# Package 'smplot2' 

May 22, 2024
Type PackageTitle Creating and Annotating a Composite Plot in 'ggplot2'
Version 0.2.2
Maintainer Seung Hyun (Sam) Min [seung.min@mail.mcgill.ca](mailto:seung.min@mail.mcgill.ca)
Description Provides functions for creating and annotating a composite plot in 'ggplot2'. Offers back-ground themes and shortcut plotting functions that produce figures that are appropri-ate for the format of scientific journals. Some methods are de-scribed in Min and Zhou (2021) [doi:10.3389/fgene.2021.802894](doi:10.3389/fgene.2021.802894).
Encoding UTF-8
RoxygenNote 7.3.1
License GPL-2
Imports ggplot2, ggpubr, cowplot, tibble, dplyr, gghalves, utils, pwr,sdamr, stats, Hmisc, zoo, patchwork
URL https://smin95.github.io/dataviz/
BugReports https://github.com/smin95/smplot2/issues
NeedsCompilation no
Author Seung Hyun (Sam) Min [aut, cre]
Repository CRAN
Date/Publication 2024-05-22 07:10:03 UTC
$R$ topics documented:
sm_add_arrow ..... 2
sm_add_legend ..... 4
sm_add_line ..... 5
sm_add_point ..... 6
sm_add_polygon ..... 7
sm_add_rect ..... 8
sm_add_text ..... 9
sm_auc ..... 10
sm_auc_all ..... 11
sm_bar ..... 12
sm_bland_altman ..... 13
sm_boxplot ..... 14
sm_ci ..... 16
sm_classic ..... 16
sm_color ..... 17
sm_common_axis ..... 18
sm_common_legend ..... 19
sm_common_title ..... 20
sm_common_xlabel ..... 21
sm_common_ylabel ..... 21
sm_corr_avgErr ..... 22
sm_effsize ..... 23
sm_forest ..... 24
sm_forest_annot ..... 26
sm_hgrid ..... 27
sm_hist ..... 28
sm_hvgrid ..... 29
sm_hvgrid_minor ..... 30
sm_minimal ..... 31
sm_palette ..... 31
sm_panel_label ..... 32
sm_plot_clean ..... 33
sm_pointplot ..... 34
sm_power ..... 35
sm_put_together ..... 36
sm_raincloud ..... 38
sm_slope ..... 40
sm_slope_all ..... 42
sm_slope_theme ..... 43
sm_statBlandAlt ..... 44
sm_statCorr ..... 45
sm_stdErr ..... 46
sm_vgrid ..... 47
sm_violin ..... 48
Index ..... 50

sm_add_arrow

Adding an arrow annotation in the combined plot

## Description

Adding an arrow annotation in the combined plot

## Usage

```
sm_add_arrow(
        x,
        y,
        xend,
        yend,
        color = "black",
        linewidth = 0.5,
        type = "open",
        arrowlength = 1,
)
```


## Arguments

$x \quad$ Starting location of the arrow along the x -axis of the combined figure. The middle origin is at 0.5 .
$y \quad$ Starting location of the arrow along the $y$-axis of the combined figure. The middle origin is at 0.5 .
xend $\quad$ Final location of the arrow along the $x$-axis of the combined figure. The middle origin is at 0.5 .
yend $\quad$ Final location of the arrow along the $y$-axis of the combined figure. The middle origin is at 0.5 .
color Color of the line. Default is set to black.
linewidth Thickness of the arrow Default is set to 0.5 .
type Type of the arrow. Default is set to "open". Other choices include "both' and "closed".
arrowlength A unit specifying the length of the arrow head (from tip to base).
... Other parameters of the arrow line that will be transferred to the function annotate()

## Value

Prints an arrow to the combined plot.

## Examples

```
library(ggplot2)
library(smplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '#0f993d', color = 'white',
    size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white', size = 3) +
    sm_hvgrid() -> p2
```

```
    combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
    combined_fig + sm_add_arrow(x = 0.4, y = 0.4, xend = 0.6, yend = 0.6)
```

sm_add_legend Adding a common legend on a combined figure

## Description

Adding a common legend on a combined figure

## Usage

```
    sm_add_legend(
        combined_plot,
        x,
        y,
        sampleplot,
        legend,
        direction = "vertical",
        border = TRUE,
        legend_spacing = 0.5,
        border_color = "black",
        font_size = 12
    )
```


## Arguments

combined_plot Combined figure, an output from sm_put_together().
$x \quad$ Location of the legend along the $x$-axis of the combined figure. The middle origin is at 0.5 .
y Location of the legend along the $y$-axis of the combined figure. The middle origin is at 0.5 .
sampleplot A variable containing one sample ggplot2 from which the legend can be derived.
legend $\quad$ Pre-specified layer of legend created with sm_common_legend().
direction Direction of the legend: 'horizontal' or 'vertical'.
border If set TRUE, border around the legend will be created. If set FALSE, the border will be removed.
legend_spacing Spacing within the legend.
border_color Color of the legend border
font_size Text size of the legend

## Value

It prints a legend on a a combined plot. It can be used to create a common legend for subplots.

## Examples

```
library(ggplot2)
library(smplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg,
fill = as.factor(cyl))) +
    geom_point(shape = 21, color = 'white',
        size = 3) +
    sm_classic(legends=FALSE) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg,
            fill = as.factor(cyl))) +
    geom_point(shape = 21, color = 'white',
        size = 3) +
    sm_hvgrid(legends=FALSE) -> p2
combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
sm_add_legend(combined_fig, x = 0.1, y = 0.1, sampleplot = p1)
```

sm_add_line

Adding a line annotation in the combined plot

## Description

Adding a line annotation in the combined plot

## Usage

sm_add_line (x, y, xend, yend, color = "black", linewidth = 0.5, ...)

## Arguments

x
$y \quad$ Starting location of the line along the $y$-axis of the combined figure. The middle origin is at 0.5 . Values from 0 to 1 .
xend $\quad$ Final location of the line along the x -axis of the combined figure. The middle origin is at 0.5 . Values from 0 to 1 .
yend $\quad$ Final location of the line along the $y$-axis of the combined figure. The middle origin is at 0.5 . Values from 0 to 1 .
color Color of the line. Default is set to black.
linewidth Thickness of the line. Default is set to 0.5 .
$\ldots \quad$ Other parameters of the line that will be transferred to the function annotate() from ggplot2.

## Value

Prints a line to the combined plot.

## Examples

```
library(ggplot2)
library(smplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '#0f993d', color = 'white',
    size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white', size = 3) +
    sm_hvgrid() -> p2
combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
combined_fig + sm_add_line(x = 0.4, y = 0.4, xend = 0.6, yend = 0.6)
```

sm_add_point
Add a point annotation onto the combined plot

## Description

Add a point annotation onto the combined plot

## Usage

sm_add_point (x, y, size $=10$, shape = 16, color = "black", ...)

## Arguments

$x \quad$ Location of the point annotation along the $x$-axis of the combined figure. Default is the middle origin (0.5). Values from 0 to 1 .
$y \quad$ Location of the point annotation along the $y$-axis of the combined figure. Default is the middle origin (0.5). Values from 0 to 1.
size Size of the point
shape $\quad$ Shape of the point. Default is set to circle without border (16).
color Color of the point. Default is set to black.
... Other parameters of point that get passed to geom_point().

## Value

Prints a point in the combined plot.

## Examples

```
library(ggplot2)
library(smplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '#0f993d', color = 'white',
    size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white', size = 3) +
    sm_hvgrid() -> p2
combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
combined_fig + sm_add_point(color='red', size = 10, x = .5, y= .5)
```

```
sm_add_polygon Adding a polygon patch on the combined figure
```


## Description

Areas captured by the coorinates of $x$ and $y$ will be filled. The lengths of $x$ and $y$ have no limitations as long as their lengths are matched.

## Usage

sm_add_polygon(x, y, fill = "gray80", color = "black", linewidth = 0.5, ...)

