plotted. Default is FALSE.

asterisks. Only applies if p. style = "asterisk".

Numeric vector of length 3, indicating the treshold for annotating p-values with

p.threshold

ci.lvl		Numeric, the level of the confidence intervals (error bars). Use ci.lvl = NA to remove error bars. For stanreg-models, ci.lvl defines the (outer) probability for the <i>credible interval</i> that is plotted (see ci). By default, stanreg-models are printed with two intervals: the "inner" interval, which defaults to the 50%-CI; and the "outer" interval, which defaults to the 89%-CI. ci.lvl affects only the outer interval in such cases. See prob.inner and prob.outer under theargument for more details.
vline.c	color	Color of the vertical "zero effect" line. Default color is inherited from the current theme.
digits		Numeric, amount of digits after decimal point when rounding estimates or values.
grid		Logical, if TRUE, multiple plots are plotted as grid layout.
auto.la	abel	Logical, if TRUE (the default), plot-labels are based on value and variable labels, if the data is labelled. See <pre>get_label</pre> and <pre>get_term_labels</pre> for details. If FALSE, original variable names and value labels (factor levels) are used.
prefix.	labels.	Indicates whether the value labels of categorical variables should be prefixed, e.g. with the variable name or variable label. See argument prefix in get_term_labels for details.

## Value

A ggplot-object.

# **Examples**

```
data(efc)
# fit three models
fit1 <- lm(barthtot ~ c160age + c12hour + c161sex + c172code, data = efc)</pre>
fit2 <- lm(neg_c_7 \sim c160age + c12hour + c161sex + c172code, data = efc)
fit3 <- lm(tot_sc_e ~ c160age + c12hour + c161sex + c172code, data = efc)
# plot multiple models
plot_models(fit1, fit2, fit3, grid = TRUE)
# plot multiple models with legend labels and
# point shapes instead of value labels
plot_models(
  fit1, fit2, fit3,
  axis.labels = c(
    "Carer's Age", "Hours of Care", "Carer's Sex", "Educational Status"
  m.labels = c("Barthel Index", "Negative Impact", "Services used"),
  show.values = FALSE, show.p = FALSE, p.shape = TRUE
# plot multiple models from nested lists argument
all.models <- list()
all.models[[1]] \leftarrow fit1
all.models[[2]] \leftarrow fit2
```

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```
all.models[[3]] <- fit3
plot_models(all.models)
# plot multiple models with different predictors (stepwise inclusion),
# standardized estimates
fit1 <- lm(mpg ~ wt + cyl + disp + gear, data = mtcars)
fit2 <- update(fit1, . ~ . + hp)
fit3 <- update(fit2, . ~ . + am)
plot_models(fit1, fit2, fit3, std.est = "std2")</pre>
```

plot\_residuals

Plot predicted values and their residuals

## **Description**

This function plots observed and predicted values of the response of linear (mixed) models for each coefficient and highlights the observed values according to their distance (residuals) to the predicted values. This allows to investigate how well actual and predicted values of the outcome fit across the predictor variables.

## Usage

```
plot_residuals(fit, geom.size = 2, remove.estimates = NULL,
    show.lines = TRUE, show.resid = TRUE, show.pred = TRUE,
    show.ci = FALSE)
```

## Arguments

fit

Fitted linear (mixed) regression model (including objects of class gls or plm).

geom.size

size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.

remove.estimates

Numeric vector with indices (order equals to row index of coef(fit)) or character vector with coefficient names that indicate which estimates should be removed from the table output. The first estimate is the intercept, followed by the model predictors. *The intercept cannot be removed from the table output!* remove.estimates = c(2:4) would remove the 2nd to the 4th estimate (1st to 3rd predictor after intercept) from the output. remove.estimates = "est\_name" would remove the estimate *est\_name*. Default is NULL, i.e. all estimates are printed.

show.lines

Logical, if TRUE, a line connecting predicted and residual values is plotted. Set this argument to FALSE, if plot-building is too time consuming.

show.resid

Logical, if TRUE, residual values are plotted.

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show.pred Logical, if TRUE, predicted values are plotted.

show.ci Logical, if TRUE), adds notches to the box plot, which are used to compare

groups; if the notches of two boxes do not overlap, medians are considered to be

significantly different.

#### Value

A ggplot-object.

#### Note

The actual (observed) values have a coloured fill, while the predicted values have a solid outline without filling.

# **Examples**

```
data(efc)
# fit model
fit <- lm(neg_c_7 ~ c12hour + e17age + e42dep, data = efc)
# plot residuals for all independent variables
plot_residuals(fit)
# remove some independent variables from output
plot_residuals(fit, remove.estimates = c("e17age", "e42dep"))</pre>
```

plot\_scatter

Plot (grouped) scatter plots

#### **Description**

Display scatter plot of two variables. Adding a grouping variable to the scatter plot is possible. Furthermore, fitted lines can be added for each group as well as for the overall plot.

# Usage

```
plot_scatter(data, x, y, grp, title = "", legend.title = NULL,
  legend.labels = NULL, dot.labels = NULL, axis.titles = NULL,
  dot.size = 1.5, label.size = 3, colors = "metro ui",
  fit.line = NULL, fit.grps = NULL, show.rug = FALSE,
  show.legend = TRUE, show.ci = FALSE, wrap.title = 50,
  wrap.legend.title = 20, wrap.legend.labels = 20, jitter = 0.05,
  emph.dots = FALSE, grid = FALSE)
```

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#### **Arguments**

A data frame, or a grouped data frame.

x Name of the variable for the x-axis.

y Name of the variable for the y-axis.

grp Optional, name of the grouping-variable. If not missing, the scatter plot will be

grouped. See 'Examples'.

title Character vector, used as plot title. By default, get\_dv\_labels is called to

retrieve the label of the dependent variable, which will be used as title. Use

title = "" to remove title.

legend.title Character vector, used as legend title for plots that have a legend.

legend.labels character vector with labels for the guide/legend.

dot.labels Character vector with names for each coordinate pair given by x and y, so text

labels are added to the plot. Must be of same length as x and y. If dot.labels has a different length, data points will be trimmed to match dot.labels. If

dot.labels = NULL (default), no labels are printed.

axis.titles character vector of length one or two, defining the title(s) for the x-axis and

y-axis.

dot.size Numeric, size of the dots that indicate the point estimates.

label.size Size of text labels if argument dot.labels is used.

colors May be a character vector of color values in hex-format, valid color value names (see demo("colors")) or a name of a pre-defined color palette. Following op-

tions are valid for the colors argument:

• If not specified, a default color brewer palette will be used, which is suitable for the plot style.

- If "gs", a greyscale will be used.
- If "bw", and plot-type is a line-plot, the plot is black/white and uses different line types to distinguish groups (see this package-vignette).
- If colors is any valid color brewer palette name, the related palette will be used. Use display.brewer.all to view all available palette names.
- There are some pre-defined color palettes in this package, see sjPlot-themes for details.
- Else specify own color values or names as vector (e.g. colors = "#00ff00" or colors = c("firebrick", "blue")).

fit.line, fit.grps

Specifies the method to add a fitted line accross the data points. Possible values are for instance "lm", "glm", "loess" or "auto". If NULL, no line is plotted. fit.line adds a fitted line for the complete data, while fit.grps adds a fitted line for each subgroup of grp.

show.rug Logical, if TRUE, a marginal rug plot is displayed in the graph.

show.legend For Marginal Effects plots, shows or hides the legend.

show.ci Logical, if TRUE), adds notches to the box plot, which are used to compare groups; if the notches of two boxes do not overlap, medians are considered to be

significantly different.

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wrap.title Numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.

wrap.legend.title

numeric, determines how many chars of the legend's title are displayed in one line and when a line break is inserted.

wrap.legend.labels

numeric, determines how many chars of the legend labels are displayed in one

line and when a line break is inserted.

jitter Numeric, between 0 and 1. If show.data = TRUE, you can add a small amount

of random variation to the location of each data point. jitter then indicates the width, i.e. how much of a bin's width will be occupied by the jittered values.

emph. dots Logical, if TRUE, overlapping points at same coordinates will be becomme larger,

so point size indicates amount of overlapping.

grid Logical, if TRUE, multiple plots are plotted as grid layout.

#### Value

A ggplot-object. For grouped data frames, a list of ggplot-objects for each group in the data.

## **Examples**

```
# load sample date
library(sjmisc)
library(sjlabelled)
data(efc)
# simple scatter plot
plot_scatter(efc, e16sex, neg_c_7)
# simple scatter plot, increased jittering
plot_scatter(efc, e16sex, neg_c_7, jitter = .4)
# grouped scatter plot
plot_scatter(efc, c160age, e17age, e42dep)
# grouped scatter plot with marginal rug plot
# and add fitted line for complete data
plot_scatter(
  efc, c12hour, c160age, c172code,
  show.rug = TRUE, fit.line = "lm"
# grouped scatter plot with marginal rug plot
# and add fitted line for each group
plot_scatter(
  efc, c12hour, c160age, c172code,
  show.rug = TRUE, fit.grps = "loess",
  grid = TRUE
)
```

save\_plot 37

save_plot	Save ggplot-figure for print publication	

# Description

Convenient function to save the last ggplot-figure in high quality for publication.

### Usage

```
save_plot(filename, fig = last_plot(), width = 12, height = 9,
    dpi = 300, theme = theme_get(), label.color = "black",
    label.size = 2.4, axis.textsize = 0.8, axis.titlesize = 0.75,
    legend.textsize = 0.6, legend.titlesize = 0.65,
    legend.itemsize = 0.5)
```

## **Arguments**

filename	Name of the output file; filename must end with one of the following accepted file types: ".png", ".jpg", ".svg" or ".tif".	
fig	The plot that should be saved. By default, the last plot is saved.	
width	Width of the figure, in centimetres.	
height	Height of the figure, in centimetres.	
dpi	Resolution in dpi (dots per inch). Ignored for vector formats, such as ".svg".	
theme	The default theme to use when saving the plot.	
label.color	Color value for labels (axis, plot, etc.).	
label.size	Fontsize of value labels inside plot area.	
axis.textsize	Fontsize of axis labels.	
axis.titlesize legend.textsize	Fontsize of axis titles.	
	Fontsize of legend labels.	
legend.titlesize		
	Fontsize of legend title.	
legend.itemsize		
	Size of legend's item (legend key), in centimetres.	

# Note

This is a convenient function with some default settings that should come close to most of the needs for fontsize and scaling in figures when saving them for printing or publishing. It uses cairographics anti-aliasing (see png).

For adjusting plot appearance, see also sjPlot-themes.

set\_theme

Set global theme options for sjp-functions

### **Description**

Set global theme options for sjp-functions.

#### Usage

```
set_theme(base = theme_grey(), theme.font = NULL,
 title.color = "black", title.size = 1.2, title.align = "left",
 title.vjust = NULL, geom.outline.color = NULL,
 geom.outline.size = 0, geom.boxoutline.size = 0.5,
 geom.boxoutline.color = "black", geom.alpha = 1, geom.linetype = 1,
 geom.errorbar.size = 0.7, geom.errorbar.linetype = 1,
 geom.label.color = NULL, geom.label.size = 4, geom.label.alpha = 1,
 geom.label.angle = 0, axis.title.color = "grey30",
 axis.title.size = 1.1, axis.title.x.vjust = NULL,
 axis.title.y.vjust = NULL, axis.angle.x = 0, axis.angle.y = 0,
 axis.angle = NULL, axis.textcolor.x = "grey30",
 axis.textcolor.y = "grey30", axis.textcolor = NULL,
 axis.linecolor.x = NULL, axis.linecolor.y = NULL,
 axis.linecolor = NULL, axis.line.size = 0.5, axis.textsize.x = 1,
 axis.textsize.y = 1, axis.textsize = NULL, axis.tickslen = NULL,
 axis.tickscol = NULL, axis.ticksmar = NULL, axis.ticksize.x = NULL,
 axis.ticksize.y = NULL, panel.backcol = NULL,
 panel.bordercol = NULL, panel.col = NULL,
 panel.major.gridcol = NULL, panel.minor.gridcol = NULL,
 panel.gridcol = NULL, panel.gridcol.x = NULL,
 panel.gridcol.y = NULL, panel.major.linetype = 1,
 panel.minor.linetype = 1, plot.backcol = NULL,
 plot.bordercol = NULL, plot.col = NULL, plot.margins = NULL,
 legend.pos = "right", legend.just = NULL, legend.inside = FALSE,
 legend.size = 1, legend.color = "black", legend.title.size = 1,
 legend.title.color = "black", legend.title.face = "bold",
 legend.backgroundcol = "white", legend.bordercol = "white",
 legend.item.size = NULL, legend.item.backcol = "grey90",
 legend.item.bordercol = "white")
```

# **Arguments**

base	base theme where theme is built on. By default, all metrics from theme_gray() are used. See 'Details'.
theme.font	base font family for the plot.
title.color	Color of plot title. Default is "black".
title.size	size of plot title. Default is 1.3.

alignment of plot title. Must be one of "left" (default), "center" or "right". title.align You may use initial letter only. numeric, vertical adjustment for plot title. title.vjust geom.outline.color Color of geom outline. Only applies, if geom.outline.size is larger than 0. geom.outline.size size of bar outlines. Default is 0.1. Use size of 0 to remove geom outline. geom.boxoutline.size size of outlines and median bar especially for boxplots. Default is 0.5. Use size of 0 to remove boxplot outline. geom.boxoutline.color Color of outlines and median bar especially for boxplots. Only applies, if geom.boxoutline.size is larger than 0. geom.alpha specifies the transparancy (alpha value) of geoms geom.linetype linetype of line geoms. Default is 1 (solid line). geom.errorbar.size size (thickness) of error bars. Default is 0.8 geom.errorbar.linetype linetype of error bars. Default is 1 (solid line). geom.label.color Color of geom's value and annotation labels geom.label.size size of geom's value and annotation labels geom.label.alpha alpha level of geom's value and annotation labels geom.label.angle angle of geom's value and annotation labels axis.title.color Color of x- and y-axis title labels axis.title.size size of x- and y-axis title labels axis.title.x.vjust numeric, vertical adjustment of x-axis-title. axis.title.y.vjust numeric, vertical adjustment of y-axis-title. angle for x-axis labels axis.angle.x axis.angle.y angle for y-axis labels axis.angle angle for x- and y-axis labels. If set, overrides both axis.angle.x and axis.angle.y axis.textcolor.x Color for x-axis labels. If not specified, a default dark gray color palette will be used for the labels. axis.textcolor.y Color for y-axis labels. If not specified, a default dark gray color palette will be

used for the labels.

axis.textcolor Color for both x- and y-axis labels. If set, overrides both axis.textcolor.x and axis.textcolor.y axis.linecolor.x Color of x-axis border axis.linecolor.y Color of y-axis border axis.linecolor Color for both x- and y-axis borders. If set, overrides both axis.linecolor.x and axis.linecolor.y. axis.line.size size (thickness) of axis lines. Only affected, if axis.linecolor is set. axis.textsize.x size of x-axis labels axis.textsize.y size of y-axis labels axis.textsize size for both x- and y-axis labels. If set, overrides both axis.textsize.x and axis.textsize.y. length of axis tick marks axis.tickslen axis.tickscol Color of axis tick marks margin between axis labels and tick marks axis.ticksmar axis.ticksize.x size of tick marks at x-axis. axis.ticksize.y size of tick marks at y-axis. panel.backcol Color of the diagram's background panel.bordercol Color of whole diagram border (panel border) Color of both diagram's border and background. If set, overrides both panel.bordercol panel.col and panel.backcol. panel.major.gridcol Color of the major grid lines of the diagram background panel.minor.gridcol Color of the minor grid lines of the diagram background panel.gridcol Color for both minor and major grid lines of the diagram background. If set, overrides both panel.major.gridcol and panel.minor.gridcol. panel.gridcol.x See panel.gridcol. panel.gridcol.y See panel.gridcol. panel.major.linetype line type for major grid lines panel.minor.linetype line type for minor grid lines plot.backcol Color of the plot's background plot.bordercol Color of whole plot's border (panel border)

plot.col Color of both plot's region border and background. If set, overrides both plot.backcol

and plot.bordercol.

numeric vector of length 4, indicating the top, right, bottom and left margin of plot.margins

the plot region.

legend.pos position of the legend, if a legend is drawn.

> legend outside plot Use "bottom", "top", "left" or "right" to position the legend above, below, on the left or right side of the diagram. Right posi-

tioning is default.

legend inside plot If legend.inside = TRUE, legend can be placed inside plot. Use "top left", "top right", "bottom left" and "bottom right" to position legend in any of these corners, or a two-element numeric vector

with values from 0-1. See also legend.inside.

legend.just justification of legend, relative to its position ("center" or two-element nu-

> meric vector with values from 0-1. By default (outside legend), justification is centered. If legend is inside and justification not specified, legend justification

is set according to legend position.

logical, use TRUE to put legend inside the plotting area. See legend.pos. legend.inside

legend.size text size of the legend. Default is 1. Relative size, so recommended values are

from 0.3 to 2.5

legend.color Color of the legend labels

legend.title.size

text size of the legend title

legend.title.color

Color of the legend title

legend.title.face

font face of the legend title. By default, "bold" face is used.

legend.backgroundcol

fill color of the legend's background. Default is "white", so no visible back-

ground is drawn.

legend.bordercol

Color of the legend's border. Default is "white", so no visible border is drawn.

legend.item.size

size of legend's item (legend key), in centimetres.

legend.item.backcol

fill color of the legend's item-background. Default is "grey90".

legend.item.bordercol

Color of the legend's item-border. Default is "white".

### Value

The customized theme object, or NULL, if a ggplot-theme was used.

#### See Also

sjPlot-themes

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### **Examples**

