Package ‘ggpubr’

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**Type**  Package
**Title**  'ggplot2' Based Publication Ready Plots
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**Description**  The 'ggplot2' package is excellent and flexible for elegant data visualization in R. However the default generated plots requires some formatting before we can send them for publication. Furthermore, to customize a 'ggplot', the syntax is opaque and this raises the level of difficulty for researchers with no advanced R programming skills. 'ggpubr' provides some easy-to-use functions for creating and customizing 'ggplot2'- based publication ready plots.

**License**  GPL (>= 2)
**LazyData**  TRUE
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**Imports**  ggrepel (>= 0.9.2), grid, ggsci, stats, utils, tidyr (>= 1.3.0), purrr, dplyr (>= 0.7.1), cowplot (>= 1.1.1), ggsignif, scales, gridExtra, glue, polynom, rlang (>= 0.7.2), tibble, magrittr

**Suggests**  grDevices, knitr, RColorBrewer, gtable, testthat
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**BugReports**  https://github.com/kassambara/ggpubr/issues
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R topics documented:


NeedsCompilation no

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add_summary

Add Summary Statistics onto a ggplot.

Description

add summary statistics onto a ggplot.

Usage

add_summary(
  p,
  fun = "mean_se",
  error.plot = "pointrange",
  color = "black",
  fill = "white",
  group = 1,
  width = NULL,
  shape = 19,
  size = 1,
  linetype = 1,
  show.legend = NA,
  ci = 0.95,
  data = NULL,
  position = position_dodge(0.8)
)

mean_se_(x, error.limit = "both")

mean_sd(x, error.limit = "both")

mean_ci(x, ci = 0.95, error.limit = "both")

mean_range(x, error.limit = "both")
add_summary

median_iqr(x, error.limit = "both")
median_hilow(x, ci = 0.95, error.limit = "both")
median_q1q3(x, error.limit = "both")
median_mad(x, error.limit = "both")
median_range(x, error.limit = "both")

Arguments

p a ggplot on which you want to add summary statistics.
fun a function that is given the complete data and should return a data frame with variables ymin, y, and ymax. Allowed values are one of: "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range".
error.plot plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange".
color point or outline color.
fill fill color. Used only when error.plot = "crossbar".
group grouping variable. Allowed values are 1 (for one group) or a character vector specifying the name of the grouping variable. Used only for adding statistical summary per group.
width numeric value between 0 and 1 specifying bar or box width. Example width = 0.8. Used only when error.plot is one of c("crossbar", "errorbar").
shape point shape. Allowed values can be displayed using the function show_point_shapes().
size numeric value in [0-1] specifying point and line size.
linetype line type.
show.legend logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.
ci the percent range of the confidence interval (default is 0.95).
data a data.frame to be displayed. If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot.
position position adjustment, either as a string, or the result of a call to a position adjustment function. Used to adjust position for multiple groups.
x a numeric vector.
error.limit allowed values are one of ("both", "lower", "upper", "none") specifying whether to plot the lower and/or the upper limits of error interval.
Functions

- add_summary(): add summary statistics onto a ggplot.
- mean_se(): returns the mean and the error limits defined by the standard error. We used the name mean_se() to avoid masking mean_se().
- mean_sd(): returns the mean and the error limits defined by the standard deviation.
- mean_ci(): returns the mean and the error limits defined by the confidence interval.
- mean_range(): returns the mean and the error limits defined by the range = max - min.
- median_iqr(): returns the median and the error limits defined by the interquartile range.
- median_hilow(): computes the sample median and a selected pair of outer quantiles having equal tail areas. This function is a reformatted version of Hmisc::median.hilow(). The confidence limits are computed as follow: lower.limits = (1-ci)/2 percentiles; upper.limits = (1+ci)/2 percentiles. By default (ci = 0.95), the 2.5th and the 97.5th percentiles are used as the lower and the upper confidence limits, respectively. If you want to use the 25th and the 75th percentiles as the confidence limits, then specify ci = 0.5 or use the function median_q1q3().
- median_q1q3(): computes the sample median and, the 25th and 75th percentiles. Wrapper around the function median_hilow() using ci = 0.5.
- median_mad(): returns the median and the error limits defined by the median absolute deviation.
- median_range(): returns the median and the error limits defined by the range = max - min.