## Arguments

$x \quad$ A vector containing $x$ coordinates from each point of the polygon. The first element of $x$ is matched to the firt element of $y$. Values from 0 to 1 , where 0.5 is the origin.
$y \quad$ A vector containing y coordinates from each point of the polygon. The first element of $x$ is matched to the firt element of $y$. Values from 0 to 1 , where 0.5 is the origin.
fill Fill color for the rectangle to be used
color Border line of the rectangle
linewidth Linewidth of border of the rectangle
... Other parameters to be used in annotate() from ggplot2.

## Value

Prints a patch of rectangle onto a combined plot.

## Examples

```
library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white',
        size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white', size = 3) +
    sm_hvgrid() -> p2
combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
combined_fig + sm_add_polygon(x = c(.2,.8,.8), y = c(.2,.8,.2))
```


## Description

Areas captured by xmin, ymin, xmax and ymax will become a patch once the function gets called.

## Usage

```
sm_add_rect (
    xmin,
    ymin,
    xmax,
    ymax,
    fill = "gray80",
    color = "black",
    linewidth = 0.5,
    )
```


## Arguments

| xmin | Starting x value of the rectangle patch. Values from 0 to 1. |
| :--- | :--- |
| ymin | Starting $y$ value of the rectangle patch. Values from 0 to 1. |
| xmax | Ending $x$ value of the rectangle patch. Values from 0 to 1. |
| ymax | Ending $x$ value of the rectangle patch. Values from 0 to 1. |
| fill | Fill color for the rectangle to be used |
| color | Border line of the rectangle |
| linewidth | Linewidth of border of the rectangle |
| $\ldots$ | Other parameters to be used in annotate() from ggplot2. |

## Value

Prints a patch of rectangle onto a combined plot.

## Examples

```
library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white',
            size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white', size = 3) +
    sm_hvgrid() -> p2
combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
combined_fig + sm_add_rect(xmin = .5, ymin = .5, xmax =.6, ymax =.6)
```

```
sm_add_text Add a text annotation onto the combined plot
```


## Description

Add a text annotation onto the combined plot

## Usage

```
sm_add_text(
    label,
    x = 0.5,
    y = 0.5,
    angle = 0,
    color = "black",
    fontface = "plain",
    size = 10,
)
```


## Arguments

label Text label in strings.
$x \quad$ Location of the text annotation along the $x$-axis of the combined figure. Default is the middle origin (0.5). Values from 0 to 1.
y
Location of the text annotation along the $y$-axis of the combined figure. Default is the middle origin (0.5). Values from 0 to 1.
angle Angle of the text. Default is set to 0 (i.e., horizontal orientation).

| color | Color of the text. Default is set to 'black'. |
| :--- | :--- |
| fontface | The default is to set the text as plain This can be changed, to either "plain", <br> "bold", "italic", "bold.italic" . |
| size | Size of the text annotation |
| $\ldots$ | Other parameters of the text that will be transferred to the function annotate() <br> from ggplot2. |

## Value

Prints a text in the combined plot.

## Examples

```
library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white',
            size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
    geom_point(shape = 21, fill = '#0f993d', color = 'white', size = 3) +
    sm_hvgrid() -> p2
combined_fig <- sm_put_together(list(p1,p2), ncol=2,nrow=1)
combined_fig + sm_add_text(label='My label', x = .5, y= .5)
``` gration)

\section*{Description}

This is equivalent to Matlab's trapz function.

\section*{Usage}
sm_auc (x, y)

\section*{Arguments}
x
y

This is the scalar spacing of the coordinate. When the argument for ' \(x\) ' is not provided, it will calculates the approximate integral value for ' Y ' with unit spacing based on the length of ' \(y\) '.
Numerical data. The length of x and y must be equal.

\section*{Value}

A vector that is the value from the trapezoidal integration is returned.

\section*{Examples}
```

library(smplot2)
X = c(1, 2, 3,4,5)
Y1 = c(2, 3,4, 2, 3)
Y2 = c(3,3, 3, 3, 3)
sm_auc(X,Y2)
sm_auc(X, Y1)

```

\section*{Description}

This function returns a data frame containing AUCs from a data frame that contains the original raw data. One of the two arguments 'groups' and 'conditions' must be filled. The function will throw an error if both arguments are empty.

\section*{Usage}
sm_auc_all(data, subjects, groups, conditions, x, values)

\section*{Arguments}
data Name of the variable that stores the data frame that contains the columns with the specified column names.
subjects The name of the column of the data frame that contains subjects. It must be strings.
groups The name of the column of the data frame that contains each group. It must be strings.
conditions The name of the column of the data frame that contains each condition. It must be strings.
\(x \quad\) The name of the column of the data frame that contains the \(x\)-axis points/x coordinates from which the AUC can be calculated. It must be strings. The column must not have characters.
values The name of the column of the data frame that contains the actual data, which are the \(y\)-axis points from which the AUC can be calculated. It must be strings.

\section*{Value}

Returns a data frame containing area under curve from each subject and experimental condition and/or group.

\section*{Examples}
```

library(smplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,0,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Value <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c(1, 2), each = length(day1))
Condition <- rep('Control', length(day1)*2)
df <- cbind(Subject, Value, Condition, Day)
sm_auc_all(data = df, subjects = 'Subject',values = 'Value',
conditions = 'Condition',x = 'Day')

```
sm_bar A bar plot with jittered individual points

\section*{Description}

A bar plot with jittered individual points

\section*{Usage}
```

sm_bar(
...,
bar.params = list(width = 0.7, alpha = 1, color = "transparent", fill = "gray80"),
err.params = list(size = 1, color = "black"),
point.params = list(size = 2.5, alpha = 0.65, shape = 16),
errorbar_type = "se",
point_jitter_width = 0.12,
points = TRUE,
borders = TRUE,
legends = FALSE,
seed = NULL
)

```

\section*{Arguments}
...
bar. params
err.params
point. params List of parameters for individual points, such as color, alpha, fill etc
errorbar_type This argument determines the errorbar type. If it is set to 'se', standard error bar will be shown. If it is set to 'sd' (default), the error bar will display standard deviation. If it is set to 'ci', the error bar will display \(95 \%\) confidence interval.
```

point_jitter_width

```

A numerical value that determines the degree of the jitter for each point. If its 0 , all the points will have no jitter (aligned along the y -axis).
points TRUE if points need to be shown. FALSE if points need to be hidden.
borders If the border needs to be displayed, the input should be TRUE. If the border is not needed, the input should be FALSE.
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.
seed Random seed

\section*{Value}

A bar graph generated using ggplot2

\section*{Examples}
```

library(smplot2)
library(ggplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,2,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Data <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c('Day 1', 'Day 2'), each = length(day1))
df <- cbind(Subject, Data, Day)

# with aesthetic defaults of smplot

ggplot(data = df, mapping = aes(x = Day, y = Value, color = Day)) +
sm_bar() +
scale_color_manual(values = sm_color('blue','orange'))

```
sm_bland_altman A Bland Altman plot

\section*{Description}

This function generates a Bland-Altman plot. This function requires two paired data sets as input (same length), and uses sm_statBlandAlt() to compute statistical values necessary for a Bland Altman plot. For more information on these values, please type ?sm_statBlandAlt.
The plot automatically uses sm_classic() theme. The upper dashed line indicates the upper limit (mean_diff \(+1.96 *\) sd), the middle dashed line indicates the mean difference between the two samples, and the lower dashed line indicates the lower limit (mean_diff - 1.96*sd).
To add a legend, you will need to add sm_classic(legends = TRUE). To customise the figure, you can add more geom objects.

\section*{Usage}
```

sm_bland_altman(first, second, point_size = 3.3, diff_ci = TRUE, ...)