```
## Not run:
library(sjmisc)
data(efc)
# set sjPlot-defaults, a slightly modification
# of the ggplot base theme
set_theme()
# legends of all plots inside
set_theme(legend.pos = "top left", legend.inside = TRUE)
sjp.xtab(efc$e42dep, efc$e16sex)
# Use classic-theme. you may need to
# load the ggplot2-library.
library(ggplot2)
set_theme(base = theme_classic())
plot_frq(efc$e42dep)
# adjust value labels
set_theme(
  geom.label.size = 3.5,
  geom.label.color = "#3366cc",
  geom.label.angle = 90
)
# hjust-aes needs adjustment for this
update_geom_defaults('text', list(hjust = -0.1))
sjp.xtab(efc$e42dep, efc$e16sex, vjust = "center", hjust = "center")
# Create own theme based on classic-theme
set_theme(
  base = theme_classic(), axis.linecolor = "grey50",
  axis.textcolor = "#6699cc"
plot_frq(efc$e42dep)
## End(Not run)
```

sjc.cluster

Compute hierarchical or kmeans cluster analysis

## **Description**

Compute hierarchical or kmeans cluster analysis and return the group association for each observation as vector.

### Usage

```
sjc.cluster(data, groupcount = NULL, method = c("hclust", "kmeans"),
```

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```
distance = c("euclidean", "maximum", "manhattan", "canberra", "binary",
"minkowski"), agglomeration = c("ward", "ward.D", "ward.D2", "single",
"complete", "average", "mcquitty", "median", "centroid"),
iter.max = 20, algorithm = c("Hartigan-Wong", "Lloyd", "MacQueen"))
```

## **Arguments**

data

A data frame with variables that should be used for the cluster analysis.

groupcount

Amount of groups (clusters) used for the cluster solution. May also be a set of initial (distinct) cluster centres, in case method = "kmeans" (see kmeans for details on centers argument). If groupcount = NULL and method = "kmeans", the optimal amount of clusters is calculated using the gap statistics (see sjc.kgap). For method = "hclust", groupcount needs to be specified. Following functions may be helpful for estimating the amount of clusters:

- Use sjc.elbow to determine the group-count depending on the elbowcriterion.
- If method = "kmeans", use sjc.kgap to determine the group-count according to the gap-statistic.
- If method = "hclust" (hierarchical clustering, default), use sjc.dend to inspect different cluster group solutions.
- Use sjc.grpdisc to inspect the goodness of grouping (accuracy of classification).

method

Method for computing the cluster analysis. By default ("kmeans"), a kmeans cluster analysis will be computed. Use "hclust" to compute a hierarchical cluster analysis. You can specify the initial letters only.

distance

Distance measure to be used when method = "hclust" (for hierarchical clustering). Must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". See dist. If is method = "kmeans" this argument will be ignored.

agglomeration

Agglomeration method to be used when method = "hclust" (for hierarchical clustering). This should be one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid". Default is "ward" (see hclust). If method = "kmeans" this argument will be ignored. See 'Note'.

iter.max

Maximum number of iterations allowed. Only applies, if method = "kmeans". See kmeans for details on this argument.

algorithm

Algorithm used for calculating kmeans cluster. Only applies, if method = "kmeans". May be one of "Hartigan-Wong" (default), "Lloyd" (used by SPSS), or "MacQueen". See kmeans for details on this argument.

#### Value

The group classification for each observation as vector. This group classification can be used for sjc.grpdisc-function to check the goodness of classification. The returned vector includes missing values, so it can be appended to the original data frame data.

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#### Note

Since R version > 3.0.3, the "ward" option has been replaced by either "ward.D" or "ward.D2", so you may use one of these values. When using "ward", it will be replaced by "ward.D2".

To get similar results as in SPSS Quick Cluster function, following points have to be considered:

- Use the /PRINT INITIAL option for SPSS Quick Cluster to get a table with initial cluster centers.
- 2. Create a matrix of this table, by consecutively copying the values, one row after another, from the SPSS output into a matrix and specify nrow and ncol arguments.
- 3. Use algorithm="Lloyd".
- 4. Use the same amount of iter.max both in SPSS and this sjc.qclus.

This ensures a fixed initial set of cluster centers (as in SPSS), while kmeans in R always selects initial cluster sets randomly.

#### References

Maechler M, Rousseeuw P, Struyf A, Hubert M, Hornik K (2014) cluster: Cluster Analysis Basics and Extensions. R package.

### **Examples**

```
# Hierarchical clustering of mtcars-dataset
groups <- sjc.cluster(mtcars, 5)

# K-means clustering of mtcars-dataset
groups <- sjc.cluster(mtcars, 5, method="k")</pre>
```

sjc.dend

Compute hierarchical cluster analysis and visualize group classification

#### **Description**

Computes a hierarchical cluster analysis and plots a hierarchical dendrogram with highlighted rectangles around the classified groups. Can be used, for instance, as visual tool to verify the elbowcriterion (see sjc.elbow).

### Usage

```
sjc.dend(data, groupcount, distance = "euclidean",
   agglomeration = "ward")
```

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### Arguments

data

A data frame with variables that should be used for the cluster analysis.

groupcount

The amount of groups (clusters) that should be used.

- Use sjc.elbow-function to determine the group-count depending on the elbow-criterion.
- Use sjc.grpdisc-function to inspect the goodness of grouping (accuracy of classification).

Solutions for multiple cluster groups can be plotted, for instance with "groupcount = c(3:6)".

distance

Distance measure to be used when method = "hclust" (for hierarchical clustering). Must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". See dist. If is method = "kmeans" this argument

will be ignored.

agglomeration

Agglomeration method to be used when method = "hclust" (for hierarchical clustering). This should be one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid". Default is "ward" (see hclust). If method = "kmeans" this argument will be ignored. See 'Note'.

#### Note

Since R version > 3.0.3, the "ward" option has been replaced by either "ward.D" or "ward.D2", so you may use one of these values. When using "ward", it will be replaced by "ward.D2".

#### **Examples**

```
# Plot dendrogram of hierarchical clustering of mtcars-dataset
# and show group classification
sjc.dend(mtcars, 5)
# Plot dendrogram of hierarchical clustering of mtcars-dataset
# and show group classification for 2 to 4 groups
sjc.dend(mtcars, 2:4)
```

sjc.elbow

Compute elbow values of a k-means cluster analysis

#### **Description**

Plot elbow values of a k-means cluster analysis. This function computes a k-means cluster analysis on the provided data frame and produces two plots: one with the different elbow values and a second plot that maps the differences between each "step" (i.e. between elbow values) on the y-axis. An increase in the second plot may indicate the elbow criterion.

#### Usage

```
sjc.elbow(data, steps = 15, show.diff = FALSE)
```

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## **Arguments**

data	data frame containing all variables that should be used for determining the elbow criteria
steps	maximum group-count for the k-means cluster analysis for which the elbow-criterion should be displayed. Default is 15.
show.diff	logical, if TRUE, an additional plot with the differences between each fusion step of the Elbow criterion calculation is shown. This plot may help identifying the "elbow". Default for this argument is FALSE.

## **Examples**

```
# plot elbow values of mtcars dataset
sjc.elbow(mtcars)
```

sjc.grpdisc	Compute a linear discriminant analysis on classified cluster groups

# Description

Computes linear discriminant analysis on classified cluster groups. This function plots a bar graph indicating the goodness of classification for each group.

# Usage

```
sjc.grpdisc(data, groups, groupcount, clss.fit = TRUE)
```

# Arguments

data	A data frame with variables that should be used for the cluster analysis.
groups	group classification of the cluster analysis that was returned from the $\verb+sjc.cluster-function$
groupcount	amount of groups (clusters) that should be used. Use sjc.elbow to determine the group-count depending on the elbow-criterion.
clss.fit	logical, if TRUE (default), a vertical line indicating the overall goodness of classification is added to the plot, so one can see whether a certain group is below or above the average classification goodness.

#### Value

(Invisibly) returns an object with

- data: the used data frame for plotting,
- plot: the ggplot object,
- accuracy: a vector with the accuracy of classification for each group,
- total.accuracy: the total accuracy of group classification.

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### **Examples**

```
# retrieve group classification from hierarchical cluster analysis
# on the mtcars data set (5 groups)
groups <- sjc.cluster(mtcars, 5)

# plot goodness of group classification
sjc.grpdisc(mtcars, groups, 5)</pre>
```

sjc.kgap

Compute gap statistics for k-means-cluster

### **Description**

An implementation of the gap statistic algorithm from Tibshirani, Walther, and Hastie's "Estimating the number of clusters in a data set via the gap statistic". This function calls the clusGap-function of the cluster-package to calculate the data for the plot.

### Usage

```
sjc.kgap(x, max = 10, B = 100, SE.factor = 1,
method = "Tibs2001SEmax", plotResults = TRUE)
```

### **Arguments**

Χ	matrix, where rows are observations and columns are individual dimensions, to
	compute and plot the gap statistic (according to a uniform reference distribu-

tion).

max maximum number of clusters to consider, must be at least two. Default is 10.

B integer, number of Monte Carlo ("bootstrap") samples. Default is 100.

SE. factor [When method contains "SE"] Determining the optimal number of clusters, Tib-

shirani et al. proposed the "1 S.E."-rule. Using an SE.factor f, the "f S.E."-rule

is used, more generally.

method character string indicating how the "optimal" number of clusters, k^, is com-

puted from the gap statistics (and their standard deviations), or more generally how the location  $k^{\wedge}$  of the maximum of f[k] should be determined. Default is

"Tibs2001SEmax". Possible value are:

"globalmax" simply corresponds to the global maximum, i.e., is which.max(f).

"firstmax" gives the location of the first local maximum.

"Tibs2001SEmax" uses the criterion, Tibshirani et al(2001) proposed: "the smallest k such that  $f(k) >= f(k+1) - s_k+1$ ". Note that this chooses k = 1 when all standard deviations are larger than the differences f(k+1) - f(k).

"firstSEmax" is the location of the first f() value which is not larger than the first local maximum minus SE.factor \* SE.f[], i.e, within an "f S.E." range of that maximum (see also SE.factor).

"globalSEmax" (used in Dudoit and Fridlyand (2002), supposedly following Tibshirani's proposition) is the location of the first f() value which is not larger than the global maximum minus SE.factor \* SE.f[], i.e, within an "f S.E." range of that maximum (see also SE.factor).

plotResults

logical, if TRUE (default), a graph visualiting the gap statistic will be plotted. Use FALSE to omit the plot.

#### Value

An object containing the used data frame for plotting, the ggplot object and the number of found cluster.

#### References

- Tibshirani R, Walther G, Hastie T (2001) Estimating the number of clusters in a data set via gap statistic. J. R. Statist. Soc. B, 63, Part 2, pp. 411-423
- Maechler, M., Rousseeuw, P., Struyf, A., Hubert, M., Hornik, K.(2013). cluster: Cluster Analysis Basics and Extensions. R package version 1.14.4. (web)

#### See Also

```
sjc.elbow
```

# Examples

```
## Not run:
# plot gap statistic and determine best number of clusters
# in mtcars dataset
sjc.kgap(mtcars)
# and in iris dataset
sjc.kgap(iris[,1:4])
## End(Not run)
```

sjc.qclus

Compute quick cluster analysis

# Description

Compute a quick kmeans or hierarchical cluster analysis and displays "cluster characteristics" as plot.

#### Usage

```
sjc.qclus(data, groupcount = NULL, groups = NULL,
 method = c("kmeans", "hclust"), distance = c("euclidean", "maximum",
 "manhattan", "canberra", "binary", "minkowski"),
 agglomeration = c("ward", "ward.D", "ward.D2", "single", "complete",
 "average", "mcquitty", "median", "centroid"), iter.max = 20,
 algorithm = c("Hartigan-Wong", "Lloyd", "MacQueen"),
 show.accuracy = FALSE, title = NULL, axis.labels = NULL,
 wrap.title = 40, wrap.labels = 20, wrap.legend.title = 20,
 wrap.legend.labels = 20, facet.grid = FALSE,
 geom.colors = "Paired", geom.size = 0.5, geom.spacing = 0.1,
 show.legend = TRUE, show.grpcnt = TRUE, legend.title = NULL,
 legend.labels = NULL, coord.flip = FALSE, reverse.axis = FALSE)
```

### **Arguments**

data

A data frame with variables that should be used for the cluster analysis.

groupcount

Amount of groups (clusters) used for the cluster solution. May also be a set of initial (distinct) cluster centres, in case method = "kmeans" (see kmeans for details on centers argument). If groupcount = NULL and method = "kmeans", the optimal amount of clusters is calculated using the gap statistics (see sjc.kgap). For method = "hclust", groupcount needs to be specified. Following functions may be helpful for estimating the amount of clusters:

- Use sjc.elbow to determine the group-count depending on the elbowcriterion.
- If method = "kmeans", use sjc.kgap to determine the group-count according to the gap-statistic.
- If method = "hclust" (hierarchical clustering, default), use sjc.dend to inspect different cluster group solutions.
- Use sjc.grpdisc to inspect the goodness of grouping (accuracy of classification).

groups

Optional, by default, this argument is NULL and will be ignored. However, to plot existing cluster groups, specify groupcount and groups. groups is a vector of same length as nrow(data) and indicates the group classification of the cluster analysis. The group classification can be computed with the sjc.cluster function. See 'Examples'.

method

Method for computing the cluster analysis. By default ("kmeans"), a kmeans cluster analysis will be computed. Use "hclust" to compute a hierarchical cluster analysis. You can specify the initial letters only.

distance

Distance measure to be used when method = "hclust" (for hierarchical clustering). Must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". See dist. If is method = "kmeans" this argument will be ignored.

agglomeration

Agglomeration method to be used when method = "hclust" (for hierarchical clustering). This should be one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid". Default is "ward" (see hclust). If method = "kmeans" this argument will be ignored. See 'Note'.

iter.max	Maximum number of iterations allowed. Only applies, if method = "kmeans". See kmeans for details on this argument.
algorithm	Algorithm used for calculating kmeans cluster. Only applies, if method = "kmeans". May be one of "Hartigan-Wong" (default), "Lloyd" (used by SPSS), or "MacQueen". See kmeans for details on this argument.
show.accuracy	Logical, if TRUE, the sjc.grpdisc function will be called, which computes a linear discriminant analysis on the classified cluster groups and plots a bar graph indicating the goodness of classification for each group.
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
wrap.legend.ti	tle
	numeric, determines how many chars of the legend's title are displayed in one line and when a line break is inserted.
wrap.legend.la	
	numeric, determines how many chars of the legend labels are displayed in one line and when a line break is inserted.
facet.grid	TRUE to arrange the lay out of of multiple plots in a grid of an integrated single plot. This argument calls facet_wrap or facet_grid to arrange plots. Use plot_grid to plot multiple plot-objects as an arranged grid with grid.arrange.
geom.colors	user defined color for geoms. See 'Details' in sjp. grpfrq.
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.
geom.spacing	the spacing between geoms (i.e. bar spacing)
show.legend	logical, if TRUE, and depending on plot type and function, a legend is added to the plot.
show.grpcnt	Logical, if TRUE (default), the count within each cluster group is added to the legend labels (e.g. "Group 1 (n=87)").
legend.title	character vector, used as title for the plot legend.
legend.labels	character vector with labels for the guide/legend.
coord.flip	logical, if TRUE, the x and y axis are swapped.
reverse.axis	Logical, if TRUE, the values on the x-axis are reversed.