Examples

```r
# Basic violin plot
p <- ggviolin(ToothGrowth, x = "dose", y = "len", add = "none")
p

# Add mean_sd
add_summary(p, "mean_sd")
```

Annotate figures including: i) ggplots, ii) arranged ggplots from ggarrange(), grid.arrange() and plot_grid().
Usage
annotate_figure(
  p,
  top = NULL,
  bottom = NULL,
  left = NULL,
  right = NULL,
  fig.lab = NULL,
  fig.lab.pos = c("top.left", "top", "top.right", "bottom.left", "bottom",
                   "bottom.right"),
  fig.lab.size,
  fig.lab.face
)

Arguments
p  (arranged) ggplots.
top, bottom, left, right
   optional string, or grob.
fig.lab  figure label (e.g.: "Figure 1").
fig.lab.pos  position of the figure label, can be one of "top.left", "top", "top.right", "bottom.left", "bottom", "bottom.right". Default is "top.left".
fig.lab.size  optional size of the figure label.
fig.lab.face  optional font face of the figure label. Allowed values include: "plain", "bold", "italic", "bold.italic".

Author(s)
Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

See Also
ggarrange()

Examples

data("ToothGrowth")
df <- ToothGrowth
df$dose <- as.factor(df$dose)

  # Create some plots
  # :::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::
  # Box plot
  bxp <- ggboxplot(df, x = "dose", y = "len",
                   color = "dose", palette = "jco")
  # Dot plot
  dp <- ggdotplot(df, x = "dose", y = "len",
                  color = "dose", palette = "jco")
# Density plot
dens <- ggdensity(df, x = "len", fill = "dose", palette = "jco")

# Arrange and annotate
figure <- ggarrange(bxp, dp, dens, ncol = 2, nrow = 2)
annotate_figure(figure,
  top = text_grob("Visualizing Tooth Growth", color = "red", face = "bold", size = 14),
  bottom = text_grob("Data source: \n ToothGrowth data set", color = "blue",
                      hjust = 1, x = 1, face = "italic", size = 10),
  left = text_grob("Figure arranged using ggpubr", color = "green", rot = 90),
  right = text_grob(bquote("Superscript: ("*kg-NH[3]-ha^-1-yr^-1*x")"), rot = 90),
  fig.lab = "Figure 1", fig.lab.face = "bold"
)

---

as_ggplot Storing grid.arrange() arrangeGrob() and plots

### Description
Transform the output of `arrangeGrob()` and `grid.arrange()` to a an object of class `ggplot`.

### Usage

```r
as_ggplot(x)
```

### Arguments

- **x**: an object of class `gtable` or `grob` as returned by the functions `arrangeGrob()` and `grid.arrange()`.

### Value
an object of class `ggplot`.

### Examples

```r
# Create some plots
bxp <- ggboxplot(iris, x = "Species", y = "Sepal.Length")
vp <- ggviolin(iris, x = "Species", y = "Sepal.Length",
              add = "mean_sd")

# Arrange the plots in one page
# Returns a gtable (grob) object
library(gridExtra)
gt <- arrangeGrob(bxp, vp, ncol = 2)

# Transform to a ggplot and print
```
as_npc

as_ggplot(gt)

---

**as_npc**  
*Convert Character Coordinates into Normalized Parent Coordinates (NPC)*

**Description**

Convert character coordinates to npc units and shift postions to avoid overlaps when grouping is active. If numeric validate npc values.

**Usage**

```r
as_npc(
  value,
  group = 1L,
  step = 0.1,
  margin.npc = 0.05,
  axis = c("xy", "x", "y")
)
```

```r
as_npcx(value, group = 1L, step = 0.1, margin.npc = 0.05)
```

```r
as_npcy(value, group = 1L, step = 0.1, margin.npc = 0.05)
```

**Arguments**

- `value` numeric (in [0-1]) or character vector of coordinates. If character, should be one of `c("right", "left", "bottom", "top", "center", "centre", "middle")`.
- `group` integer ggplot's group id. Used to shift coordinates to avoid overlaps.
- `step` numeric value in [0-1]. The step size for shifting coordinates in npc units. Considered as horizontal step for x-axis and vertical step for y-axis. For y-axis, the step value can be negative to reverse the order of groups.
- `margin.npc` numeric [0-1] The margin added towards the nearest plotting area edge when converting character coordinates into npc.
- `axis` the concerned axis. Should be one of `c("xy", "x", "y")`.