```

\section*{Arguments}
first Data from the first repetition/session
second Data from the second repetition/session
point_size The size of the individual points. The default is set to 3.3.
diff_ci If set TRUE, then it will draw a shaded region that represents the 95 confidence interval of the difference between the two sessions from one-sample t-test. If the region (i.e. confidence interval) overlaps with zero, then there is no significant bias/difference between the two sessions/datasets. If it does not overlap with 0 , then the measurement variability is significantly large.
... Parameters of geom_point(), such as 'color', 'fill', 'shape', etc.

\section*{Value}

Prints a figure, which is the Bland-Altman plot (ggplot2 object).

\section*{Examples}
```

library(smplot2)
library(tibble)
first <- rnorm(20)
second <- rnorm(20)
df <- as_tibble(cbind(first,second))
sm_bland_altman(df$first, df$second)
\# when all 3 dashed lines are not shown, extend the range of the y-axis.

```
    sm_boxplot A boxplot with jittered individual points

\section*{Description}

A boxplot with jittered individual points

\section*{Usage}
```

    sm_boxplot(
    boxplot. params = list(notch = FALSE, fill = "gray95", color = "black", size = 0.5,
        width = 0.5, outlier.shape = NA),
    point. params = list(alpha = 0.65),
    point_jitter_width = 0.12,
    points = TRUE,
    ```
```

        borders = TRUE,
        legends = FALSE,
        seed = NULL
    )

```

\section*{Arguments}
\begin{tabular}{|c|c|}
\hline & A generic aesthetic parameter across points and the boxplot. This is optional. \\
\hline boxplot.params & List of parameters for boxplot, such as color, alpha, fill etc \\
\hline point.params & List of parameters for individual points, such as color, alpha, fill etc \\
\hline point_jitter_w & \\
\hline & A numerical value that determines the degree of the jitter for each point. If its 0 , all the points will have no jitter (aligned along the \(y\)-axis). \\
\hline points & TRUE if points need to be shown. FALSE if points need to be hidden. \\
\hline borders & If the border needs to be displayed, the input should be TRUE. If the border is not needed, the input should be FALSE. \\
\hline legends & If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE. \\
\hline seed & Random seed \\
\hline
\end{tabular}

\section*{Value}

A boxplot generated using ggplot2

\section*{Examples}
```

library(ggplot2)
library(smplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,2,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Data <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c('Day 1', 'Day 2'), each = length(day1))
df <- cbind(Subject, Data, Day)

# with the default aesthetics of smplot

ggplot(data = df, mapping = aes(x = Day, y = Value, color = Day)) +
sm_boxplot() +
scale_color_manual(values = sm_color('blue','orange'))

# Without the default aesthetics of smplot

ggplot(data = df, mapping = aes(x = Day, y = Value, color = Day)) +
sm_boxplot(boxplot.params = list()) +
scale_color_manual(values = sm_color('blue','orange'))

```

\section*{Description}

This function computes the confidence interval.

\section*{Usage}
sm_ci(data, alpha \(=0.05\), low \(=\) TRUE \()\)

\section*{Arguments}
\begin{tabular}{ll} 
data & Numerical vector of data \\
alpha & Default is set to 0.05, so that \(95 \%\) confidence interval is computed. \\
low & If its TRUE, it will compute the low tail of the confidence interval. If its FALSE, \\
& it will compute the high tail of the confidence interval.
\end{tabular}

\section*{Value}

Prints a double vector that is a single end of the specified confidence interval.

\section*{Examples}
library(smplot2)
set.seed(1)
a <- rnorm \((100,1,1)\)
sm_ci(a)
sm_ci(a, low=FALSE)
```

sm_classic A SM classical theme.

```

\section*{Description}

It has x and y axis but no grids.

\section*{Usage}
sm_classic(legends = FALSE)

\section*{Arguments}
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.

\section*{Value}

Returns a background theme as a ggplot2 object.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_classic()

```
sm_color SM custom palette of colors

\section*{Description}

This is a custom color palette that SM recommends for data visualization. It returns up to 20 different colors with a high visibility.

\section*{Usage}
sm_color(...)

\section*{Arguments}

The input has to be a character string.of a color name. There are 20 colors available from the SM palette: ''blue' ','crimson' ', 'green' ', 'purple' ', ', orange' ', ''skyblue' ', ''pink'‘, ''limegreen'‘, ''lightpurple'‘, ''brown'‘, ''red'‘, '’light-
 ''yellow', ''lightgreen'"

\section*{Value}

A character/string of hex codes

\section*{Examples}
```

library(smplot2)
sm_color('crimson')
sm_color('crimson', 'green', 'blue')

```

\section*{Description}

This function is used to create a composite figure.

\section*{Usage}
sm_common_axis(location, hmargin = 1, wmargin = 1)

\section*{Arguments}
\begin{tabular}{|c|c|}
\hline location & Location of the panel. ''topleft' ': removes x -axis title, x -axis ticklabel, y -axis title. ''topright' ': removes \(x\)-axis title, \(x\)-axis ticklabel, \(y\)-axis title, \(y\)-axis ticklabel. ''bottomleft'' \(:\) removes \(x\)-axis title, \(y\)-axis title. ''bottomright'' \(:\) removes x -axis title, y -axis title, y -axis ticklabel. ''topcenter' ': removes x -axis title, x axis ticklabel, \(y\)-axis title, \(y\)-axis ticklabel. ''bottomcenter' ': removes \(x\)-axis title, \(y\)-axis title, \(y\)-axis ticklabel. ''single' ': keeps all ticks but removes title ''centerleft'' : removes some ticks and titles ''centerright' ' : removes some ticks and titles ''center' ': removes everything \\
\hline hmargin & The amount of height of blank space between subplots. It sets the size of the empty space (i.e., margin) between panels. T he default is set to 1 , which should reduce the empty space (right and left side of each panel) between the panels. \\
\hline wmargin & The amount of width of blank space between subplots. It sets the size of the empty space (i.e., margin) between panels. \(T\) he default is set to 1 , which should reduce the empty space (right and left side of each panel) between the panels. \\
\hline
\end{tabular}

\section*{Value}

Returns a ggplot2 output with ticks removed.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,2,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Data <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c('Day 1', 'Day 2'), each = length(day1))
df <- cbind(Subject, Data, Day)

# with aesthetic defaults of smplot

ggplot(data = df, mapping = aes(x = Day, y = Value, color = Day)) +
sm_bar() +
scale_color_manual(values = sm_color('blue','orange')) +

```
```

sm_common_axis('bottomleft')

```
sm_common_legend Creating a common legend for subplots on a separate panel

\section*{Description}

Creating a common legend for subplots on a separate panel

\section*{Usage}
sm_common_legend(
\(x=0.5\),
\(y=0.5\),
title = FALSE,
direction = "vertical",
border = TRUE,
legend_spacing = 0.5,
border_color = "black",
textRatio = 1
)

\section*{Arguments}
x
y

\section*{title}

\section*{direction}
border If set TRUE, border around the legend will be created. If set FALSE, the border will be removed.
legend_spacing Spacing within the legend.
border_color Color of the legend border
textRatio Size of the text relative to the plot's default. It has been set to 1.2. The larger the textRatio, the larger the texts in the legend.

\section*{Value}

It prints a legend on a blank plot. It can be used to create a common legend for subplots.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg,
fill = as.factor(cyl))) +
geom_point(shape = 21, color = 'white', size = 3) +
sm_common_legend(x = .5, y = 0.5 , direction='horizontal',
border=FALSE)

```

\section*{Description}

Common title for combined subplots

\section*{Usage}
sm_common_title(title \(=" "\), size \(=17, x=0.5, y=0.5\), fontface \(=\) "bold")

\section*{Arguments}
\[
\begin{array}{ll}
\text { title } & \text { The input should be string. } \\
\text { size } & \text { Text size of the title. } \\
\mathrm{x} & \text { Location of the title along the } \mathrm{x} \text {-axis. Default is the middle origin (0.5). } \\
\mathrm{y} & \text { Location of the legend along the } y \text {-axis. Default is the middle origin (0.5). } \\
\text { fontface } & \begin{array}{l}
\text { The default is to set the text of the title as bold. This can be changed, to either } \\
\text { "plain", "bold", "italic", "bold.italic" . }
\end{array}
\end{array}
\]

\section*{Value}

It prints title on a blank layer of ploting.