#### **Details**

Following steps are computed in this function:

1. If method = "kmeans", this function first determines the optimal group count via gap statistics (unless argument groupcount is specified), using the sjc.kgap function.

- 2. A cluster analysis is performed by running the sjc.cluster function to determine the cluster groups.
- 3. Then, all variables in data are scaled and centered. The mean value of these z-scores within each cluster group is calculated to see how certain characteristics (variables) in a cluster group differ in relation to other cluster groups.
- 4. These results are plotted as graph.

This method can also be used to plot existing cluster solution as graph witouth computing a new cluster analysis. See argument groups for more details.

### Value

(Invisibly) returns an object with

- data: the used data frame for plotting,
- plot: the ggplot object,
- groupcount: the number of found cluster (as calculated by sjc.kgap)
- classification: the group classification (as calculated by sjc.cluster), including missing values, so this vector can be appended to the original data frame.
- accuracy: the accuracy of group classification (as calculated by sjc.grpdisc).

## Note

```
See 'Note' in sjc.cluster
```

### References

Maechler M, Rousseeuw P, Struyf A, Hubert M, Hornik K (2014) cluster: Cluster Analysis Basics and Extensions. R package.

## **Examples**

```
## Not run:
# k-means clustering of mtcars-dataset
sjc.qclus(mtcars)

# k-means clustering of mtcars-dataset with 4 pre-defined
# groups in a faceted panel
sjc.qclus(airquality, groupcount = 4, facet.grid = TRUE)
## End(Not run)

# k-means clustering of airquality data
# and saving the results. most likely, 3 cluster
# groups have been found (see below).
```

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```
airgrp <- sjc.qclus(airquality)
# "re-plot" cluster groups, without computing
# new k-means cluster analysis.
sjc.qclus(airquality, groupcount = 3, groups = airgrp$classification)</pre>
```

sjp.aov1

Plot One-Way-Anova tables

## **Description**

Plot One-Way-Anova table sum of squares (SS) of each factor level (group) against the dependent variable. The SS of the factor variable against the dependent variable (variance within and between groups) is printed to the model summary.

## Usage

```
sjp.aov1(var.dep, var.grp, meansums = FALSE, title = NULL,
   axis.labels = NULL, rev.order = FALSE,
   string.interc = "(Intercept)", axis.title = "", axis.lim = NULL,
   geom.colors = c("#3366a0", "#aa3333"), geom.size = 3,
   wrap.title = 50, wrap.labels = 25, grid.breaks = NULL,
   show.values = TRUE, digits = 2, y.offset = 0.15, show.p = TRUE,
   show.summary = FALSE)
```

## **Arguments**

var.dep	Dependent variable. Will be used with following formula: aov(var.dep ~ var.grp)
var.grp	Factor with the cross-classifying variable, where var.dep is grouped into the categories represented by var.grp.
meansums	Logical, if TRUE, the values reported are the true group mean values. If FALSE (default), the values are reported in the standard way, i.e. the values indicate the difference of the group mean in relation to the intercept (reference group).
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
rev.order	Logical, if TRUE, order of categories (groups) is reversed.
string.interc	Character vector that indicates the reference group (intercept), that is appended to the value label of the grouping variable. Default is "(Intercept)".
axis.title	Character vector of length one or two (depending on the plot function and type), used as title(s) for the x and y axis. If not specified, a default labelling is chosen. <b>Note:</b> Some plot types may not support this argument sufficiently. In such cases, use the returned ggplot-object and add axis titles manually with labs. Use axis.title = "" to remove axis titles.

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axis.lim	Numeric vector of length 2, defining the range of the plot axis. Depending on plot type, may effect either x- or y-axis, or both. For multiple plot outputs (e.g., from type = "eff" or type = "slope" in plot_model), axis.lim may also be a list of vectors of length 2, defining axis limits for each plot (only if non-faceted).
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
grid.breaks	numeric; sets the distance between breaks for the axis, i.e. at every grid.breaks'th position a major grid is being printed.
show.values	Logical, whether values should be plotted or not.
digits	Numeric, amount of digits after decimal point when rounding estimates or values.
y.offset	numeric, offset for text labels when their alignment is adjusted to the top/bottom of the geom (see hjust and vjust).
show.p	Logical, adds significance levels to values, or value and variable labels.
show.summary	logical, if TRUE (default), a summary with chi-squared statistics (see chisq.test), Cramer's V or Phi-value etc. is shown. If a cell contains expected values lower than five (or lower than 10 if df is 1), the Fisher's excact test (see fisher.test) is computed instead of chi-squared test. If the table's matrix is larger than 2x2, Fisher's excact test with Monte Carlo simulation is computed.

# Value

A ggplot-object.

# Examples

```
data(efc)
# note: "var.grp" does not need to be a factor.
# coercion to factor is done by the function
sjp.aov1(efc$c12hour, efc$e42dep)
```

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sjp.chi2	Plot Pearson's Chi2-Test of multiple contingency tables
• .	

# Description

Plot p-values of Pearson's Chi2-tests for multiple contingency tables as ellipses or tiles. Requires a data frame with dichotomous (dummy) variables. Calculation of Chi2-matrix taken from Tales of R.

# Usage

```
sjp.chi2(df, title = "Pearson's Chi2-Test of Independence",
   axis.labels = NULL, wrap.title = 50, wrap.labels = 20,
   show.legend = FALSE, legend.title = NULL)
```

## **Arguments**

df	A data frame with (dichotomous) factor variables.
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If $title = ""$ , no title is printed. For effect-plots, may also be a character vector of length $> 1$ , to define titles for each sub-plot or facet.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
show.legend	logical, if TRUE, and depending on plot type and function, a legend is added to the plot.
legend.title	character vector, used as title for the plot legend.

### Value

A ggplot-object.

### See Also

Tales of R.

# **Examples**

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```
as.factor(sample(1:2, 100, replace=TRUE)))
# create variable labels
items <- list(c("Item 1", "Item 2", "Item 3", "Item 4", "Item 5"))
# plot Chi2-contingency-table
sjp.chi2(mydf, axis.labels = items)</pre>
```

sjp.corr

Plot correlation matrix

## **Description**

Plot correlation matrix as ellipses or tiles.

# Usage

```
sjp.corr(data, title = NULL, axis.labels = NULL, sort.corr = TRUE,
  decimals = 3, na.deletion = c("listwise", "pairwise"),
  corr.method = c("pearson", "spearman", "kendall"),
  geom.colors = "RdBu", wrap.title = 50, wrap.labels = 20,
  show.legend = FALSE, legend.title = NULL, show.values = TRUE,
  show.p = TRUE, p.numeric = FALSE)
```

### **Arguments**

data	Matrix with correlation coefficients as returned by the cor-function, or a data. frame of variables where correlations between columns should be computed.
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
sort.corr	Logical, if TRUE (default), the axis labels are sorted according to the correlation strength. If FALSE, axis labels appear in order of how variables were included in the cor-computation or data frame.
decimals	Indicates how many decimal values after comma are printed when the values labels are shown. Default is 3. Only applies when show.values = TRUE.
na.deletion	Indicates how missing values are treated. May be either "listwise" (default) or "pairwise". May be abbreviated.
corr.method	Indicates the correlation computation method. May be one of "spearman" (default), "pearson" or "kendall". May be abbreviated.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.

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wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
show.legend	logical, if TRUE, and depending on plot type and function, a legend is added to the plot.
legend.title	character vector, used as title for the plot legend.
show.values	Logical, whether values should be plotted or not.
show.p	Logical, adds significance levels to values, or value and variable labels.
p.numeric	Logical, if TRUE, the p-values are printed as numbers. If FALSE (default), asterisks are used.

#### **Details**

Required argument is either a data. frame or a matrix with correlation coefficients as returned by the cor-function. In case of ellipses, the ellipses size indicates the strength of the correlation. Furthermore, blue and red colors indicate positive or negative correlations, where stronger correlations are darker.

# Value

(Insisibily) returns the ggplot-object with the complete plot (plot) as well as the data frame that was used for setting up the ggplot-object (df) and the original correlation matrix (corr.matrix).

## Note

If data is a matrix with correlation coefficients as returned by the cor-function, p-values can't be computed. Thus, show.p and p.numeric only have an effect if data is a data.frame.

# See Also

```
sjt.corr
```

## **Examples**

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sjp.fa

Plot FA results

## Description

Performes a maximum likelihood factor analysis on a data frame or matrix and plots the factor solution as ellipses or tiles.

In case a data frame is used as argument, the cronbach's alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension.

### Usage

```
sjp.fa(data, rotation = c("promax", "varimax"), method = c("ml",
   "minres", "wls", "gls", "pa", "minchi", "minrank"), nmbr.fctr = NULL,
   fctr.load.tlrn = 0.1, digits = 2, title = NULL,
   axis.labels = NULL, type = c("bar", "circle", "tile"),
   geom.size = 0.6, geom.colors = "RdBu", wrap.title = 50,
   wrap.labels = 30, show.values = TRUE, show.cronb = TRUE)
```

# Arguments

data

A data frame that should be used to compute a FA, or a fa object.

rotation

Rotation of the factor loadings. May be "varimax" for orthogonal rotation or "promax" for oblique transformation (default). Requires the "GPArotation" package.

method

the factoring method to be used. "ml" will do a maximum likelihood factor analysis (default). "minres" will do a minimum residual (OLS), "wls" will do a weighted least squares (WLS) solution, "gls" does a generalized weighted least squares (GLS), "pa" will do the principal factor solution, "minchi" will minimize the sample size weighted chi square when treating pairwise correlations with different number of subjects per pair. "minrank" will do a minimum rank factor analysis.

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nmbr.fctr	Number of factors used for calculating the rotation. By default, this value is NULL and the amount of factors is calculated according to a parallel analysis.
fctr.load.tlrn	Specifies the minimum difference a variable needs to have between factor loadings (components) in order to indicate a clear loading on just one factor and not diffusing over all factors. For instance, a variable with 0.8, 0.82 and 0.84 factor loading on 3 possible factors can not be clearly assigned to just one factor and thus would be removed from the principal component analysis. By default, the minimum difference of loading values between the highest and 2nd highest factor should be 0.1
digits	Amount of decimals for estimates
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
type	Plot type resp. geom type. May be one of following: "circle" or "tile" circular or tiled geoms, or "bar" for a bar plot. You may use initial letter only for this argument.
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
show.values	Logical, whether values should be plotted or not.
show.cronb	Logical, if TRUE (default), the cronbach's alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension. Only applies when data is a data frame.

### Value

(Invisibly) returns a structure with

- the rotated factor loading matrix (rotate)
- the column indices of removed variables (for more details see next list item) (removed.colindex)
- an updated data frame containing all factors that have a clear loading on a specific scale in case data was a data frame (See argument fctr.load.tlrn for more details) (removed.df)
- the factor.index, i.e. the column index of each variable with the highest factor loading for each factor,
- the ggplot-object (plot),
- the data frame that was used for setting up the ggplot-object (df).

#### Note

This method for factor analysis relies on the functions fa and fa. parallel from the psych package.

## **Examples**

```
library(GPArotation)
data(efc)
# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")
# recveive last item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")

# use data frame as argument, let sjp.fa() compute FA
sjp.fa(efc[, start:end])
sjp.fa(efc[, start:end], type = "tile")</pre>
```

sjp.grpfrq

Plot grouped or stacked frequencies

### Description

Plot grouped or stacked frequencies of variables as bar/dot, box or violin plots, or line plot.

## Usage

```
sjp.grpfrq(var.cnt, var.grp, type = c("bar", "dot", "line", "boxplot",
    "violin"), bar.pos = c("dodge", "stack"), weight.by = NULL,
    intr.var = NULL, title = "", title.wtd.suffix = NULL,
    legend.title = NULL, axis.titles = NULL, axis.labels = NULL,
    legend.labels = NULL, intr.var.labels = NULL, wrap.title = 50,
    wrap.labels = 15, wrap.legend.title = 20, wrap.legend.labels = 20,
    geom.size = NULL, geom.spacing = 0.15, geom.colors = "Paired",
    show.values = TRUE, show.n = TRUE, show.prc = TRUE,
    show.axis.values = TRUE, show.ci = FALSE, show.grpcnt = FALSE,
    show.legend = TRUE, show.na = FALSE, show.summary = FALSE,
    drop.empty = TRUE, auto.group = NULL, ylim = NULL,
    grid.breaks = NULL, expand.grid = FALSE, inner.box.width = 0.15,
    inner.box.dotsize = 3, smooth.lines = FALSE, emph.dots = TRUE,
    summary.pos = "r", facet.grid = FALSE, coord.flip = FALSE,
    y.offset = NULL, vjust = "bottom", hjust = "center")
```

### **Arguments**

var.cnt Vector of counts, for which frequencies or means will be plotted or printed.

var.grp Factor with the cross-classifying variable, where var.cnt is grouped into the categories represented by var.grp.

type Specifies the plot type. May be abbreviated.

"bar" for simple bars (default)

"dot" for a dot plot

"histogram" for a histogram (does not apply to grouped frequencies)

"line" for a line-styled histogram with filled area

"density" for a density plot (does not apply to grouped frequencies)

"boxplot" for box plot
"violin" for violin plots

bar.pos Indicates whether bars should be positioned side-by-side (default), or stacked

(bar.pos = "stack"). May be abbreviated.

weight.by Vector of weights that will be applied to weight all cases. Must be a vector of

same length as the input vector. Default is NULL, so no weights are used.

intr.var An interaction variable which can be used for box plots. Divides each cat-

egory indicated by var.grp into the factors of intr.var, so that each category of var.grp is subgrouped into intr.var's categories. Only applies when

type = "boxplot" or type = "violin".

title character vector, used as plot title. Depending on plot type and function, will be

set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.

title.wtd.suffix

Suffix (as string) for the title, if weight.by is specified, e.g. title.wtd.suffix=" (weighted)".