**Details**

The `as_npc()` function is an adaptation from `ggpmisc::compute_npc()`.

**Value**

A numeric vector with values in the range [0-1] representing npc coordinates.
Functions

- `as_npc()`: converts x or y coordinate values into npc. Input values should be numeric or one of the following values `c('right', 'left', 'bottom', 'top', 'center', 'centre', 'middle')`.

- `as_npcx()`: converts x coordinate values into npc. Input values should be numeric or one of the following values `c('right', 'left', 'center', 'centre', 'middle')`. Wrapper around `as_npc(axis = "x")`.

- `as_npcy()`: converts y coordinate values into npc. Input values should be numeric or one of the following values `c('bottom', 'top', 'center', 'centre', 'middle')`. Wrapper around `as_npc(axis = "y")`.

See Also

npc_to_data_coord, get_coord.

Examples

```r
as_npc(c("left", "right"))
as_npc(c("top", "right"))
```

## Description

Change axis scale.

- `xscale`: change x axis scale.
- `yscale`: change y axis scale.

## Usage

```r
xscale(.scale, .format = FALSE)
yscale(.scale, .format = FALSE)
```

## Arguments

- `.scale`: axis scale. Allowed values are one of `c("none", "log2", "log10", "sqrt", "percent", "dollar", "scientific")`; e.g.: `.scale="log2"`.
- `.format`: logical value. If TRUE, axis tick mark labels will be formatted when `.scale = "log2"` or "log10".
Examples

# Basic scatter plots
data(cars)
p <- ggscatter(cars, x = "speed", y = "dist")
p
# Set log scale
p + yscale("log2", .format = TRUE)

Description

Add background image to ggplot2.

Usage

background_image(raster.img)

Arguments

raster.img  
raster object to display, as returned by the function readPNG() [in png package] 
and readJPEG() [in jpeg package].

Author(s)

Alboukadel Kassambara <alboukadel.kassambara@gmail.com>

Examples

## Not run:
install.packages("png")

# Import the image
img.file <- system.file(file.path("images", "background-image.png"),
                        package = "ggpubr")
img <- png::readPNG(img.file)

# Plot with background image
ggplot(iris, aes(Species, Sepal.Length))+
background_image(img)+
geom_boxplot(aes(fill = Species), color = "white")+
fill_palette("jco")

## End(Not run)
bgcolor  

*Change ggplot Panel Background Color*

**Description**

Change ggplot panel background color.

**Usage**

```
bgcolor(color)
```

**Arguments**

- `color`  background color.

**See Also**

`border()`.

**Examples**

```r
# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
p <- ggboxplot(df, x = "dose", y = "len")
p

# Change panel background color
p +
  bgcolor("#BFD5E3") +
  border("#BFD5E3")

```

---

**border**  

*Set ggplot Panel Border Line*

**Description**

Change or set ggplot panel border.

**Usage**

```
border(color = "black", size = 0.8, linetype = NULL)
```
compare_means

Arguments

- **color**: border line color.
- **size**: numeric value specifying border line size.
- **linetype**: line type. An integer (0:8), a name (blank, solid, dashed, dotted, dotdash, long-dash, twodash). See `show_line_types`.

Examples

```r
# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
p <- ggboxplot(df, x = "dose", y = "len")
p

# Add border
p + border()
```

Description

Performs one or multiple mean comparisons.

Usage

```r
compare_means(
  formula,
  data,
  method = "wilcox.test",
  paired = FALSE,
  group.by = NULL,
  ref.group = NULL,
  symnum.args = list(),
  p.adjust.method = "holm",
  ...
)
```

Arguments

- **formula**: a formula of the form `x ~ group` where `x` is a numeric variable giving the data values and `group` is a factor with one or multiple levels giving the corresponding groups. For example, `formula = TP53 ~ cancer_group`.

  It's also possible to perform the test for multiple response variables at the same time. For example, `formula = c(TP53, PTEN) ~ cancer_group`. 

data: a data.frame containing the variables in the formula.

method: the type of test. Default is `wilcox.test`. Allowed values include:

- `t.test` (parametric) and `wilcox.test` (non-parametric). Perform comparison between two groups of samples. If the grouping variable contains more than two levels, then a pairwise comparison is performed.
- `anova` (parametric) and `kruskal.test` (non-parametric). Perform one-way ANOVA test comparing multiple groups.

paired: a logical indicating whether you want a paired test. Used only in `t.test` and in `wilcox.test`.

group.by: a character vector containing the name of grouping variables.

ref.group: a character string specifying the reference group. If specified, for a given grouping variable, each of the group levels will be compared to the reference group (i.e. control group). `ref.group` can also be ".all.". In this case, each of the grouping variable levels is compared to all (i.e. basemean).

symnum.args: a list of arguments to pass to the function `symnum` for symbolic number coding of p-values. For example, `symnum.args <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", ",**", ",*", "ns").