\section*{Examples}
```

library(smplot2)
sm_common_title('My title')

```

\section*{Description}

Common x-axis label (title) for combined subplots

\section*{Usage}
sm_common_xlabel(label = "", size = 17, x = 0.5, y = 0.5, fontface = "plain")

\section*{Arguments}
\begin{tabular}{ll} 
label & The input should be string. \\
size & Text size of the label. \\
\(x\) & Location of the label along the x-axis. Default is the middle origin (0.5). \\
\(y\) & Location of the label along the y-axis. Default is the middle origin (0.5). \\
fontface & The default is to set the text of the title as plain This can be changed, to either \\
& "plain", "bold", "italic", "bold.italic" .
\end{tabular}

\section*{Value}

It returns a layer with the specified common \(x\)-axis label for combined plot.

\section*{Examples}
library (smplot2)
sm_common_xlabel('My x-axis')
sm_common_ylabel Common y-axis label (title) for combined subplots

\section*{Description}

Common y-axis label (title) for combined subplots

\section*{Usage}
sm_common_ylabel(
label = "",
size = 17,
\(x=0.5\),
\(y=0.52\),
fontface = "plain",
angle \(=90\)
)

\section*{Arguments}
\begin{tabular}{ll} 
label & The input should be string. \\
size & Text size of the label. \\
\(x\) & Location of the label along the \(x\)-axis. Default is the middle origin \((0.5)\). \\
\(y\) & Location of the label along the \(y\)-axis. Default is the middle origin \((0.5)\). \\
fontface & \begin{tabular}{l} 
The default is to set the text of the title as plain This can be changed, to either \\
"plain", "bold", "italic", "bold.italic" .
\end{tabular} \\
angle & \begin{tabular}{l} 
Orientation of the \(y\)-axis title. Default is 90 degrees.
\end{tabular}
\end{tabular}

\section*{Value}

It returns a layer with the specified common \(y\)-axis label for combined plot.

\section*{Examples}
library (smplot2)
sm_common_ylabel('My y-axis')
```

sm_corr_avgErr Superimposition of the average point with horizontal and vertical er-

``` ror bars in the correlation plot

\section*{Description}

Superimposition of the average point with horizontal and vertical error bars in the correlation plot

\section*{Usage}
```

sm_corr_avgErr(
data,
x,
y,
point.params = list(size = 2.5),
errh.params = list(height = 0),
errv.params = list(width = 0),
errorbar_type = "se",
)

```

\section*{Arguments}
data Data frame variable that is used for plotting.
x
Column of the data frame that represents the \(x\)-axis.
\(y \quad\) Column of the data frame that represents the \(y\)-axis.
point. params List of parameters for the mean point, such as color, alpha, fill etc
\begin{tabular}{ll} 
errh.params & List of parameters for the horizontal error bar, such as color, alpha, fill etc \\
errv.params & \begin{tabular}{l} 
List of parameters for the vertical points, such as color, alpha, fill etc
\end{tabular} \\
errorbar_type & \begin{tabular}{l} 
This argument determines the error bar type. If it is set to 'se', standard error \\
bar will be shown. If it is set to 'sd', the error bar will display standard deviation. \\
If it is set to 'ci' (default), the error bar will display \(95 \%\) confidence interval.
\end{tabular} \\
\(\ldots\) & A generic aesthetic parameter across points and error bars. This is optional.
\end{tabular}

\section*{Value}

A point with error bars representing the average will be returned.

\section*{Examples}
```

library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, size = 3) +
sm_corr_avgErr(mtcars, drat,mpg, errorbar_type = 'se',
color = sm_color('red'))

```
sm_effsize Cohen's \(d\)-effect size

\section*{Description}

Cohen's \(d\) is a measure of the effect size. It is often reported with p-values (ex. from a t-test or posthoc pairwise comparisons).

\section*{Usage}
sm_effsize(group1, group2, absolute = TRUE)

\section*{Arguments}
group1 Numeric vector containing data from one sample (i.e., group 1) that is to be compared with another group.
group2 Numeric vector containing data from another sample (i.e., group 2) that is to be compared with the former group.
absolute If set TRUE, the function will print the absolute value of the effect size. If set FALSE, the function will print effect size of group2 - group1. For example, it will be positive if group2 has a larger mean than group 1.

\section*{Value}

Returns a double vector that is the effect size between two samples.

\section*{Examples}
```

library(smplot2)
group1 <- rnorm(10,0,1)
group2 <- rnorm(10,1,1)
sm_effsize(group1, group2)

```
sm_forest Forest plot

\section*{Description}

Forest plot

\section*{Usage}
```

sm_forest(
point.params = list(size = 2.5, alpha = 0.3),
avgPoint.params = list(size = 5.5, shape = 18),
err.params = list(color = "black"),
ref.params = list(size = 0.4, color = "gray80", linetype = "dashed"),
xintercept = 0,
sep_level = 2,
point_jitter_width = 0,
errorbar_type = "ci",
points = TRUE,
refLine = TRUE,
borders = TRUE,
legends = FALSE,
seed = NULL
)

```

\section*{Arguments}
... A generic aesthetic parameter across points, lines and error bars. This is optional.
point. params List of parameters for individual points, such as color, alpha, fill etc avgPoint.params

List of parameters for the average point, such as color, alpha, fill etc
err.params List of parameters for the error bar from the average point, such as color, alpha etc
ref. params List of parameters for the vertical reference line, such as color, alpha etc
xintercept Location of the vertical reference line along the \(x\) coordinate.
```

sep_level A numerical value that controls the level of the separation between the individual points and the average point. If it's 0 , all of these are clustered together. If it's higher (and more positive), the text annotations will increasingly go below the mean point. Default is set to 2 . The values can be negative so that the points can be above the mean point. There is no limit of the range for this argument.
point_jitter_width
A numerical value that determines the degree of the jitter for each point. If its 0 , all the points will have no jitter (aligned along the $y$-axis).
errorbar_type This argument determines the error bar type. If it is set to 'se', standard error bar will be shown. If it is set to 'sd', the error bar will display standard deviation. If it is set to 'ci' (default), the error bar will display $95 \%$ confidence interval.
points If points is set TRUE, individual points are shown. If FALSE, they are not shown.
refLine If it is set TRUE, the reference line at a specified location along the $x$-axis is shown. If it is set FALSE, it is not shown.
borders If the border needs to be displayed, the input should be TRUE. If the border is not needed, the input should be FALSE.
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.
seed Random seed

```

\section*{Value}

A forest plot generated using ggplot2

\section*{Examples}
```

library(smplot2)
library(ggplot2)
day1 = rnorm(20,0,1)
day2 = rnorm(20,5,1)
day3 = rnorm(20,6,1.5)
day4 = rnorm(20,7,2)
Subject <- rep(paste0('S',seq(1:20)), 4)
Data <- data.frame(Value = matrix(c(day1,day2,day3,day4),ncol=1))
Day <- rep(c('Day 1', 'Day 2', 'Day 3', 'Day 4'), each = length(day1))
df2 <- cbind(Subject, Data, Day)
ggplot(data = df2, aes(x = Value, y = Day, color = Day, fill = Day)) +
sm_forest(sep_level = 2, point_jitter_width = .12,
errorbar_type = 'ci',
point.params = list(alpha=0.2, size= 2.5)) +
scale_color_manual(values = sm_palette(4))

```
```

sm_forest_annot Annotation of the error range on the forest plot

```

\section*{Description}

Annotation of the error range on the forest plot

\section*{Usage}
```

sm_forest_annot(
data,
x ,
$y$,
errorbar_type = "ci",
text.params = list(size = 4, color = "black"),
sep_level = 2,
)

```

\section*{Arguments}
data Data frame variable that is used for plotting.
\(x \quad\) Column of the data frame that represents the \(x\)-axis.
\(y \quad\) Column of the data frame that represents the \(y\)-axis.
errorbar_type This argument determines the errorbar type. If it is set to 'se', standard error bar will be shown. If it is set to 'sd' (default), the error bar will display standard deviation. If it is set to 'ci', the error bar will display \(95 \%\) confidence interval.
text. params List of parameters for the text annotation, such as color, size etc
sep_level A numerical value that controls the level of the separation between the text annotation and the average point. If it's 0 , all of these are clustered together. If it's higher (and more positive), the text annotations will increasingly go above the mean point. Default is set to 2 . The values can be negative so that the texts can be below the mean point. There is no limit of the range for this argument. Ideally, this should equal to the sep_level in sm_forest().
... Parameters for the text annotation, such as size and color etc.