Default is NULL, so title will not have a suffix when cases are weighted.

legend.title character vector, used as title for the plot legend.

axis.titles character vector of length one or two, defining the title(s) for the x-axis and

y-axis.

axis.labels character vector with labels used as axis labels. Optional argument, since in

most cases, axis labels are set automatically.

legend.labels character vector with labels for the guide/legend.

intr.var.labels

a character vector with labels for the x-axis breaks when having interaction variables included. These labels replace the axis.labels. Only applies, when using box or violin plots (i.e. type = "boxplot" or "violin") and intr.var

is not NULL.

wrap.title numeric, determines how many chars of the plot title are displayed in one line

and when a line break is inserted.

wrap.labels numeric, determines how many chars of the value, variable or axis labels are

displayed in one line and when a line break is inserted.

wrap.legend.title

numeric, determines how many chars of the legend's title are displayed in one

line and when a line break is inserted.

wrap.legend.labels

numeric, determines how many chars of the legend labels are displayed in one line and when a line break is inserted.

geom.size size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes. geom.spacing the spacing between geoms (i.e. bar spacing) user defined color for geoms. See 'Details' in sjp.grpfrq. geom.colors show.values Logical, whether values should be plotted or not. show.n logical, if TRUE, adds total number of cases for each group or category to the labels. show.prc logical, if TRUE (default), percentage values are plotted to each bar If FALSE, percentage values are removed. show.axis.values logical, whether category, count or percentage values for the axis should be printed or not. show.ci Logical, if TRUE), adds notches to the box plot, which are used to compare groups; if the notches of two boxes do not overlap, medians are considered to be significantly different. show.grpcnt logical, if TRUE, the count within each group is added to the category labels (e.g. "Cat 1 (n=87)"). Default value is FALSE. show.legend logical, if TRUE, and depending on plot type and function, a legend is added to the plot. logical, if TRUE, NA's (missing values) are added to the output. show.na logical, if TRUE (default), a summary with chi-squared statistics (see chisq.test), show.summary Cramer's V or Phi-value etc. is shown. If a cell contains expected values lower than five (or lower than 10 if df is 1), the Fisher's excact test (see fisher.test) is computed instead of chi-squared test. If the table's matrix is larger than 2x2, Fisher's excact test with Monte Carlo simulation is computed. Logical, if TRUE and the variable's values are labelled, values that have no obserdrop.empty vations are still printed in the table (with frequency 0). If FALSE, values / factor levels with no occurence in the data are omitted from the output. auto.group numeric value, indicating the minimum amount of unique values in the count variable, at which automatic grouping into smaller units is done (see group\_var). Default value for auto. group is NULL, i.e. auto-grouping is off. See group\_var for examples on grouping. numeric vector of length two, defining lower and upper axis limits of the y scale. ylim By default, this argument is set to NULL, i.e. the y-axis fits to the required range of the data. numeric; sets the distance between breaks for the axis, i.e. at every grid. breaks'th grid.breaks position a major grid is being printed. logical, if TRUE, the plot grid is expanded, i.e. there is a small margin between expand.grid axes and plotting region. Default is FALSE.

width of the inner box plot that is plotted inside of violin plots. Only applies if

type = "violin". Default value is 0.15

inner.box.width

 ${\color{blue} \mathsf{emph.dots}} \qquad \qquad \mathsf{logical, if \ TRUE, the \ groups \ of \ dots \ in \ a \ dot-plot \ are \ highlighted \ with \ a \ shaded}$ 

rectangle.

summary.pos position of the model summary which is printed when show.summary is TRUE.

Default is "r", i.e. it's printed to the upper right corner. Use "1" for upper left

corner.

facet.grid TRUE to arrange the lay out of of multiple plots in a grid of an integrated single plot. This argument calls facet\_wrap or facet\_grid to arrange plots. Use

plot\_grid to plot multiple plot-objects as an arranged grid with grid.arrange.

coord.flip logical, if TRUE, the x and y axis are swapped.

y. offset numeric, offset for text labels when their alignment is adjusted to the top/bottom

of the geom (see hjust and vjust).

vjust character vector, indicating the vertical position of value labels. Allowed are

same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which

align text towards and away from the center of the plot respectively.

hjust character vector, indicating the horizontal position of value labels. Allowed are

same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which

align text towards and away from the center of the plot respectively.

#### **Details**

geom.colors may be a character vector of color values in hex-format, valid color value names (see demo("colors") or a name of a color brewer palette. Following options are valid for the geom.colors argument:

- If not specified, a default color brewer palette will be used, which is suitable for the plot style (i.e. diverging for likert scales, qualitative for grouped bars etc.).
- If "gs", a greyscale will be used.
- If "bw", and plot-type is a line-plot, the plot is black/white and uses different line types to distinguish groups (see this package-vignette).
- If geom.colors is any valid color brewer palette name, the related palette will be used. Use display.brewer.all to view all available palette names.
- Else specify own color values or names as vector (e.g. geom.colors = c("#f00000", "#00ff00")).

#### Value

A ggplot-object.

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### **Examples**

```
data(efc)
sjp.grpfrq(efc$e17age, efc$e16sex, show.values = FALSE)
# boxplot
sjp.grpfrq(efc$e17age, efc$e42dep, type = "box")
# grouped bars
sjp.grpfrq(efc$e42dep, efc$e16sex, title = NULL)
# box plots with interaction variable
sjp.grpfrq(efc$e17age, efc$e42dep, intr.var = efc$e16sex, type = "box")
# Grouped bar plot
sjp.grpfrq(efc$neg_c_7, efc$e42dep, show.values = FALSE)
# same data as line plot
sjp.grpfrq(efc$neg_c_7, efc$e42dep, type = "line")
# show ony categories where we have data (i.e. drop zero-counts)
library(dplyr)
efc <- dplyr::filter(efc, e42dep %in% c(3,4))
sjp.grpfrq(efc$c161sex, efc$e42dep, drop.empty = TRUE)
# show all categories, even if not in data
sjp.grpfrq(efc$c161sex, efc$e42dep, drop.empty = FALSE)
```

sjp.pca

Plot PCA results

## **Description**

Performs a principle component analysis on a data frame or matrix (with varimax or oblimin rotation) and plots the factor solution as ellipses or tiles.

In case a data frame is used as argument, the cronbach's alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension.

### Usage

```
sjp.pca(data, rotation = c("varimax", "quartimax", "promax", "oblimin",
   "simplimax", "cluster", "none"), nmbr.fctr = NULL,
   fctr.load.tlrn = 0.1, plot.eigen = FALSE, digits = 2,
   title = NULL, axis.labels = NULL, type = c("bar", "circle",
   "tile"), geom.size = 0.6, geom.colors = "RdBu", wrap.title = 50,
   wrap.labels = 30, show.values = TRUE, show.cronb = TRUE)
```

sjp.pca

# Arguments

data	A data frame that should be used to compute a PCA, or a prcomp object.
rotation	Rotation of the factor loadings. May be one of "varimax", "quartimax", "promax", "oblimin", "si or "none".
nmbr.fctr	Number of factors used for calculating the rotation. By default, this value is NULL and the amount of factors is calculated according to the Kaiser-criteria.
fctr.load.tlrn	Specifies the minimum difference a variable needs to have between factor loadings (components) in order to indicate a clear loading on just one factor and not diffusing over all factors. For instance, a variable with 0.8, 0.82 and 0.84 factor loading on 3 possible factors can not be clearly assigned to just one factor and thus would be removed from the principal component analysis. By default, the minimum difference of loading values between the highest and 2nd highest factor should be 0.1
plot.eigen	If TRUE, a plot showing the Eigenvalues according to the Kaiser criteria is plotted to determine the number of factors.
digits	Amount of decimals for estimates
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
type	Plot type resp. geom type. May be one of following: "circle" or "tile" circular or tiled geoms, or "bar" for a bar plot. You may use initial letter only for this argument.
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
show.values	Logical, whether values should be plotted or not.
show.cronb	Logical, if TRUE (default), the cronbach's alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension. Only applies when data is a data frame.

# Value

(Invisibly) returns a structure with

- the rotated factor loading matrix (varim)
- the column indices of removed variables (for more details see next list item) (removed.colindex)

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 an updated data frame containing all factors that have a clear loading on a specific scale in case data was a data frame (See argument fctr.load.tlrn for more details) (removed.df)

- the factor.index, i.e. the column index of each variable with the highest factor loading for each factor,
- the ggplot-object (plot),
- the data frame that was used for setting up the ggplot-object (df).

### **Examples**

```
library(sjmisc)
data(efc)
# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")</pre>
# recveive last item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")</pre>
# manually compute PCA
pca <- prcomp(</pre>
 na.omit(efc[, start:end]),
 retx = TRUE,
 center = TRUE,
 scale. = TRUE
# plot results from PCA as circles, including Eigenvalue-diagnostic.
# note that this plot does not compute the Cronbach's Alpha
sjp.pca(pca, plot.eigen = TRUE, type = "circle", geom.size = 10)
# use data frame as argument, let sjp.pca() compute PCA
sjp.pca(efc[, start:end])
sjp.pca(efc[, start:end], type = "tile")
```

sjp.poly

Plot polynomials for (generalized) linear regression

## **Description**

This function plots a scatter plot of a term poly.term against a response variable x and adds - depending on the amount of numeric values in poly.degree - multiple polynomial curves. A loess-smoothed line can be added to see which of the polynomial curves fits best to the data.

### Usage

```
sjp.poly(x, poly.term, poly.degree, poly.scale = FALSE, fun = NULL,
axis.title = NULL, geom.colors = NULL, geom.size = 0.8,
show.loess = TRUE, show.loess.ci = TRUE, show.p = TRUE,
show.scatter = TRUE, point.alpha = 0.2, point.color = "#404040",
loess.color = "#808080")
```

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### **Arguments**

Х	A vector, representing the response variable of a linear (mixed) model; or a linear (mixed) model as returned by lm or lmer.
poly.term	If x is a vector, poly.term should also be a vector, representing the polynomial term (independent variabl) in the model; if x is a fitted model, poly.term should be the polynomial term's name as character string. See 'Examples'.
poly.degree	Numeric, or numeric vector, indicating the degree of the polynomial. If poly.degree is a numeric vector, multiple polynomial curves for each degree are plotted. See 'Examples'.
poly.scale	Logical, if TRUE, poly.term will be scaled before linear regression is computed. Default is FALSE. Scaling the polynomial term may have an impact on the resulting p-values.
fun	Linear function when modelling polynomial terms. Use fun = " $lm$ " for linear models, or fun = " $glm$ " for generalized linear models. When x is not a vector, but a fitted model object, the function is detected automatically. If x is a vector, fun defaults to " $lm$ ".
axis.title	Character vector of length one or two (depending on the plot function and type), used as title(s) for the x and y axis. If not specified, a default labelling is chosen. <b>Note:</b> Some plot types may not support this argument sufficiently. In such cases, use the returned ggplot-object and add axis titles manually with labs. Use axis.title = "" to remove axis titles.
geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.
show.loess	Logical, if TRUE, an additional loess-smoothed line is plotted.
show.loess.ci	Logical, if TRUE, a confidence region for the loess-smoothed line will be plotted.
show.p	Logical, if TRUE (default), p-values for polynomial terms are printed to the console.
show.scatter	Logical, if TRUE (default), adds a scatter plot of data points to the plot.
point.alpha	Alpha value of point-geoms in the scatter plots. Only applies, if $show.scatter = TRUE$ .
point.color	Color of of point-geoms in the scatter plots. Only applies, if show. scatter = TRUE.
loess.color	Color of the loess-smoothed line. Only applies, if show.loess = TRUE.

# **Details**

For each polynomial degree, a simple linear regression on x (resp. the extracted response, if x is a fitted model) is performed, where only the polynomial term poly.term is included as independent variable. Thus,  $lm(y \sim x + I(x^2) + ... + I(x^i))$  is repeatedly computed for all values in poly.degree, and the predicted values of the reponse are plotted against the raw values of poly.term. If x is a fitted model, other covariates are ignored when finding the best fitting polynomial.

This function evaluates raw polynomials, not orthogonal polynomials. Polynomials are computed

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using the poly function, with argument raw = TRUE.

To find out which polynomial degree fits best to the data, a loess-smoothed line (in dark grey) can be added (with show.loess = TRUE). The polynomial curves that comes closest to the loess-smoothed line should be the best fit to the data.

#### Value

A ggplot-object.

### **Examples**

```
library(sjmisc)
data(efc)
# linear fit. loess-smoothed line indicates a more
# or less cubic curve
sjp.poly(efc$c160age, efc$quol_5, 1)
# quadratic fit
sjp.poly(efc$c160age, efc$quol_5, 2)
# linear to cubic fit
sjp.poly(efc$c160age, efc$quol_5, 1:4, show.scatter = FALSE)
# fit sample model
fit <- lm(tot_sc_e ~ c12hour + e17age + e42dep, data = efc)</pre>
# inspect relationship between predictors and response
plot_model(fit, type = "slope")
# "e17age" does not seem to be linear correlated to response
# try to find appropiate polynomial. Grey line (loess smoothed)
# indicates best fit. Looks like x^4 has the best fit,
# however, only x^3 has significant p-values.
sjp.poly(fit, "e17age", 2:4, show.scatter = FALSE)
## Not run:
# fit new model
fit <- lm(tot_sc_e ~ c12hour + e42dep + e17age + I(e17age^2) + I(e17age^3),
          data = efc)
# plot marginal effects of polynomial term
plot_model(fit, type = "pred", terms = "e17age")
## End(Not run)
```

sjp.stackfrq

Plot stacked proportional bars

### **Description**

Plot items (variables) of a scale as stacked proportional bars. This function is useful when several items with identical scale/categoroies should be plotted to compare the distribution of answers.

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## Usage

```
sjp.stackfrq(items, title = NULL, legend.title = NULL,
  legend.labels = NULL, axis.titles = NULL, axis.labels = NULL,
  weight.by = NULL, sort.frq = NULL, wrap.title = 50,
  wrap.labels = 30, wrap.legend.title = 30, wrap.legend.labels = 28,
  geom.size = 0.5, geom.colors = "Blues", show.values = TRUE,
  show.n = TRUE, show.prc = TRUE, show.legend = TRUE,
  grid.breaks = 0.2, expand.grid = FALSE, digits = 1,
  vjust = "center", coord.flip = TRUE)
```

# Arguments

guments		
items	Data frame, with each column representing one item.	
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.	
legend.title	character vector, used as title for the plot legend.	
legend.labels	character vector with labels for the guide/legend.	
axis.titles	character vector of length one or two, defining the title(s) for the x-axis and y-axis.	
axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.	
weight.by	Vector of weights that will be applied to weight all cases. Must be a vector of same length as the input vector. Default is NULL, so no weights are used.	
sort.frq	Indicates whether the items should be ordered by by highest count of first or last category of items.	
	"first.asc" to order ascending by lowest count of first category,	
	"first.desc" to order descending by lowest count of first category,	
	"last.asc" to order ascending by lowest count of last category,	
	"last.desc" to order descending by lowest count of last category,	
	NULL (default) for no sorting.	
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.	
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.	
wrap.legend.title		
wrap.legend.la	numeric, determines how many chars of the legend's title are displayed in one line and when a line break is inserted.	
wi ap. regenu. Tai	numeric, determines how many chars of the legend labels are displayed in one	
	line and when a line break is inserted.	
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller	

values than dot sizes.

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geom.colors	user defined color for geoms. See 'Details' in sjp.grpfrq.
show.values	Logical, whether values should be plotted or not.
show.n	logical, if TRUE, adds total number of cases for each group or category to the labels.
show.prc	Logical, if TRUE (default), the percentage values at the x-axis are shown.
show.legend	logical, if TRUE, and depending on plot type and function, a legend is added to the plot.
grid.breaks	numeric; sets the distance between breaks for the axis, i.e. at every grid.breaks'th position a major grid is being printed.
expand.grid	logical, if TRUE, the plot grid is expanded, i.e. there is a small margin between axes and plotting region. Default is FALSE.
digits	Numeric, amount of digits after decimal point when rounding estimates or values.
vjust	character vector, indicating the vertical position of value labels. Allowed are same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which align text towards and away from the center of the plot respectively.
coord.flip	logical, if TRUE, the x and y axis are swapped.

### Value

A ggplot-object.

### Note

Thanks to Forrest Stevens for bug fixes.

# **Examples**

```
# Data from the EUROFAMCARE sample dataset
library(sjmisc)
data(efc)
# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")
# recveive first item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")
# auto-detection of labels
sjp.stackfrq(efc[, start:end])
```

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sjp.xtab	Plot contingency tables	

## **Description**

Plot proportional crosstables (contingency tables) of two variables as ggplot diagram.