In other words, we use the following convention for symbols indicating statistical significance:

- `ns`: \( p > 0.05 \)
- `*`: \( p \leq 0.05 \)
- `**: \( p \leq 0.01 \)
- `***`: \( p \leq 0.001 \)
- `****`: \( p \leq 0.0001 \)

p.adjust.method: method for adjusting p values (see `p.adjust`). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don’t want to adjust the p value (not recommended), use `p.adjust.method = "none"`. Note that, when the formula contains multiple variables, the p-value adjustment is done independently for each variable.

Value

return a data frame with the following columns:

- `.y.`: the y variable used in the test.
- `group1,group2`: the compared groups in the pairwise tests. Available only when method = "t.test" or method = "wilcox.test".
- `p`: the p-value.

• p.format: the formatted p-value.
• p.signif: the significance level.
• method: the statistical test used to compare groups.

Examples

# Load data
#:::::::::::::::::::::::::::::::::::::::
data("ToothGrowth")
df <- ToothGrowth

# One-sample test
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ 1, df, mu = 0)

# Two-samples unpaired test
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ supp, df)

# Two-samples paired test
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ supp, df, paired = TRUE)

# Compare supp levels after grouping the data by "dose"
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ supp, df, group.by = "dose")

# pairwise comparisons
#:::::::::::::::::::::::::::::::::::::::::
# As dose contains more than two levels =>
# pairwise test is automatically performed.
compare_means(len ~ dose, df)

# Comparison against reference group
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ dose, df, ref.group = "0.5")

# Comparison against all
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ dose, df, ref.group = "all.")

# Anova and kruskal.test
#:::::::::::::::::::::::::::::::::::::::::
compare_means(len ~ dose, df, method = "anova")
compare_means(len ~ dose, df, method = "kruskal.test")
**Description**

Create aes mapping to make programming easy with ggplot2.

**Usage**

```r
create_aes(.list, parse = TRUE)
```

**Arguments**

- `.list` : a list of aesthetic arguments; for example `.list = list(x = "dose", y = "len", color = "dose")`.
- `parse` : logical. If TRUE, parse the input as an expression.

**Examples**

```r
# Simple aes creation
create_aes(list(x = "Sepal.Length", y = "Petal.Length" ))

# Parse an expression
x <- "log2(Sepal.Length)"
y <- "log2(Petal.Length)"
create_aes(list(x = x, y = y ), parse = TRUE)

# Create a ggplot
mapping <- create_aes(list(x = x, y = y ), parse = TRUE)
ggplot(iris, mapping) + geom_point()
```

---

**desc_statby**

**Descriptive statistics by groups**

**Description**

Computes descriptive statistics by groups for a measure variable.

**Usage**

```r
desc_statby(data, measure.var, grps, ci = 0.95)
```

**Arguments**

- `data` : a data frame.
- `measure.var` : the name of a column containing the variable to be summarized.
- `grps` : a character vector containing grouping variables; e.g.: `grps = c("grp1", "grp2")`
- `ci` : the percent range of the confidence interval (default is 0.95).
Value

A data frame containing descriptive statistics, such as:

- **length**: the number of elements in each group
- **min**: minimum
- **max**: maximum
- **median**: median
- **mean**: mean
- **iqr**: interquartile range
- **mad**: median absolute deviation (see ?MAD)
- **sd**: standard deviation of the sample
- **se**: standard error of the mean. It’s calculated as the sample standard deviation divided by the root of the sample size.
- **ci**: confidence interval of the mean
- **range**: the range = max - min
- **cv**: coefficient of variation, sd/mean
- **var**: variance, sd^2

Examples

```r
# Load data
data("ToothGrowth")

# Descriptive statistics
res <- desc_statby(ToothGrowth, measure.var = "len",
                    grps = c("dose", "supp"))
head(res[, 1:10])
```

Description

Differential gene expression analysis results obtained from comparing the RNAseq data of two different cell populations using DESeq2

Usage

```r
data("diff_express")
```
Format

A data frame with 36028 rows and 5 columns.

name  gene names
baseMean  mean expression signal across all samples
log2FoldChange  log2 fold change
padj  Adjusted p-value
detection_call  a numeric vector specifying whether the genes is expressed (value = 1) or not (value = 0).