\section*{Value}

Annotations showing the range of uncertainty will printed on the forest plot.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
day1 = rnorm(20,0,1)

```
```

day2 = rnorm(20,5,1)
day3 = rnorm(20,6,1.5)
day4 = rnorm(20,7,2)
Subject <- rep(paste0('S',seq(1:20)), 4)
Data <- data.frame(Value = matrix(c(day1,day2,day3,day4),ncol=1))
Day <- rep(c('Day 1', 'Day 2', 'Day 3', 'Day 4'), each = length(day1))
df2 <- cbind(Subject, Data, Day)
ggplot(data = df2, aes(x = Value, y = Day, color = Day)) +
sm_forest(point_jitter_width = 0.12, sep_level = 3) +
scale_color_manual(values = sm_palette(4)) +
sm_forest_annot(data = df2, x = Value, y = Day, sep_level = 3)

```

\section*{Description}

A graph with a horizontal grid is plotted. Border can be added or removed. This is useful for plotting a bar graph.

\section*{Usage}
sm_hgrid(legends = FALSE, borders = TRUE)

\section*{Arguments}
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.
borders If the border needs to be displayed, the input should be TRUE. If the border is not needed, the input should be FALSE.

\section*{Value}

Returns a background theme with major horizontal grids (ggplot2 output).

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_hgrid()

```

\section*{Description}

Histogram with kernel density estimation (Gaussian) and rugs

\section*{Usage}
```

sm_hist(
...,
hist.params = list(binwidth = 1/2, fill = sm_color("blue"), color = "white", alpha =
0.4),
density.params = list(color = sm_color("blue"), size = 0.8, fill = "transparent"),
rug.params = list(color = sm_color("blue"), alpha = 0.8, size = 0.4),
histogram = TRUE,
density = TRUE,
rug = TRUE,
borders = FALSE,
legends = FALSE
)

```

\section*{Arguments}
\begin{tabular}{ll}
\(\ldots\). & A generic aesthetic parameter across points and the boxplot. This is optional. \\
hist. params & List of parameters for the histogram, such as binwidth, color, alpha, fill etc. \\
density.params \\
rug. params & \begin{tabular}{l} 
List of parameters for the density estimation, such as color, size, alpha etc
\end{tabular} \\
histogram & \begin{tabular}{l} 
List of parameters for the rugs, such as color, size, alpha etc \\
TRUE if the histogram needs to be shown. FALSE if the histogram needs to be \\
hidden.
\end{tabular} \\
density & \begin{tabular}{l} 
TRUE if the density plot needs to be shown. FALSE if the density plot needs to \\
be hidden.
\end{tabular} \\
rug & \begin{tabular}{l} 
TRUE if the rugs need to be shown. FALSE if the rugs need to be hidden.
\end{tabular} \\
borders & \begin{tabular}{l} 
If the border needs to be displayed, the input should be TRUE. If the border is \\
not needed, the input should be FALSE.
\end{tabular} \\
legends & \begin{tabular}{l} 
If the legend needs to be displayed, the input should be TRUE. If the legend is \\
not needed, the input should be FALSE.
\end{tabular}
\end{tabular}

\section*{Value}

Returns a histogram generated using ggplot2.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
set.seed(2)
data=data.frame(value=rnorm(1000))
data2 = data.frame(value=rnorm(1000,5,1))
data$day <- 'day1'
data2$day <- 'day2'
rbind(data,data2) -> df
ggplot(data = data, aes(x=value)) +
sm_hist()
ggplot(data = df, aes(x=value, fill=day, color = day)) +
sm_hist(hist.params = list(binwidth = 1/2, alpha = 0.3),
density.params = list(fill='transparent', size = 0.8),
rug.params = list(alpha = 0.8)) +
scale_color_manual(values = sm_palette(2)) +
scale_fill_manual(values = sm_palette(2))

```
sm_hvgrid Minimalistic theme with major horizontal and vertical grids

\section*{Description}

This theme has major vertical and horizontal grids. This is useful for plotting correlations. sm_corr_theme() is exactly the same as sm_hvgrid().

\section*{Usage}
sm_hvgrid(legends = TRUE, borders = TRUE)

\section*{Arguments}
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.
borders If the border needs to be displayed, the input should be 'TRUE'. If the border is not needed, the input should be 'FALSE'.

\section*{Value}

Returns a background theme that has both horizontal and vertical major grids (ggplot2 output).

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_hvgrid()

```
sm_hvgrid_minor A theme with horizontal and vertical major and minor grids

\section*{Description}

This theme has vertical and horizontal grids.

\section*{Usage}
sm_hvgrid_minor(legends = TRUE, borders = TRUE)

\section*{Arguments}
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.
borders If the border needs to be displayed, the input should be 'TRUE'. If the border is not needed, the input should be 'FALSE'.

\section*{Value}

Returns a background theme that has both horizontal and vertical major and minor grids (ggplot2 output).

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_hvgrid_minor()

```
sm_minimal SM minimal theme (no grid) with borders.

\section*{Description}

This theme has no major grid.

\section*{Usage}
sm_minimal(legends = FALSE)

\section*{Arguments}
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.

\section*{Value}

Returns a background theme that has no grids (ggplot2 output).

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_minimal()

```
sm_palette SM custom palette of colors

\section*{Description}

This is a custom color palette that SM recommends for data visualization. It returns up to 20 different colors with a high contrast.

\section*{Usage}
```

    sm_palette(colorNum = 10)
    ```

\section*{Arguments}
colorNum \(\quad\) Number of colors (1-20).

\section*{Value}

Returns a hex code in string vector. The input determines the length of the output.

\section*{Examples}
library(smplot2)
\#' sm_palette(3) \# returns 3 colors

Writing a label for each panel of a combined figure

\section*{Description}

Writing a label for each panel of a combined figure

\section*{Usage}
```

sm_panel_label(
all_plots,
x,
y,
panel_tag = "1",
panel_pretag,
panel_posttag,
text_size = 5.5,
text_color = "black",
fontface = "plain",
)

```

\section*{Arguments}
all_plots all_plots should be a list vector, which should contain all panels that are to be combined into one figure.
\(x \quad\) Location of the label along the \(x\)-axis ( 0 to 1 ). 0.5 is the middle origin.
\(y \quad\) Location of the label along the \(y\)-axis ( 0 to 1 ). 0.5 is the middle origin.
panel_tag A character vector that defines how each panel is enumerated. Options include: 'a', ' A ', ' 1 ', ' I ' or ' i '.' a ' is for lowercase letters. ' A ' is for uppercase letters. ' 1 ' is for integers. ' I ' is for upper case roman numerals. ' i ' is for lower case roman numerals. Each panel will display a unique string based on the set enumeration.
panel_pretag A character vector that is identical across panels BEFORE the panel_tag.
panel_posttag A character vector that is identical across panels AFTER the panel_tag.
text_size Text size of the panel label
text_color Text color of the panel label
\begin{tabular}{ll} 
fontface & \begin{tabular}{l} 
Fontface of the panel label. Options include "plain", "bold", "italic" and others \\
that are provided by ggplot2.
\end{tabular} \\
\(\ldots\) & \begin{tabular}{l} 
Additional parameters for adjusting the appearance of the sticker. Same param- \\
eters for annotate() from ggplot2.
\end{tabular}
\end{tabular}

\section*{Value}

It returns a list of plots with panel labels.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white',
size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white', size = 3) +
sm_hvgrid() -> p2
sm_panel_label(list(p1,p2), x = 0.1, y = 0.9,
panel_tag ='1', panel_pretag = 'S', text_size = 4, text_color = 'black')