### Usage

```
sjp.xtab(x, grp, type = c("bar", "line"), margin = c("col", "cell",
   "row"), bar.pos = c("dodge", "stack"), title = "",
   title.wtd.suffix = NULL, axis.titles = NULL, axis.labels = NULL,
   legend.title = NULL, legend.labels = NULL, weight.by = NULL,
   rev.order = FALSE, show.values = TRUE, show.n = TRUE,
   show.prc = TRUE, show.total = TRUE, show.legend = TRUE,
   show.summary = FALSE, summary.pos = "r", drop.empty = TRUE,
   string.total = "Total", wrap.title = 50, wrap.labels = 15,
   wrap.legend.title = 20, wrap.legend.labels = 20, geom.size = 0.7,
   geom.spacing = 0.1, geom.colors = "Paired", dot.size = 3,
   smooth.lines = FALSE, grid.breaks = 0.2, expand.grid = FALSE,
   ylim = NULL, vjust = "bottom", hjust = "center", y.offset = NULL,
   coord.flip = FALSE)
```

## **Arguments**

X	A vector of values (variable) describing the bars which make up the plot.	
grp	Grouping variable of same length as x, where x is grouped into the categories represented by grp.	
type	Plot type. may be either "bar" (default) for bar charts, or "line" for line diagram.	
margin	Indicates which data of the proportional table should be plotted. Use "row" for calculating row percentages, "col" for column percentages and "cell" for cell percentages. If margin = "col", an additional bar with the total sum of each column can be added to the plot (see show.total).	
bar.pos	Indicates whether bars should be positioned side-by-side (default), or stacked (bar.pos = "stack"). May be abbreviated.	
title	character vector, used as plot title. Depending on plot type and function, will be set automatically. If title = "", no title is printed. For effect-plots, may also be a character vector of length > 1, to define titles for each sub-plot or facet.	
title.wtd.suffix		
	Suffix (as string) for the title, if weight.by is specified, e.g. title.wtd.suffix=" (weighted)". Default is NULL, so title will not have a suffix when cases are weighted.	
axis.titles	character vector of length one or two, defining the title(s) for the x-axis and y-axis.	

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axis.labels	character vector with labels used as axis labels. Optional argument, since in most cases, axis labels are set automatically.
legend.title	character vector, used as title for the plot legend.
legend.labels	character vector with labels for the guide/legend.
weight.by	Vector of weights that will be applied to weight all cases. Must be a vector of same length as the input vector. Default is NULL, so no weights are used.
rev.order	Logical, if TRUE, order of categories (groups) is reversed.
show.values	Logical, whether values should be plotted or not.
show.n	logical, if TRUE, adds total number of cases for each group or category to the labels.
show.prc	logical, if TRUE (default), percentage values are plotted to each bar If FALSE, percentage values are removed.
show.total	When margin = "col", an additional bar with the sum within each category and it's percentages will be added to each category.
show.legend	logical, if TRUE, and depending on plot type and function, a legend is added to the plot.
show.summary	logical, if TRUE (default), a summary with chi-squared statistics (see chisq.test), Cramer's V or Phi-value etc. is shown. If a cell contains expected values lower than five (or lower than 10 if df is 1), the Fisher's excact test (see fisher.test) is computed instead of chi-squared test. If the table's matrix is larger than 2x2, Fisher's excact test with Monte Carlo simulation is computed.
summary.pos	position of the model summary which is printed when show. summary is TRUE. Default is "r", i.e. it's printed to the upper right corner. Use "1" for upper left corner.
drop.empty	Logical, if TRUE and the variable's values are labelled, values that have no observations are still printed in the table (with frequency 0). If FALSE, values / factor levels with no occurence in the data are omitted from the output.
string.total	String for the legend label when a total-column is added. Only applies if show. total = TRUE. Default is "Total".
wrap.title	numeric, determines how many chars of the plot title are displayed in one line and when a line break is inserted.
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
wrap.legend.ti	
numeric, determines how many chars of the legend's title are displayed in one line and when a line break is inserted. wrap.legend.labels	
. 0	numeric, determines how many chars of the legend labels are displayed in one line and when a line break is inserted.
geom.size	size resp. width of the geoms (bar width, line thickness or point size, depending on plot type and function). Note that bar and bin widths mostly need smaller values than dot sizes.
geom.spacing	the spacing between geoms (i.e. bar spacing)

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geom.colors user defined color for geoms. See 'Details' in sjp.grpfrq. dot.size Dot size, only applies, when argument type = "line". smooth.lines prints a smooth line curve. Only applies, when argument type = "line". grid.breaks numeric; sets the distance between breaks for the axis, i.e. at every grid. breaks'th position a major grid is being printed. logical, if TRUE, the plot grid is expanded, i.e. there is a small margin between expand.grid axes and plotting region. Default is FALSE. ylim numeric vector of length two, defining lower and upper axis limits of the y scale. By default, this argument is set to NULL, i.e. the y-axis fits to the required range of the data. vjust character vector, indicating the vertical position of value labels. Allowed are same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which align text towards and away from the center of the plot respectively. hjust character vector, indicating the horizontal position of value labels. Allowed are same values as for vjust aesthetics from ggplot2: "left", "center", "right", "bottom", "middle", "top" and new options like "inward" and "outward", which align text towards and away from the center of the plot respectively. numeric, offset for text labels when their alignment is adjusted to the top/bottom y.offset of the geom (see hjust and vjust). coord.flip logical, if TRUE, the x and y axis are swapped.

#### Value

A ggplot-object.

### **Examples**

```
# create 4-category-items
grp <- sample(1:4, 100, replace = TRUE)</pre>
# create 3-category-items
x <- sample(1:3, 100, replace = TRUE)
# plot "cross tablulation" of x and grp
sjp.xtab(x, grp)
# plot "cross tablulation" of x and y, including labels
sjp.xtab(x, grp, axis.labels = c("low", "mid", "high"),
         legend.labels = c("Grp 1", "Grp 2", "Grp 3", "Grp 4"))
# plot "cross tablulation" of x and grp
# as stacked proportional bars
sjp.xtab(x, grp, margin = "row", bar.pos = "stack",
         show.summary = TRUE, coord.flip = TRUE)
# example with vertical labels
library(sjmisc)
library(sjlabelled)
```

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```
data(efc)
set_theme(geom.label.angle = 90)
sjp.xtab(efc$e42dep, efc$e16sex, vjust = "center", hjust = "bottom")
# grouped bars with EUROFAMCARE sample dataset
# dataset was importet from an SPSS-file,
# see ?sjmisc::read_spss
data(efc)
efc.val <- get_labels(efc)</pre>
efc.var <- get_label(efc)</pre>
sjp.xtab(efc$e42dep, efc$e16sex, title = efc.var['e42dep'],
         axis.labels = efc.val[['e42dep']], legend.title = efc.var['e16sex'],
        legend.labels = efc.val[['e16sex']])
sjp.xtab(efc$e16sex, efc$e42dep, title = efc.var['e16sex'],
         axis.labels = efc.val[['e16sex']], legend.title = efc.var['e42dep'],
        legend.labels = efc.val[['e42dep']])
# auto-detection of labels works here
# so no need to specify labels. For
# title-auto-detection, use NULL
# -----
sjp.xtab(efc$e16sex, efc$e42dep, title = NULL)
sjp.xtab(efc$e16sex, efc$e42dep, margin = "row",
        bar.pos = "stack", coord.flip = TRUE)
```

sjplot

Wrapper to create plots and tables within a pipe-workflow

## **Description**

This function has a pipe-friendly argument-structure, with the first argument always being the data, followed by variables that should be plotted or printed as table. The function then transforms the input and calls the requested sjp.- resp. sjt.-function to create a plot or table.

Both sjplot() and sjtab() support grouped data frames.

#### **Usage**

```
sjplot(data, ..., fun = c("grpfrq", "xtab", "aov1", "likert",
    "stackfrq"))
sjtab(data, ..., fun = c("xtab", "stackfrq"))
```

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## Arguments

data	A data frame. May also be a grouped data frame (see 'Note' and 'Examples').
	Names of variables that should be plotted, and also further arguments passed down to the <b>sjPlot</b> -functions. See 'Examples'.
fun	Plotting function. Refers to the function name of <b>sjPlot</b> -functions. See 'Details' and 'Examples'.

#### **Details**

Following fun-values are currently supported:

```
"aov1" calls sjp. aov1. The first two variables in data are used (and required) to create the plot.
```

"grpfrq" calls sjp.grpfrq. The first two variables in data are used (and required) to create the plot.

"likert" calls plot\_likert. data must be a data frame with items to plot.

"stackfrq" calls sjp.stackfrq or sjt.stackfrq. data must be a data frame with items to create the plot or table.

"xtab" calls sjp.xtab or sjt.xtab. The first two variables in data are used (and required) to create the plot or table.

#### Value

See related sjp. and sjt.-functions.

#### Note

The ...-argument is used, first, to specify the variables from data that should be plotted, and, second, to name further arguments that are used in the subsequent plotting functions. Refer to the online-help of supported plotting-functions to see valid arguments.

data may also be a grouped data frame (see group\_by) with up to two grouping variables. Plots are created for each subgroup then.

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```
group_by(e16sex, c172code) %>%
  select(e42dep, n4pstu, e16sex, c172code) %>%
  sjtab(fun = "xtab", use.viewer = FALSE) # open all tables in browser
## End(Not run)
```

sjPlot-themes

Modify plot appearance

## **Description**

Set default plot themes, use pre-defined color scales or modify plot or table appearance.

## Usage

```
theme_sjplot(base_size = 12, base_family = "")
theme_sjplot2(base_size = 12, base_family = "")
theme_blank(base_size = 12, base_family = "")
theme_538(base_size = 12, base_family = "")
font_size(title, axis_title.x, axis_title.y, labels.x, labels.y, offset.x, offset.y, base.theme)
label_angle(angle.x, angle.y, base.theme)
legend_style(inside, pos, justify, base.theme)
scale_color_sjplot(palette = "metro ui", discrete = TRUE, reverse = FALSE, ...)
scale_fill_sjplot(palette = "metro ui", discrete = TRUE, reverse = FALSE, ...)
sjplot_pal(palette = "metro ui", n = NULL)
show_sjplot_pals()
css_theme(css.theme = "regression")
```

# Arguments

```
base_sizebase_familybase font family.titleFont size for plot titles.
```

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axis_title.x	Font size for x-axis titles.
axis_title.y	Font size for y-axis titles.
labels.x	Font size for x-axis labels.
labels.y	Font size for y-axis labels.
offset.x	Offset for x-axis titles.
offset.y	Offset for y-axis titles.
base.theme	Optional ggplot-theme-object, which is needed in case multiple functions should be combined, e.g. theme_sjplot() + label_angle(). In such cases, use label_angle(base.theme = theme_sjplot()).
angle.x	Angle for x-axis labels.
angle.y	Angle for y-axis labels.
inside	Logical, use TRUE to put legend inside the plotting area. See also pos.
pos	Position of the legend, if a legend is drawn.
	<b>Legend outside plot</b> Use "bottom", "top", "left" or "right" to position the legend above, below, on the left or right side of the diagram.
	<b>Legend inside plot</b> If inside = TRUE, legend can be placed inside plot. Use "top left", "top right", "bottom left" and "bottom right" to position legend in any of these corners, or a two-element numeric vector with values from 0-1. See also inside.
justify	Justification of legend, relative to its position ("center" or two-element numeric vector with values from 0-1.
palette	Character name of color palette.
discrete	Logical, if TRUE, a discrete colour palette is returned. Else, a gradient palette is returned, where colours of the requested palette are interpolated using colorRampPalette.
reverse	Logical, if TRUE, order of returned colours is reversed.
	Further arguments passed down to ggplot's scale()-functions.
n	Numeric, number of colors to be returned. By default, the complete colour palette is returned.
css.theme	Name of the CSS pre-set theme-style. Can be used for table-functions.

## **Details**

When using the colors argument in function calls (e.g. plot\_model()) or when calling one of the predefined scale-functions (e.g. scale\_color\_sjplot()), there are pre-defined colour palettes in this package. Use show\_sjplot\_pals() to show all available colour palettes.

```
# prepare data
library(sjmisc)
data(efc)
efc <- to_factor(efc, c161sex, e42dep, c172code)
m <- lm(neg_c_7 ~ pos_v_4 + c12hour + e42dep + c172code, data = efc)</pre>
```

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```
# create plot-object
p <- plot_model(m)

# change theme
p + theme_sjplot()

# change font-size
p + font_size(axis_title.x = 30)

# apply color theme
p + scale_color_sjplot()

# show all available colour palettes
show_sjplot_pals()

# get colour values from specific palette
sjplot_pal(pal = "breakfast club")</pre>
```

sjt.corr

Summary of correlations as HTML table

## Description

Shows the results of a computed correlation as HTML table. Requires either a data.frame or a matrix with correlation coefficients as returned by the cor-function.

## Usage

```
sjt.corr(data, na.deletion = c("listwise", "pairwise"),
  corr.method = c("pearson", "spearman", "kendall"), title = NULL,
  var.labels = NULL, wrap.labels = 40, show.p = TRUE,
  p.numeric = FALSE, fade.ns = TRUE, val.rm = NULL, digits = 3,
  triangle = "both", string.diag = NULL, CSS = NULL,
  encoding = NULL, file = NULL, use.viewer = TRUE,
  remove.spaces = TRUE)
```

## **Arguments**

data	Matrix with correlation coefficients as returned by the cor-function, or a data. frame of variables where correlations between columns should be computed.
na.deletion	Indicates how missing values are treated. May be either "listwise" (default) or "pairwise". May be abbreviated.
corr.method	Indicates the correlation computation method. May be one of "spearman" (default), "pearson" or "kendall". May be abbreviated.
title	String, will be used as table caption.
var.labels	Character vector with variable names, which will be used to label variables in the output.

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wrap.labels Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.

show.p Logical, if TRUE, p-values are also printed.

p.numeric Logical, if TRUE, the p-values are printed as numbers. If FALSE (default), aster-

isks are used.

fade.ns Logical, if TRUE (default), non-significant correlation-values appear faded (by

using a lighter grey text color). See 'Note'.

val.rm Specify a number between 0 and 1 to suppress the output of correlation values

that are smaller than val.rm. The absolute correlation values are used, so a correlation value of -.5 would be greater than val.rm = .4 and thus not be omitted. By default, this argument is NULL, hence all values are shown in the table. If a correlation value is below the specified value of val.rm, it is still printed to the HTML table, but made "invisible" with white foreground color. You can use the CSS argument ("css.valueremove") to change color and appearance of those correlation value that are smaller than the limit specified by

val.rm.

digits Amount of decimals for estimates

triangle Indicates whether only the upper right (use "upper"), lower left (use "lower")

or both (use "both") triangles of the correlation table is filled with values. De-

fault is "both". You can specify the inital letter only.

string.diag A vector with string values of the same length as ncol(data) (number of cor-

related items) that can be used to display content in the diagonal cells where row and column item are identical (i.e. the "self-correlation"). By defauilt, this

argument is NULL and the diagnal cells are empty.

CSS A list with user-defined style-sheet-definitions, according to the official CSS

syntax. See 'Details' or this package-vignette.

encoding String, indicating the charset encoding used for variable and value labels. De-

fault is NULL, so encoding will be auto-detected depending on your platform (e.g., "UTF-8" for Unix and "Windows-1252" for Windows OS). Change encoding if specific chars are not properly displayed (e.g. German umlauts).

file Destination file, if the output should be saved as file. If NULL (default), the output

will be saved as temporary file and openend either in the IDE's viewer pane or

the default web browser.

use.viewer Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE

or no viewer available, the HTML table is opened in a web browser.

remove. spaces Logical, if TRUE, leading spaces are removed from all lines in the final string that

contains the html-data. Use this, if you want to remove parantheses for html-tags. The html-source may look less pretty, but it may help when exporting

html-tables to office tools.

## Value

Invisibly returns

• the web page style sheet (page.style),

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- the web page content (page.content),
- the complete html-output (page.complete) and
- the html-table with inline-css for use with knitr (knitr)

for further use.

#### Note

If data is a matrix with correlation coefficients as returned by the cor-function, p-values can't be computed. Thus, show.p, p.numeric and fade.ns only have an effect if data is a data.frame.