Examples

data(diff_express)

# Default plot
ggmaplot(diff_express, main = expression("Group 1" %->% "Group 2"),
  fdr = 0.05, fc = 2, size = 0.4,
  palette = c("#B31B21", "#1465AC", "darkgray"),
  genenames = as.vector(diff_express$name),
  legend = "top", top = 20,
  font.label = c("bold", 11),
  font.legend = "bold",
  font.main = "bold",
  ggtheme = ggplot2::theme_minimal())

# Add rectangle around labels
ggmaplot(diff_express, main = expression("Group 1" %->% "Group 2"),
  fdr = 0.05, fc = 2, size = 0.4,
  palette = c("#B31B21", "#1465AC", "darkgray"),
  genenames = as.vector(diff_express$name),
  legend = "top", top = 20,
  font.label = c("bold", 11), label.rectangle = TRUE,
  font.legend = "bold",
  font.main = "bold",
  ggtheme = ggplot2::theme_minimal())

---

Facet

Facet a ggplot into Multiple Panels

Description

Create multi-panel plots of a data set grouped by one or two grouping variables. Wrapper around facet_wrap
Usage

```r
facet(
  p,
  facet.by,
  nrow = NULL,
  ncol = NULL,
  scales = "fixed",
  short.panel.labs = TRUE,
  labeller = "label_value",
  panel.labs = NULL,
  panel.labs.background = list(color = NULL, fill = NULL),
  panel.labs.font = list(face = NULL, color = NULL, size = NULL, angle = NULL),
  panel.labs.font.x = panel.labs.font,
  panel.labs.font.y = panel.labs.font,
  strip.position = "top",
  ...
)
```

Arguments

- **p** a `ggplot`
- **facet.by** character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
- **nrow, ncol** Number of rows and columns in the panel. Used only when the data is faceted by one grouping variable.
- **scales** should axis scales of panels be fixed ("fixed", the default), free ("free"), or free in one dimension ("free_x", "free_y").
- **short.panel.labs** logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.
- **labeller** Character vector. An alternative to the argument `short.panel.labs`. Possible values are one of "label_both" (panel labelled by both grouping variable names and levels) and "label_value" (panel labelled with only grouping levels).
- **panel.labs** a list of one or two character vectors to modify facet panel labels. For example, `panel.labs = list(sex = c("Male", "Female"))` specifies the labels for the "sex" variable. For two grouping variables, you can use for example `panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2"))`.
- **panel.labs.background** a list to customize the background of panel labels. Should contain the combination of the following elements:
  - color, linetype, size: background line color, type and size
  - fill: background fill color.

For example, `panel.labs.background = list(color = "blue", fill = "pink", linetype = "dashed", size = 0.5)`. 
font

panel.labs.font

A list of aesthetics indicating the size (e.g.: 14), the face/style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") and the orientation angle (e.g.: 45) of panel labels.

panel.labs.font.x, panel.labs.font.y

Same as panel.labs.font but for only x and y direction, respectively.

strip.position

Used only in facet_wrap(). By default, the labels are displayed on the top of the plot. Using strip.position it is possible to place the labels on either of the four sides by setting strip.position = c("top", "bottom", "left", "right")

Examples

p <- ggboxplot(ToothGrowth, x = "dose", y = "len", color = "supp")
print(p)

facet(p, facet.by = "supp")

# Customize

facet(p + theme_bw(), facet.by = "supp",
  short.panel.labs = FALSE, # Allow long labels in panels
  panel.labs.background = list(fill = "steelblue", color = "steelblue")
)

Change the Appearance of Titles and Axis Labels

Description

Change the appearance of the main title, subtitle, caption, axis labels and text, as well as the legend title and texts. Wrapper around element_text().

Usage

font(object, size = NULL, color = NULL, face = NULL, family = NULL, ...)