```
sm_plot_clean Remove xticklabels and yticklabels in selected panels for proper sub-
    plotting

\section*{Description}

Remove xticklabels and yticklabels in selected panels for proper subplotting

\section*{Usage}
sm_plot_clean(all_plots, ncol, nrow, wmargin = wmargin, hmargin = hmargin)

\section*{Arguments}
all_plots all_plots should be list, which should contain all panels that are to be combined into one figure.
ncol Number of columns in the combined plot
nrow Number of rows in the combined plot
wmargin The amount of width of blank space between subplots. It sets the size of the empty space (i.e., margin) between panels. T he default is set to 1 , which should reduce the empty space (right and left side of each panel) between the panels.
```

hmargin The amount of height of blank space between subplots. It sets the size of the
empty space (i.e., margin) between panels. T he default is set to 1, which should
reduce the empty space (right and left side of each panel) between the panels.

```

\section*{Value}

Returns a list of plots with new layouts.

\section*{Examples}
```

library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white',
size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white', size = 3) +
sm_hvgrid() -> p2
sm_plot_clean(list(p1,p2), ncol=2,nrow=1,wmargin=-2, hmargin=-2)

```
```

sm_pointplot

```

Point plot with optional shadow

\section*{Description}

This is a common plot with mean point, standard error (se, sd or 95 uniquely shadow, which is a faint display of individual points behind the mean.

\section*{Usage}
```

sm_pointplot(
...,
avgPoint.params = list(size = 2.5),
avgLine.params = list(),
point.params = list(alpha = 0.35, color = "gray", fill = "gray"),
line.params = list(alpha = 0.35, color = "gray"),
err.params = list(size = 1),
errorbar_type = "se",
show_shadow = FALSE,
group = NULL,
borders = TRUE,
legends = FALSE
)

```

\section*{Arguments}
\begin{tabular}{|c|c|}
\hline & A generic aesthetic parameter across points, lines and errorbars. This is optional. This will be extremely useful for dodging each line using position_dodge(). \\
\hline \multicolumn{2}{|l|}{avgPoint.params} \\
\hline & List of parameters for the average point, such as color, alpha, fill etc \\
\hline avgLine.params & List of parameters for the average line, such as color, alpha etc \\
\hline point.pa & List of parameters for the points in the shadow, such as color, alpha, fill etc \\
\hline line.params & List of parameters for the lines in the shadow, such as color, alpha etc \\
\hline err.params & List of parameters for the error bar from the average plot, such as color, alpha etc \\
\hline errorbar_typ & This argument determines the errorbar type. If it is set to 'se', standard error bar will be shown. If it is set to 'sd' (default), the error bar will display standard deviation. If it is set to 'ci', the error bar will display \(95 \%\) confidence interval. \\
\hline show_shadow & If it is TRUE, it will show the shadow. If it is FALSE, it will not show the shadow (default). \\
\hline group & If show_shadow \(=\) TRUE, this argument is required. This is the variable that each plot from the shadow should be grouped along aesthetically. It should be grouped for each individual observation, ex. sm_pointplot(group = Subject), whereby Subject is the column that holds identifers for each observation. \\
\hline borders & If the border needs to be displayed, the input should be TRUE. If the border is not needed, the input should be FALSE. \\
\hline legends & If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE. \\
\hline
\end{tabular}

\section*{Value}

Returns a pointplot generated using ggplot2.

\section*{Examples}
library (smplot2)
library (ggplot2)
\(\operatorname{ggplot}(\) data \(=\) mtcars, mapping \(=\operatorname{aes}(x=c y l, y=m p g))+\)
sm_pointplot()

\section*{sm_power}

Post-hoc power analysis using two-sample or paired t-test

\section*{Description}

Post-hoc power analysis using two-sample or paired t-test

\section*{Usage}
sm_power (group1, group2, paired, sig.level = 0.05, power = 0.8)

\section*{Arguments}
group1 Numeric vector containing data from one sample (i.e., group 1) that is to be compared with another group.
group2 Numeric vector containing data from another sample (i.e., group 2) that is to be compared with the former group.
paired A logical indicating whether your two samples (group1 and group2) are paired.
sig.level Significance level (Type I error probability). Default is set to 0.05.
power \(\quad\) Power of test ( 1 minus Type II error probability). Default is set to 0.8 .

\section*{Value}

Returns a result with a class of "power.htest" from the pwr package.

\section*{Examples}
```

library(smplot2)
group1 <- rnorm(10,0,1)
group2 <- rnorm(10,1,1)
sm_power(group1, group2, paired = TRUE)

```
sm_put_together Combining figures together

\section*{Description}

Combining figures together

\section*{Usage}
```

sm_put_together(
all_plots,
title,
xlabel,
ylabel,
legend,
ncol,
nrow,
xlabel2,
ylabel2,
tickRatio,
panel_scale = 0.9,

```
```

    wRatio = 1.1,
    hRatio = 1.1,
    hmargin = 0,
    wmargin = 0,
    remove_ticks = "some",
    wRatio2 = 1.1,
    hRatio2 = 1.1
    )

```

\section*{Arguments}
\(\left.\begin{array}{ll}\text { all_plots } & \begin{array}{l}\text { all_plots should be list, which should contain all panels that are to be combined } \\ \text { into one figure. }\end{array} \\ \text { title } \\ \text { Title layer that will determine the main title of the combined plot. This is created } \\ \text { using sm_common_title(). Optional argument. } \\ \text { xlabel layer that will determine the label of the combined plot's x-axis. This is } \\ \text { created using sm_common_xlabel(). Optional argument. } \\ \text { ylabel layer that will determine the label of the combined plot's y-axis. This is } \\ \text { created using sm_common_ylabel(). Optional argument. } \\ \text { ggplot() layer that has legend. Optional argument. }\end{array}\right\}\)
wmargin \begin{tabular}{l} 
The amount of width of blank space between subplots. It sets the size of the \\
empty space (i.e., margin) between panels. T he default is set to 0 . If its positive, \\
the blank spacing will increase. If its negative, it will get reduced between \\
panels. \\
remove_ticks \begin{tabular}{l} 
If set to 'some', x-axis ticks and y-axis ticks will be removed in inner plots. If set \\
to 'all', then all panels' ticks will be removed. If set to 'none', then all panels' \\
ticks will be kept.
\end{tabular} \\
wRatio2 \\
This adjusts the ratio of the width of the last column to those of other columns. \\
By default, if ylabel2 is provided, it is set to be 1.1 x wider than that of other \\
columns. If the value is larger than 1 , then it will be wider than that of other \\
columns. Users are encouraged to adjust this value because different computers \\
can show different looking outputs.
\end{tabular}
hRatio2 \begin{tabular}{l} 
This adjusts the ratio of the height of the first row to those of other rows By \\
default, if xlabel2 is provided, it is set to be 1.1 x taller than that of other columns.
\end{tabular}
If the value is larger than 1 , then it will be taller than that of other columns.
Users are encouraged to adjust this value because different computers can show
different looking outputs.

\section*{Value}

Returns a combined figure.

\section*{Examples}
```

library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white',
size = 3) -> p1
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white', size = 3) +
sm_hvgrid() -> p2
title <- sm_common_title('My title')
xlabel <- sm_common_xlabel('My x-axis')
ylabel <- sm_common_ylabel('My y-axis')
sm_put_together(list(p1,p2), title=title, xlabel=xlabel,
ylabel=ylabel, ncol=2,nrow=1)

```

\section*{Description}

This function visualizes a raincloud plot, which is a combination of jittered points, boxplots and violin plots. The creation of this function has been inspired by the R package called 'raincloudplots' by Jordy van Langen.
This function has been created to allow more customisation than the functions in the raincloudplots package. Also, this function automatically sorts the data given the condition that the x -axis factor levels have been sorted properly.