```
## Not run:
# plot correlation matrix using circles
sjt.corr(mydf)
# Data from the EUROFAMCARE sample dataset
library(sjmisc)
data(efc)
# retrieve variable and value labels
varlabs <- get_label(efc)</pre>
# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c83cop2")</pre>
# recveive last item of COPE-index scale
end <- which(colnames(efc) == "c88cop7")</pre>
# create data frame with COPE-index scale
mydf <- data.frame(efc[, c(start:end)])</pre>
colnames(mydf) <- varlabs[c(start:end)]</pre>
# we have high correlations here, because all items
# belong to one factor. See example from "sjp.pca".
sjt.corr(mydf, p.numeric = TRUE)
# auto-detection of labels, only lower triangle
sjt.corr(efc[, c(start:end)], triangle = "lower")
# auto-detection of labels, only lower triangle, all correlation
# values smaller than 0.3 are not shown in the table
sjt.corr(efc[, c(start:end)], triangle = "lower", val.rm = 0.3)
# auto-detection of labels, only lower triangle, all correlation
# values smaller than 0.3 are printed in blue
sjt.corr(efc[, c(start:end)], triangle = "lower",val.rm = 0.3,
         CSS = list(css.valueremove = 'color:blue;'))
## End(Not run)
```

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Summary of factor analysis as HTML table

#### **Description**

Performes a factor analysis on a data frame or matrix and displays the factors as HTML table, or saves them as file.

In case a data frame is used as parameter, the Cronbach's Alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension.

## Usage

```
sjt.fa(data, rotation = c("promax", "varimax"), method = c("ml",
 "minres", "wls", "gls", "pa", "minchi", "minrank"), nmbr.fctr = NULL,
 fctr.load.tlrn = 0.1, title = "Factor Analysis", var.labels = NULL,
 wrap.labels = 40, show.cronb = TRUE, show.comm = FALSE,
 alternate.rows = FALSE, digits = 2, CSS = NULL, encoding = NULL,
 file = NULL, use.viewer = TRUE, remove.spaces = TRUE)
```

## **Arguments**

data A data frame that should be used to compute a FA, or a fa object.

rotation Rotation of the factor loadings. May be "varimax" for orthogonal rotation or

"promax" for oblique transformation (default). Requires the "GPArotation"

package.

method the factoring method to be used. "ml" will do a maximum likelihood factor

> analysis (default). "minres" will do a minimum residual (OLS), "wls" will do a weighted least squares (WLS) solution, "gls" does a generalized weighted least squares (GLS), "pa" will do the principal factor solution, "minchi" will minimize the sample size weighted chi square when treating pairwise correlations with different number of subjects per pair. "minrank" will do a minimum

rank factor analysis.

nmbr.fctr Number of factors used for calculating the rotation. By default, this value is

NULL and the amount of factors is calculated according to a parallel analysis.

fctr.load.tlrn Specifies the minimum difference a variable needs to have between factor load-

ings (components) in order to indicate a clear loading on just one factor and not diffusing over all factors. For instance, a variable with 0.8, 0.82 and 0.84 factor loading on 3 possible factors can not be clearly assigned to just one factor and thus would be removed from the principal component analysis. By default, the minimum difference of loading values between the highest and 2nd highest

factor should be 0.1

title character vector, used as plot title. Depending on plot type and function, will be

set automatically. If title = "", no title is printed. For effect-plots, may also

be a character vector of length > 1, to define titles for each sub-plot or facet.

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var.labels	Character vector with variable names, which will be used to label variables in the output.	
wrap.labels	numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.	
show.cronb	Logical, if TRUE (default), the cronbach's alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension. Only applies when data is a data frame.	
show.comm	Logical, if TRUE, show the communality column in the table.	
alternate.rows	Logical, if TRUE, rows are printed in alternatig colors (white and light grey by default).	
digits	Amount of decimals for estimates	
CSS	A list with user-defined style-sheet-definitions, according to the official CSS syntax. See 'Details' or this package-vignette.	
encoding	Character vector, indicating the charset encoding used for variable and value labels. Default is "UTF-8". For Windows Systems, encoding = "Windows-1252" might be necessary for proper display of special characters.	
file	Destination file, if the output should be saved as file. If NULL (default), the output will be saved as temporary file and openend either in the IDE's viewer pane or the default web browser.	
use.viewer	Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE or no viewer available, the HTML table is opened in a web browser.	
remove.spaces	Logical, if TRUE, leading spaces are removed from all lines in the final string that contains the html-data. Use this, if you want to remove parantheses for html-tags. The html-source may look less pretty, but it may help when exporting html-tables to office tools.	

## Value

# Invisibly returns

- the web page style sheet (page.style),
- the web page content (page.content),
- the complete html-output (page.complete),
- the html-table with inline-css for use with knitr (knitr),
- the factor.index, i.e. the column index of each variable with the highest factor loading for each factor and
- the removed.items, i.e. which variables have been removed because they were outside of the fctr.load.tlrn's range.

for further use.

## Note

This method for factor analysis relies on the functions fa and fa.parallel from the psych package.

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## **Examples**

```
## Not run:
# Data from the EUROFAMCARE sample dataset
library(sjmisc)
library(GPArotation)
data(efc)

# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")
# recveive last item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")
# auto-detection of labels
sjt.fa(efc[, start:end])
## End(Not run)</pre>
```

sjt.itemanalysis

Summary of item analysis of an item scale as HTML table

# Description

This function performs an item analysis with certain statistics that are useful for scale or index development. The resulting tables are shown in the viewer pane resp. webbrowser or can be saved as file. Following statistics are computed for each item of a data frame:

- percentage of missing values
- · mean value
- · standard deviation
- skew
- · item difficulty
- · item discrimination
- · Cronbach's Alpha if item was removed from scale
- mean (or average) inter-item-correlation

Optional, following statistics can be computed as well:

- · kurstosis
- · Shapiro-Wilk Normality Test

If factor.groups is not NULL, the data frame df will be splitted into groups, assuming that factor.groups indicate those columns of the data frame that belong to a certain factor (see return value of function sjt.pca as example for retrieving factor groups for a scale and see examples for more details).

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## Usage

```
sjt.itemanalysis(df, factor.groups = NULL,
 factor.groups.titles = "auto", scale = FALSE,
 min.valid.rowmean = 2, alternate.rows = TRUE, sort.column = NULL,
 show.shapiro = FALSE, show.kurtosis = FALSE,
 show.corr.matrix = TRUE, CSS = NULL, encoding = NULL,
 file = NULL, use.viewer = TRUE, remove.spaces = TRUE)
```

## **Arguments**

df

A data frame with items.

factor.groups

If not NULL, df will be splitted into sub-groups, where the item analysis is carried out for each of these groups. Must be a vector of same length as ncol(df), where each item in this vector represents the group number of the related columns of df. See 'Examples'.

factor.groups.titles

Titles for each factor group that will be used as table caption for each componenttable. Must be a character vector of same length as length(unique(factor.groups)). Default is "auto", which means that each table has a standard caption Component x. Use NULL to suppress table captions.

scale

Logical, if TRUE, the data frame's vectors will be scaled when calculating the Cronbach's Alpha value (see item\_reliability). Recommended, when the variables have different measures / scales.

min.valid.rowmean

Minimum amount of valid values to compute row means for index scores. Default is 2, i.e. the return values index.scores and df.index.scores are computed for those items that have at least min.valid.rowmean per case (observation, or technically, row). See mean\_n for details.

alternate.rows Logical, if TRUE, rows are printed in alternatig colors (white and light grey by default).

sort.column

Numeric vector, indicating the index of the column that should sorted. by default, the column is sorted in ascending order. Use negative index for descending order, for instance, sort.column = -3 would sort the third column in descending order. Note that the first column with rownames is not counted.

show.shapiro

Logical, if TRUE, a Shapiro-Wilk normality test is computed for each item. See shapiro. test for details.

show.kurtosis

Logical, if TRUE, the kurtosis for each item will also be shown (see kurtosi and describe in the psych-package for more details.

show.corr.matrix

Logical, if TRUE (default), a correlation matrix of each component's index score is shown. Only applies if factor, groups is not NULL and df has more than one group. First, for each case (df's row), the sum of all variables (df's columns) is scaled (using the scale-function) and represents a "total score" for each component (a component is represented by each group of factor.groups). After that, each case (df's row) has a scales sum score for each component. Finally, a correlation of these "scale sum scores" is computed.

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CSS A list with user-defined style-sheet-definitions, according to the official CSS

syntax. See 'Details' or this package-vignette.

encoding Character vector, indicating the charset encoding used for variable and value la-

bels. Default is "UTF-8". For Windows Systems, encoding = "Windows-1252"

might be necessary for proper display of special characters.

file Destination file, if the output should be saved as file. If NULL (default), the output

will be saved as temporary file and openend either in the IDE's viewer pane or

the default web browser.

use.viewer Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE

or no viewer available, the HTML table is opened in a web browser.

remove. spaces Logical, if TRUE, leading spaces are removed from all lines in the final string that

contains the html-data. Use this, if you want to remove parantheses for html-tags. The html-source may look less pretty, but it may help when exporting

html-tables to office tools.

#### Value

## Invisibly returns

• df.list: List of data frames with the item analysis for each sub.group (or complete, if factor.groups was NULL)

- index.scores: A data frame with of standardized scale / index scores for each case (mean value of all scale items for each case) for each sub-group.
- ideal.item.diff: List of vectors that indicate the ideal item difficulty for each item in each sub-group. Item difficulty only differs when items have different levels.
- cronbach.values: List of Cronbach's Alpha values for the overall item scale for each subgroup.
- knitr.list: List of html-tables with inline-css for use with knitr for each table (sub-group)
- knitr: html-table of all complete output with inline-css for use with knitr
- complete.page: Complete html-output.

If factor.groups = NULL, each list contains only one elment, since just one table is printed for the complete scale indicated by df. If factor.groups is a vector of group-index-values, the lists contain elements for each sub-group.

## Note

- The *Shapiro-Wilk Normality Test* (see column W(p)) tests if an item has a distribution that is significantly different from normal.
- *Item difficulty* should range between 0.2 and 0.8. Ideal value is p+(1-p)/2 (which mostly is between 0.5 and 0.8).
- For *item discrimination*, acceptable values are 0.20 or higher; the closer to 1.00 the better. See item\_reliability for more details.
- In case the total *Cronbach's Alpha* value is below the acceptable cut-off of 0.7 (mostly if an index has few items), the *mean inter-item-correlation* is an alternative measure to indicate acceptability. Satisfactory range lies between 0.2 and 0.4. See also item\_intercor.

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## References

 Jorion N, Self B, James K, Schroeder L, DiBello L, Pellegrino J (2013) Classical Test Theory Analysis of the Dynamics Concept Inventory. (web)

- Briggs SR, Cheek JM (1986) The role of factor analysis in the development and evaluation of personality scales. Journal of Personality, 54(1), 106-148. doi: 10.1111/j.14676494.1986.tb00391.x
- McLean S et al. (2013) Stigmatizing attitudes and beliefs about bulimia nervosa: Gender, age, education and income variability in a community sample. International Journal of Eating Disorders. doi: 10.1002/eat.22227
- Trochim WMK (2008) Types of Reliability. (web)

```
# Data from the EUROFAMCARE sample dataset
library(sjmisc)
library(sjlabelled)
data(efc)
# retrieve variable and value labels
varlabs <- get_label(efc)</pre>
# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")</pre>
# recveive last item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")</pre>
# create data frame with COPE-index scale
mydf <- data.frame(efc[, start:end])</pre>
colnames(mydf) <- varlabs[start:end]</pre>
## Not run:
sjt.itemanalysis(mydf)
# auto-detection of labels
sjt.itemanalysis(efc[, start:end])
# Compute PCA on Cope-Index, and perform a
# item analysis for each extracted factor.
factor.groups <- sjt.pca(mydf)$factor.index</pre>
sjt.itemanalysis(mydf, factor.groups)
## End(Not run)
```

sjt.pca

## **Description**

Performes a principle component analysis on a data frame or matrix (with varimax or oblimin rotation) and displays the factor solution as HTML table, or saves them as file.

In case a data frame is used as parameter, the Cronbach's Alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension.

## Usage

```
sjt.pca(data, rotation = c("varimax", "quartimax", "promax", "oblimin",
   "simplimax", "cluster", "none"), nmbr.fctr = NULL,
   fctr.load.tlrn = 0.1, title = "Principal Component Analysis",
   var.labels = NULL, wrap.labels = 40, show.cronb = TRUE,
   show.msa = FALSE, show.var = FALSE, alternate.rows = FALSE,
   digits = 2, string.pov = "Proportion of Variance",
   string.cpov = "Cumulative Proportion", CSS = NULL, encoding = NULL,
   file = NULL, use.viewer = TRUE, remove.spaces = TRUE)
```

## **Arguments**

data	A data frame that should be used to compute a PCA, or a prcomp object.
rotation	Rotation of the factor loadings. May be one of "varimax", "quartimax", "promax", "oblimin", "si or "none".
nmbr.fctr	Number of factors used for calculating the rotation. By default, this value is NULL and the amount of factors is calculated according to the Kaiser-criteria.
fctr.load.tlrn	Specifies the minimum difference a variable needs to have between factor loadings (components) in order to indicate a clear loading on just one factor and not diffusing over all factors. For instance, a variable with 0.8, 0.82 and 0.84 factor loading on 3 possible factors can not be clearly assigned to just one factor and thus would be removed from the principal component analysis. By default, the minimum difference of loading values between the highest and 2nd highest factor should be 0.1
title	String, will be used as table caption.
var.labels	Character vector with variable names, which will be used to label variables in the output.
wrap.labels	Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
show.cronb	Logical, if TRUE (default), the cronbach's alpha value for each factor scale will be calculated, i.e. all variables with the highest loading for a factor are taken for the reliability test. The result is an alpha value for each factor dimension. Only applies when data is a data frame.
show.msa	Logical, if TRUE, shows an additional column with the measure of sampling adequacy according dor each component.
show.var	Logical, if TRUE, the proportions of variances for each component as well as cumulative variance are shown in the table footer.

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alternate.rows	Logical, if TRUE, rows are printed in alternatig colors (white and light grey by default).
digits	Amount of decimals for estimates
string.pov	String for the table row that contains the proportions of variances. By default, "Proportion of Variance" will be used.
string.cpov	String for the table row that contains the cumulative variances. By default, "Cumulative Proportion" will be used.
CSS	A list with user-defined style-sheet-definitions, according to the official CSS syntax. See 'Details' or this package-vignette.
encoding	Character vector, indicating the charset encoding used for variable and value labels. Default is "UTF-8". For Windows Systems, encoding = "Windows-1252" might be necessary for proper display of special characters.
file	Destination file, if the output should be saved as file. If NULL (default), the output will be saved as temporary file and openend either in the IDE's viewer pane or the default web browser.
use.viewer	Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE or no viewer available, the HTML table is opened in a web browser.
remove.spaces	Logical, if TRUE, leading spaces are removed from all lines in the final string that contains the html-data. Use this, if you want to remove parantheses for html-tags. The html-source may look less pretty, but it may help when exporting html-tables to office tools.

#### Value

# Invisibly returns

- the web page style sheet (page.style),
- the web page content (page.content),
- the complete html-output (page.complete),
- the html-table with inline-css for use with knitr (knitr),
- the factor.index, i.e. the column index of each variable with the highest factor loading for each factor and
- the removed.items, i.e. which variables have been removed because they were outside of the fctr.load.tlrn's range.

for further use.