Arguments

object character string specifying the plot components. Allowed values include:
  • "title" for the main title
  • "subtitle" for the plot subtitle
  • "caption" for the plot caption
  • "legend. title" for the legend title
  • "legend.text" for the legend text
  • "x", "xlab", or "x.title" for x axis label
gene_citation 21

- "y", "ylab", or "y.title" for y axis label
- "xy", "xylab", "xy.title" or "axis.title" for both x and y axis labels
- "x.text" for x axis texts (x axis tick labels)
- "y.text" for y axis texts (y axis tick labels)
- "xy.text" or "axis.text" for both x and y axis texts

size numeric value specifying the font size, (e.g.: size = 12).
color character string specifying the font color, (e.g.: color = "red").
face the font face or style. Allowed values include one of "plain", "bold", "italic", "bold.italic", (e.g.: face = "bold.italic").
family the font family.

Examples

# Load data
data("ToothGrowth")

# Basic plot
p <- ggboxplot(ToothGrowth, x = "dose", y = "len", color = "dose",
title = "Box Plot created with ggpubr",
subtitle = "Length by dose",
caption = "Source: ggpubr",
xlab = "Dose (mg)", ylab = "Teeth length")
p

# Change the appearance of titles and labels
p +
  font("title", size = 14, color = "red", face = "bold.italic")+
  font("subtitle", size = 10, color = "orange")+
  font("caption", size = 10, color = "orange")+
  font("xlab", size = 12, color = "blue")+
  font("ylab", size = 12, color = "#993333")+
  font("xy.text", size = 12, color = "gray", face = "bold")

# Change the appearance of legend title and texts
p +
  font("legend.title", color = "blue", face = "bold")+
  font("legend.text", color = "red")

---

gene_citation  Gene Citation Index

Description

Contains the mean citation index of 66 genes obtained by assessing PubMed abstracts and annotations using two key words i) Gene name + b cell differentiation and ii) Gene name + plasma cell differentiation.
gene_expression

Usage

data("gene_expression")

Description

Gene expression data extracted from TCGA using the 'RTCGA' and 'RTCGA.mRNA' R packages. It contains the mRNA expression for 3 genes - GATA3, PTEN and XBP1 - from 3 different datasets: Breast invasive carcinoma (BRCA), Ovarian serous cystadenocarcinoma (OV) and Lung squamous cell carcinoma (LUSC)

Usage

data("gene_expression")
Format

A data frame with 1305 rows and 5 columns.

<table>
<thead>
<tr>
<th>bcr_patient_barcode</th>
<th>sample ID</th>
<th>dataset</th>
<th>canc type</th>
<th>GATA3</th>
<th>GATA3 gene expression</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTEN</td>
<td>PTEN gene expression</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>XBP1</td>
<td>XBP1 gene expression</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Examples

```r
data(gene_expression)

ggboxplot(gene_expression, x = "dataset",
y = c("GATA3", "PTEN", "XBP1"),
    combine = TRUE,
ylab = "Expression",
color = "dataset", palette = "jco")
```

Description

A helper function used by ggpubr functions to execute any geom_* functions in ggplot2. Useful only when you want to call a geom_* function without specifying the arguments to put in aes(). Basic users of ggpubr don’t need this function.

Usage

```r
geom_exec(geomfunc = NULL, data = NULL, position = NULL, ...)
```

Arguments

- `geomfunc` a ggplot2 function (e.g.: geom_point)
- `data` a data frame to be used for mapping
- `position` Position adjustment, either as a string, or the result of a call to a position adjustment function.
- `...` arguments accepted by the function

Value

return a plot if geomfunc!=Null or a list(option, mapping) if geomfunc = NULL.
Examples

```r
## Not run:
ggplot() + geom_exec(geom_point, data = mtcars,
  x = "mpg", y = "wt", size = "cyl", color = "cyl")

## End(Not run)
```

---

**get_breaks**

**Easy Break Creation for Numeric Axes**

**Description**

Creates breaks for numeric axes to be used in the functions `scale_x_continuous()` and `scale_y_continuous()`. Can be used to increase the number of x and y ticks by specifying the option `n`. It's also possible to control axis breaks by specifying a step between ticks. For example, if `by = 5`, a tick mark is shown on every 5.

**Usage**

```r
get_breaks(n = NULL, by = NULL, from = NULL, to = NULL)
```

**Arguments**

- `n` number of breaks.
- `by` number: the step between breaks.
- `from` the starting value of breaks. By default, 0 is used for positive variables.
- `to` the end values of breaks. This corresponds generally to the maximum limit of the axis.