\section*{Usage}
```

sm_raincloud(
boxplot.params = list(),
violin.params = list(alpha = 0.3, color = "transparent"),
point.params = list(alpha = 1, size = 3, shape = 21, color = "transparent"),
which_side = "r",
sep_level = 2,
point_jitter_width = 0.12,
vertical = TRUE,
points = TRUE,
borders = TRUE,
legends = FALSE,
seed = NULL
)

```

\section*{Arguments}
... A generic aesthetic parameter across points, boxplot and violin. This is optional.
boxplot. params List of parameters for boxplot, such as color, alpha, fill etc
violin. params List of parameters for violin, such as color, alpha, fill etc
point. params List of parameters for individual points, such as color, alpha, fill etc
which_side String argument to specify the side of the boxplots and violinplots. The options are: 'right' and 'left'. 'mixed' has been removed from smplot due to its lack of usage.
sep_level A numerical value that controls the level of the separation among the boxplot, violin plot and the points. The value can be \(0-4\). If it's 0 , all of these are clustered together. If it's 3 , they are all separated. 1 and 2 are somewhere in the middle. Default is set to 2 .
point_jitter_width
A numerical value that determines the degree of the jitter for each point. If its 0 , all the points will have no jitter (aligned along the \(y\)-axis).
vertical The orientation of the plots. The default is set to TRUE. If you want the horizontal orientation of the plot, set this argument as FALSE.
points If the points need to be displayed, the input should be TRUE. If the points are not needed, the input should be FALSE.
\begin{tabular}{ll} 
borders & If the border needs to be displayed, the input should be TRUE. If the border is \\
not needed, the input should be FALSE.
\end{tabular}

\section*{Value}

Returns a raincloud plot generated using ggplot2.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
set.seed(2) \# generate random data
day1 = rnorm(20,0,1)
day2 = rnorm(20,5,1)
day3 = rnorm(20,6,1.5)
day4 = rnorm(20,7,2)
Subject <- rep(paste0('S',seq(1:20)), 4)
Data <- data.frame(Value = matrix(c(day1,day2,day3,day4),ncol=1))
Day <- rep(c('Day 1', 'Day 2', 'Day 3', 'Day 4'), each = length(day1))
df2 <- cbind(Subject, Data, Day)
ggplot(data=df2, aes(x = Day, y = Value, color = Day, fill = Day)) +
sm_raincloud() +
xlab('Day') +
scale_fill_manual(values = sm_palette(4))

```
    sm_slope A slope chart

\section*{Description}

This function generates a slope chart. This is very useful for comparing the effect between two time points.

For this function to work properly, ggplot()'s mapping has to be quite specific. 1. Each observation has to be grouped.
With smplot2, average plot can be displayed along with error bars, both of which are optional.

\section*{Usage}
```

sm_slope(
...,
labels,
group,
line.params = list(color = "gray53", size = 0.4, alpha = 0.4),
point.params = list(size = 2.5, shape = 21, color = "white"),
avgLine.params = list(),
avgPoint.params = list(),
err.params = list(),
xTick.params = list(position = "top", expand = c(0.17, 0.1), drop = FALSE),
errorbar_type = "sd",
many_groups = FALSE,
show_err = FALSE,
show_mean = FALSE,
legends = FALSE
)

```

\section*{Arguments}
\begin{tabular}{ll}
\(\ldots\). & \begin{tabular}{l} 
List of parameters for individual points and lines across different elements (ex- \\
cept for except for xTick.params), such as color, alpha, fill etc.
\end{tabular} \\
labels & \begin{tabular}{l} 
Labels for the ticks of the x-axis. This is a required argument. It has to be a \\
single vector containing either one or multiple elements. ex: c('Day 1', 'Day \\
\(2 ')\)
\end{tabular} \\
group & Name of the variable by which the individual data should be grouped \\
line.params & List of parameters for the individual lines, such as color, alpha etc \\
point.params & List of parameters for the individual points, such as color, alpha, fill etc \\
avgLine.params \\
avgPoint.params
\end{tabular}
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.

\section*{Value}

Returns a double vector output representing the slope from linear regression.

\section*{Examples}
```

library(ggplot2)
library(smplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,2,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Data <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c('Day 1', 'Day 2'), each = length(day1))
df <- cbind(Subject, Data, Day)
ggplot(data=df, aes(x = Day, y = Value, fill = Day)) +
sm_slope(labels = c('Day 1', 'Day 2'), group = Subject) +
scale_fill_manual(values= sm_color('blue','orange'))
ggplot(data = df, aes(x = Day, y = Value, fill = Day)) +
sm_slope(labels = c('Day 1','Day 2'),group = Subject,
point.params = list(alpha = 0.3, size = 2.5, color = 'white',
shape = 21, fill = sm_color('skyblue')),
line.params = list(color = sm_color('skyblue'),
alpha = 0.3),
avgPoint.params = list(color='transparent', shape = 21,
size = 4, fill = sm_color('blue')),
avgLine.params = list(color = sm_color('blue'), size = 1),
show_mean = TRUE)

```

\section*{Description}

This function returns a data frame containing slope (from linear regression) from a data frame that contains the original raw data.

The user can use \(\operatorname{lm}()\) from base R to compute the slope as well.

\section*{Usage}
```

sm_slope_all(data, subjects, groups, conditions, x, values)

```

\section*{Arguments}
\begin{tabular}{ll} 
data & \begin{tabular}{l} 
Name of the variable that stores the data frame that contains the columns with \\
the specified column names.
\end{tabular} \\
subjects & \begin{tabular}{l} 
The name of the column of the data frame that contains subjects. It must be \\
strings.
\end{tabular} \\
groups & \begin{tabular}{l} 
The name of the column of the data frame that contains each group. It must be \\
strings.
\end{tabular} \\
conditions & \begin{tabular}{l} 
The name of the column of the data frame that contains each condition. It must \\
be strings.
\end{tabular} \\
x & \begin{tabular}{l} 
The name of the column of the data frame that contains the \(x\)-axis points \(/ x\) coor- \\
dinates from which the slopes can be calculated. It must be strings. The column \\
must not have characters.
\end{tabular} \\
values & \begin{tabular}{l} 
The name of the column of the data frame that contains the actual data, which \\
are the y-axis points from which the slope can be calculated. It must be strings.
\end{tabular}
\end{tabular}

\section*{Value}

Returns a data frame containing slopes for each subject and each experimental condition and/or group.

\section*{Examples}
```

library(smplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,0,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Value <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c(1,2), each = length(day1))
Condition <- rep('Control', length(day1)*2)
df <- cbind(Subject, Value, Condition, Day)
sm_slope_all(data = df, subjects = 'Subject',values = 'Value',
conditions = 'Condition',x = 'Day')

```
sm_slope_theme SM plot with a theme appropriate for the slope chart

\section*{Description}

In this plot, all aspects except for the left-handed spine are missing. This format is appropriate for the slope chart.

\section*{Usage}
```

sm_slope_theme(legends = TRUE)

```

\section*{Arguments}
legends \#' If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.

\section*{Value}

Returns a background theme that is suitable for a slope chart (ggplot2 output).

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_slope_theme()

```
sm_statBlandAlt Statistics for a Bland-Altman plot

\section*{Description}

Bland-Altman plot is drawn to show measurement variability/reliabiilty of a task. This function requires two paired datasets (same length). It returns a list of difference (by element), mean, standard deviation of the difference, mean difference, upper and lower limits. These values are necessary to draw a Bland Altman plot.

The list returned from this function can be directly used as an argument for sm_bland_altman(), which draws a Bland-Altman plot using ggplot2.
Another output 'data' is a tibble with two columns: 1) Mean across each pair for each element (ex. a mean of the 1st element from the first set and 1st element from the second set), 2) Difference between each pair for every element. The output 'data' should be used as a argument for data in ggplot() when plotting.