```
## Not run:
# Data from the EUROFAMCARE sample dataset
library(sjmisc)
data(efc)

# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")</pre>
```

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```
# recveive last item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")
# auto-detection of labels
sjt.pca(efc[, start:end])
## End(Not run)</pre>
```

sjt.stackfrq

Summary of stacked frequencies as HTML table

## **Description**

Shows the results of stacked frequencies (such as likert scales) as HTML table. This function is useful when several items with identical scale/categories should be printed as table to compare their distributions (e.g. when plotting scales like SF, Barthel-Index, Quality-of-Life-scales etc.).

## Usage

```
sjt.stackfrq(items, weight.by = NULL, title = NULL,
  var.labels = NULL, value.labels = NULL, wrap.labels = 20,
  sort.frq = NULL, alternate.rows = FALSE, digits = 2,
  string.total = "N", string.na = "NA", show.n = FALSE,
  show.total = FALSE, show.na = FALSE, show.skew = FALSE,
  show.kurtosis = FALSE, digits.stats = 2, file = NULL,
  encoding = NULL, CSS = NULL, use.viewer = TRUE,
  remove.spaces = TRUE)
```

#### **Arguments**

items	Data frame, with each column representing one item.	
weight.by	Vector of weights that will be applied to weight all cases. Must be a vector of same length as the input vector. Default is NULL, so no weights are used.	
title	Character vector with table caption(s) resp. footnote(s). For tab_df(), must be a character of length 1; for tab_dfs(), a character vector of same length as x (i.e. one title or footnote per data frame).	
var.labels	Character vector with variable names, which will be used to label variables in the output.	
value.label	S Character vector (or list of character vectors) with value labels of the supplied variables, which will be used to label variable values in the output.	
wrap.labels	Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.	
sort.frq	logical, indicates whether the items should be ordered by by highest count of first or last category of items.	

- Use "first.asc" to order ascending by lowest count of first category,
- "first.desc" to order descending by lowest count of first category,

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• "last.asc" to order ascending by lowest count of last category,

• "last.desc" to order descending by lowest count of last category,

• or NULL (default) for no sorting.

alternate.rows Logical, if TRUE, rows are printed in alternatig colors (white and light grey by

default).

digits Amount of decimals for estimates

string.total label for the total N column.

string.na label for the missing column/row.

show.n logical, if TRUE, adds total number of cases for each group or category to the

labels.

show.total logical, if TRUE, an additional column with each item's total N is printed.

show. na logical, if TRUE, NA's (missing values) are added to the output.

show. skew logical, if TRUE, an additional column with each item's skewness is printed. The

skewness is retrieved from the describe-function of the psych-package.

show.kurtosis Logical, if TRUE, the kurtosis for each item will also be shown (see kurtosi and

describe in the psych-package for more details.

digits.stats amount of digits for rounding the skewness and kurtosis valuess. Default is 2,

i.e. skewness and kurtosis values have 2 digits after decimal point.

file Destination file, if the output should be saved as file. If NULL (default), the output

will be saved as temporary file and openend either in the IDE's viewer pane or

the default web browser.

encoding Character vector, indicating the charset encoding used for variable and value la-

bels. Default is "UTF-8". For Windows Systems, encoding = "Windows-1252"

might be necessary for proper display of special characters.

CSS A list with user-defined style-sheet-definitions, according to the official CSS

syntax. See 'Details' or this package-vignette.

use.viewer Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE

or no viewer available, the HTML table is opened in a web browser.

remove. spaces Logical, if TRUE, leading spaces are removed from all lines in the final string that

contains the html-data. Use this, if you want to remove parantheses for html-tags. The html-source may look less pretty, but it may help when exporting

html-tables to office tools.

#### Value

#### Invisibly returns

- the web page style sheet (page.style),
- the web page content (page.content),
- the complete html-output (page.complete) and
- the html-table with inline-css for use with knitr (knitr)

for further use.

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```
# -----
# random sample
# -----
# prepare data for 4-category likert scale, 5 items
likert_4 <- data.frame(</pre>
 as.factor(sample(1:4, 500, replace = TRUE, prob = c(0.2, 0.3, 0.1, 0.4))),
 as.factor(sample(1:4, 500, replace = TRUE, prob = c(0.5, 0.25, 0.15, 0.1)),
 as.factor(sample(1:4, 500, replace = TRUE, prob = c(0.25, 0.1, 0.4, 0.25))),
 as.factor(sample(1:4, 500, replace = TRUE, prob = c(0.1, 0.4, 0.4, 0.1))),
 as.factor(sample(1:4, 500, replace = TRUE, prob = c(0.35, 0.25, 0.15, 0.25)))
)
# create labels
levels_4 <- c("Independent", "Slightly dependent",</pre>
             "Dependent", "Severely dependent")
# create item labels
items <- c("Q1", "Q2", "Q3", "Q4", "Q5")
# plot stacked frequencies of 5 (ordered) item-scales
## Not run:
sjt.stackfrq(likert_4, value.labels = levels_4, var.labels = items)
# Data from the EUROFAMCARE sample dataset
# Auto-detection of labels
# -----
data(efc)
# recveive first item of COPE-index scale
start <- which(colnames(efc) == "c82cop1")</pre>
# recveive first item of COPE-index scale
end <- which(colnames(efc) == "c90cop9")</pre>
sjt.stackfrq(efc[, c(start:end)], alternate.rows = TRUE)
sjt.stackfrq(efc[, c(start:end)], alternate.rows = TRUE,
            show.n = TRUE, show.na = TRUE)
# User defined style sheet
sjt.stackfrq(efc[, c(start:end)], alternate.rows = TRUE,
            show.total = TRUE, show.skew = TRUE, show.kurtosis = TRUE,
            CSS = list(css.ncol = "border-left:1px dotted black;",
                       css.summary = "font-style:italic;"))
## End(Not run)
```

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## **Description**

Shows contingency tables as HTML file in browser or viewer pane, or saves them as file.

#### Usage

```
sjt.xtab(var.row, var.col, weight.by = NULL, title = NULL,
  var.labels = NULL, value.labels = NULL, wrap.labels = 20,
  show.obs = TRUE, show.cell.prc = FALSE, show.row.prc = FALSE,
  show.col.prc = FALSE, show.exp = FALSE, show.legend = FALSE,
  show.na = FALSE, show.summary = TRUE, drop.empty = TRUE,
  statistics = c("auto", "cramer", "phi", "spearman", "kendall",
  "pearson", "fisher"), string.total = "Total", digits = 1,
  tdcol.n = "black", tdcol.expected = "#339999",
  tdcol.cell = "#993333", tdcol.row = "#333399",
  tdcol.col = "#339933", emph.total = FALSE, emph.color = "#f8f8f8",
  prc.sign = " %", hundret = "100.0", CSS = NULL,
  encoding = NULL, file = NULL, use.viewer = TRUE,
  remove.spaces = TRUE, ...)
```

## **Arguments**

var.row	Variable that should be displayed in the table rows.	
var.col	Cariable that should be displayed in the table columns.	
weight.by	Vector of weights that will be applied to weight all cases. Must be a vector of same length as the input vector. Default is NULL, so no weights are used.	
title	String, will be used as table caption.	
var.labels	Character vector with variable names, which will be used to label variables in the output.	
value.labels	Character vector (or list of character vectors) with value labels of the supplied variables, which will be used to label variable values in the output.	
wrap.labels	Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.	
show.obs	Logical, if TRUE, observed values are shown	
show.cell.prc	Logical, if TRUE, cell percentage values are shown	
show.row.prc	Logical, if TRUE, row percentage values are shown	
show.col.prc	Logical, if TRUE, column percentage values are shown	
show.exp	Logical, if TRUE, expected values are also shown	
show.legend	logical, if TRUE, and depending on plot type and function, a legend is added to the plot.	
show.na	logical, if TRUE, NA's (missing values) are added to the output.	
show.summary	Logical, if TRUE, a summary row with chi-squared statistics, degrees of freedom and Cramer's V or Phi coefficient and p-value for the chi-squared statistics.	
drop.empty	Logical, if TRUE and the variable's values are labelled, values that have no observations are still printed in the table (with frequency $\emptyset$ ). If FALSE, values / factor levels with no occurence in the data are omitted from the output.	

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statistics Name of measure of association that should be computed. May be one of "auto", "cramer", "phi", "spearman", "kendall", "pearson" or "fisher". See xtab\_statistics. string.total Character label for the total column / row header Amount of decimals for estimates digits tdcol.n Color for highlighting count (observed) values in table cells. Default is black. tdcol.expected Color for highlighting expected values in table cells. Default is cyan. tdcol.cell Color for highlighting cell percentage values in table cells. Default is red. tdcol.row Color for highlighting row percentage values in table cells. Default is blue. tdcol.col Color for highlighting column percentage values in table cells. Default is green. emph.total Logical, if TRUE, the total column and row will be emphasized with a different background color. See emph.color. Logical, if emph. total = TRUE, this color value will be used for painting the emph.color background of the total column and row. Default is a light grey. prc.sign The percentage sign that is printed in the table cells, in HTML-format. Default is " %", hence the percentage sign has a non-breaking-space after the percentage value. hundret Default value that indicates the 100-percent column-sums (since rounding values may lead to non-exact results). Default is "100.0". CSS A list with user-defined style-sheet-definitions, according to the official CSS syntax. See 'Details' or this package-vignette. encoding String, indicating the charset encoding used for variable and value labels. Default is NULL, so encoding will be auto-detected depending on your platform (e.g., "UTF-8" for Unix and "Windows-1252" for Windows OS). Change encoding if specific chars are not properly displayed (e.g. German umlauts). file Destination file, if the output should be saved as file. If NULL (default), the output will be saved as temporary file and openend either in the IDE's viewer pane or the default web browser. Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE use.viewer or no viewer available, the HTML table is opened in a web browser. Logical, if TRUE, leading spaces are removed from all lines in the final string that remove.spaces contains the html-data. Use this, if you want to remove parantheses for htmltags. The html-source may look less pretty, but it may help when exporting html-tables to office tools. Other arguments, currently passed down to the test statistics functions chisq.test() or fisher.test().

#### Value

#### Invisibly returns

- the web page style sheet (page.style),
- the web page content (page.content),

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- the complete html-output (page.complete) and
- the html-table with inline-css for use with knitr (knitr)

for further use.

```
# prepare sample data set
data(efc)
# print simple cross table with labels
## Not run:
sjt.xtab(efc$e16sex, efc$e42dep)
# print cross table with manually set
# labels and expected values
sjt.xtab(
 efc$e16sex,
 efc$e42dep,
 var.labels = c("Elder's gender", "Elder's dependency"),
 show.exp = TRUE
)
# print minimal cross table with labels, total col/row highlighted
sjt.xtab(efc$e16sex, efc$e42dep, show.cell.prc = FALSE, emph.total = TRUE)
# User defined style sheet
sjt.xtab(efc$e16sex, efc$e42dep,
        CSS = list(css.table = "border: 2px solid;",
                   css.tdata = "border: 1px solid;",
                    css.horline = "border-bottom: double blue;"))
## End(Not run)
# ordinal data, use Kendall's tau
sjt.xtab(efc$e42dep, efc$quol_5, statistics = "kendall")
# calculate Spearman's rho, with continuity correction
sjt.xtab(
 efc$e42dep,
 efc$quol_5,
 statistics = "spearman",
 exact = FALSE,
 continuity = TRUE
)
```

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#### **Description**

These functions print data frames as HTML-table, showing the results in RStudio's viewer pane or in a web browser.

#### Usage

```
tab_df(x, title = NULL, footnote = NULL, col.header = NULL,
    show.type = FALSE, show.rownames = TRUE, show.footnote = FALSE,
    alternate.rows = FALSE, sort.column = NULL, encoding = "UTF-8",
    CSS = NULL, file = NULL, use.viewer = TRUE, ...)

tab_dfs(x, titles = NULL, footnotes = NULL, col.header = NULL,
    show.type = FALSE, show.rownames = TRUE, show.footnote = FALSE,
    alternate.rows = FALSE, sort.column = NULL, encoding = "UTF-8",
    CSS = NULL, file = NULL, use.viewer = TRUE, ...)
```

## **Arguments**

sort.column

file

x For tab\_df(), a data frame; and for tab\_dfs(), a list of data frames. title, titles, footnote, footnotes

Character vector with table caption(s) resp. footnote(s). For tab\_df(), must be a character of length 1; for tab\_dfs(), a character vector of same length as x (i.e. one title or footnote per data frame).

col.header Character vector with elements used as column header for the table. If NULL, column names from x are used as column header.

show. type Logical, if TRUE, adds information about the variable type to the variable column.

show.rownames Logical, if TRUE, adds a column with the data frame's rowname to the table

show.footnote Logical, if TRUE,adds a summary footnote below the table. For tab\_df(), specify the string in footnote, for tab\_dfs() provide a character vector in footnotes.

alternate.rows Logical, if TRUE, rows are printed in alternatig colors (white and light grey by default).

Numeric vector, indicating the index of the column that should sorted. by default, the column is sorted in ascending order. Use negative index for descending order, for instance, sort.column = -3 would sort the third column in descending order. Note that the first column with rownames is not counted.

encoding Character vector, indicating the charset encoding used for variable and value labels. Default is "UTF-8". For Windows Systems, encoding = "Windows-1252" might be necessary for proper display of special characters.

CSS A list with user-defined style-sheet-definitions, according to the official CSS syntax. See 'Details' or this package-vignette.

Destination file, if the output should be saved as file. If NULL (default), the output will be saved as temporary file and openend either in the IDE's viewer pane or the default web browser.

tab\_df

use.viewer Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE or no viewer available, the HTML table is opened in a web browser.

Currently not used.

#### **Details**

#### How do I use CSS-argument?

With the CSS-argument, the visual appearance of the tables can be modified. To get an overview of all style-sheet-classnames that are used in this function, see return value page.style for details. Arguments for this list have following syntax:

- 1. the class-name as argument name and
- 2. each style-definition must end with a semicolon

You can add style information to the default styles by using a + (plus-sign) as initial character for the argument attributes. Examples:

- table = 'border:2px solid red;' for a solid 2-pixel table border in red.
- summary = 'font-weight: bold; ' for a bold fontweight in the summary row.
- lasttablerow = 'border-bottom: 1px dotted blue;' for a blue dotted border of the last table row.
- colnames = '+color: green' to add green color formatting to column names.
- arc = 'color:blue;' for a blue text color each 2nd row.
- caption = '+color:red;' to add red font-color to the default table caption style.

See further examples in this package-vignette.

#### Value

A list with following items:

- the web page style sheet (page.style),
- the HTML content of the data frame (page.content),
- the complete HTML page, including header, style sheet and body (page.complete)
- the HTML table with inline-css for use with knitr (knitr)
- the file path, if the HTML page should be saved to disk (file)

## Note

The HTML tables can either be saved as file and manually opened (use argument file) or they can be saved as temporary files and will be displayed in the RStudio Viewer pane (if working with RStudio) or opened with the default web browser. Displaying resp. opening a temporary file is the default behaviour.

## **Examples**

```
## Not run:
data(iris)
data(mtcars)
tab_df(iris[1:5, ])
tab_dfs(list(iris[1:5, ], mtcars[1:5, 1:5]))

# sort 2nd column ascending
tab_df(iris[1:5, ], sort.column = 2)

# sort 2nd column descending
tab_df(iris[1:5, ], sort.column = -2)
## End(Not run)
```

tab\_model

Print regression models as HTML table

## **Description**

tab\_model() creates HTML tables from regression models.

## Usage