**Value**

a break function

**Examples**

```r
# Generate 5 breaks for a variable x
get_breaks(n = 5)(x = 1:100)

# Generate breaks using an increasing step
get_breaks(by = 10)(x = 1:100)

# Combine with ggplot scale_xx functions
library(ggplot2)

# Create a basic plot
p <- ggscatter(mtcars, x = "wt", y = "mpg")
p
```
# Increase the number of ticks
p +
  scale_x_continuous(breaks = get_breaks(n = 10)) +
  scale_y_continuous(breaks = get_breaks(n = 10))

# Set ticks according to a specific step, starting from 0
p + scale_x_continuous(
  breaks = get_breaks(by = 1.5, from = 0),
  limits = c(0, 6)
) +
  scale_y_continuous(
    breaks = get_breaks(by = 10, from = 0),
    limits = c(0, 40)
  )

---

get_coord

Checks and Returns Data Coordinates from Multiple Input Options

Description

Checks and returns selected coordinates from multiple input options, which can be either data (x-y) coordinates or npc (normalized parent coordinates).

Helper function internally used in ggpubr function to guess the type of coordinates specified by the user. For example, in the function `stat_cor()`, users can specify either the option `label.x` (data coordinates) or `label.x.npc` (npc coordinates); those coordinates are passed to `get_coord()`, which will make some checking and then return a unique coordinates for the label position.

Usage

```r
get_coord(
  group = 1L,
  data.ranges = NULL,
  coord = NULL,
  npc = "left",
  step = 0.1,
  margin.npc = 0.05
)
```

Arguments

- **group**: integer ggplot’s group id. Used to shift coordinates to avoid overlaps.
- **data.ranges**: a numeric vector of length 2 containing the data ranges (minimum and the maximum). Should be specified only when `coord = NULL` and `npc` is specified. Used to convert npc to data coordinates. Considered only when the argument `npc` is specified.
- **coord**: data coordinates (i.e., either x or y coordinates).
get_legend

Extract Legends from a ggplot object

Description

Extract the legend labels from a ggplot object.

Usage

get_legend(p, position = NULL)
get_palette

Arguments

p an object of class ggplot or a list of ggplots. If p is a list, only the first legend is returned.

position character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). To remove the legend use legend = "none".

Value

an object of class gtable.

Examples

# Create a scatter plot
p <- ggscatter(iris, x = "Sepal.Length", y = "Sepal.Width",
  color = "Species", palette = "jco",
  ggtheme = theme_minimal())
p

# Extract the legend. Returns a gtable
leg <- get_legend(p)

# Convert to a ggplot and print
as_ggplot(leg)

get_palette

Generate Color Palettes

Description

Generate a palette of k colors from ggsci palettes, RColorBrewer palettes and custom color palettes. Useful to extend RColorBrewer and ggsci to support more colors.

Usage

get_palette(palette = "default", k)

Arguments

palette Color palette. Allowed values include:
  - Grey color palettes: "grey" or "gray";
  - RColorBrewer palettes, see brewer.pal and details section. Examples of palette names include: "RdBu", "Blues", "Dark2", "Set2", ...;
  - Custom color palettes. For example, palette = c("#00AFBB", "#E7B800", 
    "#FC4E07");
  - ggsci scientific journal palettes, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

k the number of colors to generate.
get_palette

Details

RColorBrewer palettes: To display all available color palettes, type this in R:RColorBrewer::display.brewer.all(). Color palette names include:

- **Sequential palettes**, suited to ordered data that progress from low to high. Palette names include: Blues BuGn BuPu GnuBu Greens Greys OrRd PuBu PuBuGn PuRd Purples RdPu Reds YlGn YlGnBu YlOrBr YlOrRd.
- **Diverging palettes**: Gradient colors. Names include: BrBG PiYG PRGn PuOr RdBu RdGy RdYlBu RdYlGn Spectral.
- **Qualitative palettes**: Best suited to representing nominal or categorical data. Names include: Accent, Dark2, Paired, Pastel1, Pastel2, Set1, Set2, Set3.

Value

Returns a vector of color palettes.