\section*{Usage}
sm_statBlandAlt(first, second)

\section*{Arguments}
first Data from the first repetition/session
second Data from the second repetition/session

\section*{Value}

A list is returned, which has all numerical results that are relevant to drawing a Bland-Altman plot.

\section*{Examples}
```

library(smplot2)
library(tibble)
first <- rnorm(20)
second <- rnorm(20)
df <- as_tibble(cbind(first,second)) \# requires library(tidyverse)
sm_statBlandAlt(df$first, df$second)

```
```

sm_statCorr

```

Linear regression slope and statistical values from a paired correlation test

\section*{Description}

This combines two different functions: 1) 'geom_smooth()' from ggplot, and 2) 'stat_cor()' from ggpubr. 'geom_smooth()' is used to fit the best-fit model, whereas 'stat_cor()' is used to print correlation results at an optimized location.
Updates from smplot2 include more flexibility, less input arguments and its pairing with 'sm_hvgrid()' / 'sm_corr_theme()'.

\section*{Usage}
```

sm_statCorr(
...,
fit.params = list(),
corr_method = "pearson",
alternative = "two.sided",
separate_by = ",",
label_x = NULL,
label_y = NULL,
text_size = 4,
show_text = TRUE,
borders = TRUE,
legends = FALSE
)

```

\section*{Arguments}
... Arguments for the properties of regression line, such as 'linetype', 'color', etc. For more information, type ?geom_smooth
fit.params Paramters for the fitted line, such as color, linetype and alpha.
\begin{tabular}{ll} 
corr_method & \begin{tabular}{l} 
Method of the correlation test. Options include: 'pearson', 'kendall', or 'spear- \\
man'.
\end{tabular} \\
alternative & \begin{tabular}{l} 
Specifies the alternative hypothesis (H1). 'two.sided' is the standard way. 'greater' \\
is a positive association, whereas 'less' is a negative association.
\end{tabular} \\
separate_by & \begin{tabular}{l} 
This marks how the p- and r- values should be separated. The default option is: \\
,, , For more information, check out stat_cor() from the ggpubr package. \\
Location of the statistical value prints along the figure's x-axis. It asks for a \\
number within the x-axis limit.
\end{tabular} \\
label_x & \begin{tabular}{l} 
Location of the statistical value prints along the figure's y-axis. It requires a \\
number within the y-axis limit.
\end{tabular} \\
label_y & \begin{tabular}{l} 
Size (numerical value) of the texts from correlation.
\end{tabular} \\
text_size & \begin{tabular}{l} 
If the statistical result needs to be displayed, the input should be TRUE (default). \\
If the statistical result is not needed, the input should be FALSE.
\end{tabular} \\
show_text & \begin{tabular}{l} 
If the border needs to be displayed, the input should be TRUE. If the border is \\
not needed, the input should be FALSE.
\end{tabular} \\
legends & \begin{tabular}{l} 
If the legend needs to be displayed, the input should be TRUE. If the legend is \\
not needed, the input should be FALSE.
\end{tabular}
\end{tabular}

\section*{Value}

Plots a best-fitted linear regression on a correlation plot with results from correlation statistical tests.

\section*{Examples}
```

library(smplot2)
library(ggplot2)
ggplot(data = mtcars, mapping = aes(x = drat, y = mpg)) +
geom_point(shape = 21, fill = '\#0f993d', color = 'white', size = 3) +
sm_statCorr()

```
    sm_stdErr Standard error

\section*{Description}

Standard error

\section*{Usage}
sm_stdErr (data)

\section*{Arguments}
data
Numerical vector of data.

\section*{Value}

A double vector is returned with a standard error of the input (given sample).

\section*{Examples}
```

library(smplot2)
sm_stdErr(rnorm(10,0,1))

```
sm_vgrid Minimalistic theme with vertical major grids

\section*{Description}

This theme has major vertical grids.

\section*{Usage}
sm_vgrid(legends = TRUE, borders = TRUE)

\section*{Arguments}
\begin{tabular}{ll} 
legends & If the legend needs to be displayed, the input should be TRUE. If the legend is \\
not needed, the input should be FALSE. \\
borders & If the border needs to be displayed, the input should be TRUE. If the border is \\
not needed, the input should be FALSE.
\end{tabular}

\section*{Value}

Returns a background theme with major vertical grids (ggplot2 output).

\section*{Examples}
```

library(ggplot2)
library(smplot2)
ggplot(data = mpg) +
geom_point(mapping = aes(x = displ, y = hwy, color = class)) +
sm_vgrid()

```
```

sm_violin

```

A violin plot with jittered individual points

\section*{Description}

A violin plot with jittered individual points

\section*{Usage}
sm_violin(
...,
violin.params = list(fill = "gray90", color = "transparent"),
err.params = list(size = 1.2),
point.params = list(alpha = 0.2),
errorbar_type = "sd",
point_jitter_width = 0.17,
points = TRUE,
borders = TRUE,
legends = FALSE,
seed \(=\) NULL
)

\section*{Arguments}
... A generic aesthetic parameter across points and the violin plot. This is optional.
violin. params List of parameters for the violin, such as color, alpha, fill etc
err.params List of parameters for the error bar, such as color, size, alpha etc
point. params List of parameters for individual points, such as color, alpha, fill etc
errorbar_type This argument determines the errorbar type. If it is set to 'se', standard error bar will be shown. If it is set to 'sd' (default), the error bar will display standard deviation. If it is set to 'ci', the error bar will display \(95 \%\) confidence interval.
point_jitter_width
A numerical value that determines the degree of the jitter for each point. If its 0 , all the points will have no jitter (aligned along the \(y\)-axis).
points TRUE if points need to be shown. FALSE if points need to be hidden.
borders If the border needs to be displayed, the input should be TRUE. If the border is not needed, the input should be FALSE.
legends If the legend needs to be displayed, the input should be TRUE. If the legend is not needed, the input should be FALSE.
seed Random seed

\section*{Value}

Violin plot generated using ggplot2

\section*{Examples}
```

library(ggplot2)
library(smplot2)
set.seed(1) \# generate random data
day1 = rnorm(16,2,1)
day2 = rnorm(16,5,1)
Subject <- rep(paste0('S',seq(1:16)), 2)
Data <- data.frame(Value = matrix(c(day1,day2),ncol=1))
Day <- rep(c('Day 1', 'Day 2'), each = length(day1))
df <- cbind(Subject, Data, Day)

# with aesthetic defaults of smplot

ggplot(data = df, mapping = aes(x = Day, y = Value, color = Day)) +
sm_violin() +
scale_color_manual(values = sm_color('blue','orange'))

# without aesthetic defaults of smplot

ggplot(data = df, mapping = aes(x = Day, y = Value, color = Day)) +
sm_violin(violin.params = list()) +
scale_color_manual(values = sm_color('blue','orange'))

```

\section*{Index}
```

sm_add_arrow, 2
sm_add_legend,4
sm_add_line,5
sm_add_point,6
sm_add_polygon, 7
sm_add_rect, 8
sm_add_text, 9
sm_auc, 10
sm_auc_all,11
sm_bar,12
sm_bland_altman, 13
sm_boxplot,14
sm_ci,16
sm_classic,16
sm_color,17
sm_common_axis,18
sm_common_legend, 19
sm_common_title, 20
sm_common_xlabel, 21
sm_common_ylabel, 21
sm_corr_avgErr, 22
sm_effsize, 23
sm_forest, 24
sm_forest_annot,26
sm_hgrid, 27
sm_hist, 28
sm_hvgrid, 29
sm_hvgrid_minor,30
sm_minimal, 31
sm_palette, 31
sm_panel_label, 32
sm_plot_clean, 33
sm_pointplot, 34
sm_power, 35
sm_put_together, 36
sm_raincloud, 38
sm_slope,40
sm_slope_all, 42
sm_slope_theme, 43

```
```