```
tab_model(..., transform, show.intercept = TRUE, show.est = TRUE,
 show.ci = 0.95, show.ci50 = TRUE, show.se = NULL,
 show.std = NULL, show.p = TRUE, show.stat = FALSE,
 show.df = FALSE, show.zeroinf = TRUE, show.r2 = TRUE,
 show.icc = TRUE, show.re.var = TRUE, show.ngroups = TRUE,
 show.fstat = FALSE, show.aic = FALSE, show.aicc = FALSE,
 show.dev = FALSE, show.loglik = FALSE, show.obs = TRUE,
 terms = NULL, rm.terms = NULL, group.terms = TRUE,
 order.terms = NULL, title = NULL, pred.labels = NULL,
 dv.labels = NULL, wrap.labels = 25, vcov.fun = NULL,
 vcov.type = c("HC3", "const", "HC", "HC0", "HC1", "HC2", "HC4", "HC4m",
 "HC5"), vcov.args = NULL, string.pred = "Predictors",
 string.est = "Estimate", string.std = "std. Beta",
 string.ci = "CI", string.se = "std. Error", string.p = "p",
 string.df = "df", string.stat = "Statistic",
 string.resp = "Response", strings = NULL,
 ci.hyphen = " – ", minus.sign = "-",
 collapse.ci = FALSE, collapse.se = FALSE, linebreak = TRUE,
 col.order = c("est", "se", "std.est", "std.se", "ci", "std.ci",
 "ci.inner", "ci.outer", "stat", "p", "df", "response.level"),
 digits = 2, digits.p = 3, emph.p = TRUE, p.val = c("wald", "kr"),
 p.style = c("numeric", "asterisk", "both"), p.threshold = c(0.05,
 0.01, 0.001), case = "parsed", auto.label = TRUE,
 prefix.labels = c("none", "varname", "label"), bpe = "median",
 CSS = css_theme("regression"), file = NULL, use.viewer = TRUE)
```

#### **Arguments**

One or more regression models, including glm's or mixed models. May also be a list with fitted models. See 'Examples'. transform A character vector, naming a function that will be applied on estimates and confidence intervals. By default, transform will automatically use "exp" as transformation for applicable classes of model (e.g. logistic or poisson regression). Estimates of linear models remain untransformed. Use NULL if you want the raw, non-transformed estimates. show.intercept Logical, if TRUE, the intercepts are printed. show.est Logical, if TRUE, the estimates are printed. Either logical, and if TRUE, the confidence intervals is printed to the table; if show.ci FALSE, confidence intervals are omitted. Or numeric, between 0 and 1, indicating the range of the confidence intervals. show.ci50 Logical, if TRUE, for Bayesian models, a second credible interval is added to the table output. show.se Logical, if TRUE, the standard errors are also printed. If robust standard errors are required, use arguments vcov. fun, vcov. type and vcov. args (see robust for details). show.std Indicates whether standardized beta-coefficients should also printed, and if yes, which type of standardization is done. See 'Details'. Logical, if TRUE, p-values are also printed. show.p show.stat Logical, if TRUE, the coefficients' test statistic is also printed. show.df Logical, if TRUE and p.val = "kr", the p-values for linear mixed models are based on df with Kenward-Rogers approximation. These df-values are printed. See p\_value for details. show.zeroinf Logical, if TRUE and model has a zero-inflated model part, this is also printed to the table. Logical, if TRUE, the r-squared value is also printed. Depending on the model, show.r2 these might be pseudo-r-squared values, or Bayesian r-squared etc. See r2 for show.icc Logical, if TRUE, prints the intraclass correlation coefficient for mixed models. See icc for details. Logical, if TRUE, prints the random effect variances for mixed models. See show.re.var get\_variance for details. show.ngroups Logical, if TRUE, shows number of random effects groups for mixed models. Logical, if TRUE, the F-statistics for each model is printed in the table summary. show.fstat This option is not supported by all model types. show.aic Logical, if TRUE, the AIC value for each model is printed in the table summary. Logical, if TRUE, the second-order AIC value for each model is printed in the show.aicc table summary. show.dev Logical, if TRUE, shows the deviance of the model. show.loglik Logical, if TRUE, shows the log-Likelihood of the model.

show.obs Logical, if TRUE, the number of observations per model is printed in the table summary.

Character vector with names of those terms (variables) that should be printed in terms

the table. All other terms are removed from the output. If NULL, all terms are printed. Note that the term names must match the names of the model's coefficients. For factors, this means that the variable name is suffixed with the related factor level, and each category counts as one term. E.g. rm. terms = "t\_name [2,3]" would remove the terms "t\_name2" and "t\_name3" (assuming that the variable t\_name is categorical and has at least the factor levels 2 and 3). Another example for the *iris*-dataset: terms = "Species" would not work, instead use terms = "Species [versicolor, virginica]".

Character vector with names that indicate which terms should be removed from rm.terms

the output Counterpart to terms. rm. terms = "t\_name" would remove the term t\_name. Default is NULL, i.e. all terms are used. For factors, levels that should be removed from the plot need to be explicitely indicated in square brackets, and match the model's coefficient names, e.g. rm. terms = "t\_name [2,3]" would remove the terms "t\_name2" and "t\_name3" (assuming that the variable

t\_name was categorical and has at least the factor levels 2 and 3).

Logical, if TRUE (default), automatically groups table rows with factor levels of group.terms same factor, i.e. predictors of type factor will be grouped, if the factor has more than two levels. Grouping means that a separate headline row is inserted

to the table just before the predictor values.

order.terms Numeric vector, indicating in which order the coefficients should be plotted. See

examples in this package-vignette.

String, will be used as table caption. title

pred.labels Character vector with labels of predictor variables. If not NULL, pred.labels

> will be used in the first table column with the predictors' names. By default, if auto.label = TRUE and get\_term\_labels is called to retrieve the labels of the coefficients, which will be used as predictor labels. If pred.labels = "" or auto.label = FALSE, the raw variable names as used in the model formula are used as predictor labels. If pred. labels is a named vector, predictor labels (by default, the names of the model's coefficients) will be matched with the names of pred.labels. This ensures that labels always match the related predictor in the table, no matter in which way the predictors are sorted. See 'Examples'.

dv.labels Character vector with labels of dependent variables of all fitted models. See 'Examples'.

> Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.

vcov.fun Character vector, indicating the name of the vcov\*()-function from the sandwichpackage, e.g. vcov.fun = "vcovCL", if robust standard errors are required.

vcov.type Character vector, specifying the estimation type for the robust covariance matrix estimation (see vcovHC for details).

> List of named vectors, used as additional arguments that are passed down to vcov.fun.

Character vector, used as headline for the predictor column. Default is "Predictors". string.pred

wrap.labels

vcov.args

string.est	Character vector, used for the column heading of coefficients. Default is based on the response scale, e.g. for logistic regression models, "Odds Ratios" will be chosen, while for Poisson models it is "Incidence Rate Ratios" etc. Default if not specified is "Estimate".
string.std	Character vector, used for the column heading of standardized beta coefficients. Default is "std. Beta".
string.ci	Character vector, used for the column heading of confidence interval values. Default is "CI".
string.se	Character vector, used for the column heading of standard error values. Default is "std. Error".
string.p	Character vector, used for the column heading of p values. Default is "p".
string.df	Character vector, used for the column heading of degrees of freedom. Default is "df".
string.stat	Character vector, used for the test statistic. Default is "Statistic".
string.resp	Character vector, used for the column heading of of the response level for multi- nominal or categorical models. Default is "Response".
strings	Named character vector, as alternative to arguments like string.ci or string.p etc. The name (lhs) must be one of the string-indicator from the forementioned arguments, while the value (rhs) is the string that is used as column heading. E.g., strings = c(ci = "Conf.Int.", se = "std. Err") would be equivalent to setting string.ci = "Conf.Int.", string.se = "std. Err".
ci.hyphen	Character vector, indicating the hyphen for confidence interval range. May be an HTML entity. See 'Examples'.
minus.sign	string, indicating the minus sign for negative numbers. May be an HTML entity. See 'Examples'.
collapse.ci	Logical, if FALSE, the CI values are shown in a separate table column.
collapse.se	Logical, if FALSE, the SE values are shown in a separate table column.
linebreak	Logical, if TRUE and collapse.ci = FALSE or collapse.se = FALSE, inserts a line break between estimate and CI resp. SE values. If FALSE, values are printed in the same line as estimate values.
col.order	Character vector, indicating which columns should be printed and in which order. Column names that are excluded from col.order are not shown in the table output. However, column names that are included, are only shown in the table when the related argument (like show.est for "estimate") is set to TRUE or another valid value. Table columns are printed in the order as they appear in col.order.
digits	Amount of decimals for estimates
digits.p	Amount of decimals for p-values
emph.p	Logical, if TRUE, significant p-values are shown bold faced.
p.val	Character, for mixed models, indicates how p-values are computed. Use p.val = "wald" for a faster, but less precise computation. For p.val = "kr", computation of p-values is based on conditional F-tests with Kenward-Roger approximation for the degrees of freedom, using the <b>pbkrtest</b> -package. In this case, use show.df = TRUE to show the approximated degrees of freedom for each coefficient.

p.style	Character, indicating if p-values should be printed as numeric value ("numeric"), as asterisks ("asterisk") or both ("both"). May be abbreviated.
p.threshold	Numeric vector of length 3, indicating the treshold for annotating p-values with asterisks. Only applies if p. style = "asterisk".
case	Desired target case. Labels will automatically converted into the specified character case. See to_any_case for more details on this argument. By default, if case is not specified, it will be set to "parsed", unless prefix.labels is not "none". If prefix.labels is either "label" (or "l") or "varname" (or "v") and case is not specified, it will be set to NULL - this is a more convenient default when prefixing labels.
auto.label	Logical, if TRUE (the default), plot-labels are based on value and variable labels, if the data is labelled. See <a href="mailto:get_label">get_label</a> and <a href="mailto:get_labels">get_label</a> for details. If FALSE, original variable names and value labels (factor levels) are used.
prefix.labels	Indicates whether the value labels of categorical variables should be prefixed, e.g. with the variable name or variable label. See argument prefix in get_term_labels for details.
bpe	For <b>Stan</b> -models (fitted with the <b>rstanarm</b> - or <b>brms</b> -package), the Bayesian point estimate is, by default, the median of the posterior distribution. Use bpe to define other functions to calculate the Bayesian point estimate. bpe needs to be a character naming the specific function, which is passed to the fun-argument in typical_value. So, bpe = "mean" would calculate the mean value of the posterior distribution.
CSS	A list with user-defined style-sheet-definitions, according to the official CSS syntax. See 'Details' or this package-vignette.
file	Destination file, if the output should be saved as file. If NULL (default), the output will be saved as temporary file and openend either in the IDE's viewer pane or the default web browser.
use.viewer	Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE or no viewer available, the HTML table is opened in a web browser.

## **Details**

## **Standardized Estimates**

Concerning the show.std argument, show.std = "std" will print normal standardized estimates. For show.std = "std2", however, standardization of estimates follows Gelman's (2008) suggestion, rescaling the estimates by dividing them by two standard deviations instead of just one. Resulting coefficients are then directly comparable for untransformed binary predictors. For backward compatibility reasons, show.std also may be a logical value; if TRUE, normal standardized estimates are printed (same effect as show.std = "std"). Use show.std = NULL (default) or show.std = FALSE, if standardized estimats should not be printed.

# How do I use CSS-argument?

With the CSS-argument, the visual appearance of the tables can be modified. To get an overview of all style-sheet-classnames that are used in this function, see return value page.style for details. Arguments for this list have following syntax:

view\_df

- 1. the class-names with "css."-prefix as argument name and
- 2. each style-definition must end with a semicolon

You can add style information to the default styles by using a + (plus-sign) as initial character for the argument attributes. Examples:

- css.table = 'border:2px solid red;' for a solid 2-pixel table border in red.
- css.summary = 'font-weight:bold;' for a bold fontweight in the summary row.
- css.lasttablerow = 'border-bottom: 1px dotted blue;' for a blue dotted border of the last table row.
- css.colnames = '+color: green' to add green color formatting to column names.
- css.arc = 'color:blue;' for a blue text color each 2nd row.
- css.caption = '+color:red;' to add red font-color to the default table caption style.

#### Value

Invisibly returns

- the web page style sheet (page.style),
- the web page content (page.content),
- the complete html-output (page.complete) and
- the html-table with inline-css for use with knitr (knitr)

for further use.

## Note

The HTML tables can either be saved as file and manually opened (use argument file) or they can be saved as temporary files and will be displayed in the RStudio Viewer pane (if working with RStudio) or opened with the default web browser. Displaying resp. opening a temporary file is the default behaviour (i.e. file = NULL).

Examples are shown in these three vignettes: Summary of Regression Models as HTML Table, Summary of Mixed Models as HTML Table and Summary of Bayesian Models as HTML Table.

view\_df

View structure of labelled data frames

## **Description**

Save (or show) content of an imported SPSS, SAS or Stata data file, or any similar labelled data. frame, as HTML table. This quick overview shows variable ID number, name, label, type and associated value labels. The result can be considered as "codeplan" of the data frame.

view\_df

## Usage

```
view_df(x, weight.by = NULL, alternate.rows = TRUE, show.id = TRUE,
    show.type = FALSE, show.values = TRUE, show.string.values = FALSE,
    show.labels = TRUE, show.frq = FALSE, show.prc = FALSE,
    show.wtd.frq = FALSE, show.wtd.prc = FALSE, show.na = FALSE,
    max.len = 15, sort.by.name = FALSE, wrap.labels = 50,
    verbose = TRUE, CSS = NULL, encoding = NULL, file = NULL,
    use.viewer = TRUE, remove.spaces = TRUE)
```

## **Arguments**

Х	A (labelled) data frame, imported by read_spss, read_sas or read_stata function, or any similar labelled data frame (see set_label and set_labels).
weight.by	Name of variable in x that indicated the vector of weights that will be applied to weight all observations. Default is NULL, so no weights are used.
alternate.rows	Logical, if TRUE, rows are printed in alternatig colors (white and light grey by default). $\label{eq:logical}$
show.id	Logical, if TRUE (default), the variable ID is shown in the first column.
show.type	Logical, if TRUE, adds information about the variable type to the variable column. $ \\$
<pre>show.values show.string.val</pre>	Logical, if TRUE (default), the variable values are shown as additional column.
· ·	Logical, if TRUE, elements of character vectors are also shown. By default, these are omitted due to possibly overlengthy tables.
show.labels	Logical, if TRUE (default), the value labels are shown as additional column.
show.frq	Logical, if TRUE, an additional column with frequencies for each variable is shown. $ \\$
show.prc	Logical, if TRUE, an additional column with percentage of frequencies for each variable is shown.
show.wtd.frq	Logical, if TRUE, an additional column with weighted frequencies for each variable is shown. Weights strem from weight.by.
show.wtd.prc	Logical, if TRUE, an additional column with weighted percentage of frequencies for each variable is shown. Weights strem from weight.by.
show.na	logical, if TRUE, NA's (missing values) are added to the output.
max.len	Numeric, indicates how many values and value labels per variable are shown. Useful for variables with many different values, where the output can be truncated.
sort.by.name	Logical, if TRUE, rows are sorted according to the variable names. By default, rows (variables) are ordered according to their order in the data frame.
wrap.labels	Numeric, determines how many chars of the value, variable or axis labels are displayed in one line and when a line break is inserted.
verbose	Logical, if TRUE, a progress bar is displayed while creating the output.
CSS	A list with user-defined style-sheet-definitions, according to the official CSS syntax. See 'Details' or this package-vignette.

view\_df

encoding Character vector, indicating the charset encoding used for variable and value la-

bels. Default is "UTF-8". For Windows Systems, encoding = "Windows-1252"

might be necessary for proper display of special characters.

file Destination file, if the output should be saved as file. If NULL (default), the output

will be saved as temporary file and openend either in the IDE's viewer pane or

the default web browser.

use.viewer Logical, if TRUE, the HTML table is shown in the IDE's viewer pane. If FALSE

or no viewer available, the HTML table is opened in a web browser.

remove. spaces Logical, if TRUE, leading spaces are removed from all lines in the final string that

contains the html-data. Use this, if you want to remove parantheses for html-tags. The html-source may look less pretty, but it may help when exporting

html-tables to office tools.

## Value

Invisibly returns

- the web page style sheet (page.style),
- the web page content (page.content),
- the complete html-output (page.complete) and
- the html-table with inline-css for use with knitr (knitr)

for further use.

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