Examples

data("iris")
iris$Species2 <- factor(rep(c(1:10), each = 15))

# Generate a gradient of 10 colors
ggscatter(iris, x = "Sepal.Length", y = "Petal.Length", 
color = "Species2", 
palette = get_palette(c("#00AFBB", "#E7B800", "#FC4E07"), 10))

# Scatter plot with default color palette
ggscatter(iris, x = "Sepal.Length", y = "Petal.Length", 
color = "Species")

# RColorBrewer color palettes
ggscatter(iris, x = "Sepal.Length", y = "Petal.Length", 
color = "Species", palette = get_palette("Dark2", 3))

# ggsci color palettes
ggscatter(iris, x = "Sepal.Length", y = "Petal.Length", 
color = "Species", palette = get_palette("npg", 3))

# Custom color palette
ggscatter(iris, x = "Sepal.Length", y = "Petal.Length", 
color = "Species", 
palette = c("#00AFBB", "#E7B800", "#FC4E07"))

# Or use this
ggscatter(iris, x = "Sepal.Length", y = "Petal.Length", 
color = "Species", 
palette = get_palette(c("#00AFBB", "#FC4E07"), 3))
**ggadd**

Add summary statistics or a geom onto a ggplot

---

**Description**

Add summary statistics or a geometry onto a ggplot.

**Usage**

```r
ggadd(
p,  
add = NULL,  
color = "black",  
fill = "white",  
group = 1,  
width = 1,  
shape = 19,  
size = NULL,  
alpha = 1,  
nseed = 123,  
binwidth = NULL,  
dotsize = size,  
linetype = 1,  
show.legend = NA,  
error.plot = "pointrange",  
ci = 0.95,  
data = NULL,  
position = position_dodge(0.8),  
p_geom = ""  
)
```

**Arguments**

- **p**: a ggplot
- **add**: character vector specifying other plot elements to be added. Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_span", "median_range".
- **color**: point or outline color.
- **fill**: fill color. Used only when `error.plot = "crossbar"`.
- **group**: grouping variable. Allowed values are 1 (for one group) or a character vector specifying the name of the grouping variable. Used only for adding statistical summary per group.
width
numeric value between 0 and 1 specifying bar or box width. Example width = 0.8. Used only when error.plot is one of c("crossbar", "errorbar").

shape
point shape. Allowed values can be displayed using the function `show_point_shapes()`.

size
numeric value in [0-1] specifying point and line size.

alpha
numeric value specifying fill color transparency. Value should be in [0, 1], where 0 is full transparency and 1 is no transparency.

jitter
a numeric value specifying the amount of jittering. Used only when add contains "jitter".

seed
A random seed to make the jitter reproducible. Default is ‘123’. Useful if you need to apply the same jitter twice, e.g., for a point and a corresponding label. The random seed is reset after jittering. If ‘NA’, the seed is initialized with a random value; this makes sure that two subsequent calls start with a different seed. Use NULL to use the current random seed and also avoid resetting (the behaviour of ggplot 2.2.1 and earlier).

binwidth
numeric value specifying bin width. use value between 0 and 1 when you have a strong dense dotplot. For example binwidth = 0.2. Used only when add contains "dotplot".

dotsize
as size but applied only to dotplot.

linetype
line type.

show.legend
logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

error.plot
plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange".

ci
the percent range of the confidence interval (default is 0.95).

data
a data.frame to be displayed. If NULL, the default, the data is inherited from the plot data as specified in the call to `ggplot`.

position
position adjustment, either as a string, or the result of a call to a position adjustment function. Used to adjust position for multiple groups.

p_geom
the geometry of the main plot. Ex: p_geom = "geom_line". If NULL, the geometry is extracted from p. Used only by `ggline()`.

Examples

# Basic violin plot
data("ToothGrowth")
p <- ggviolin(ToothGrowth, x = "dose", y = "len", add = "none")

# Add mean +/- SD and jitter points
p %>% ggadd(c("mean_sd", "jitter"), color = "dose")

# Add box plot
p %>% ggadd(c("boxplot", "jitter"), color = "dose")
ggadjust_pvalue

Adjust p-values Displayed on a GGPlot

Description

Adjust p-values produced by `geom_pwc()` on a ggplot. This is mainly useful when using facet, where p-values are generally computed and adjusted by panel without taking into account the other panels. In this case, one might want to adjust after the p-values of all panels together.

Usage

```r
ggadjust_pvalue(
  p,
  layer = NULL,
  p.adjust.method = "holm",
  label = "p.adj",
  hide.ns = NULL,
  symnum.args = list(),
  output = c("plot", "stat_test")
)
```

Arguments

- **p** 
  a ggplot

- **layer**
  An integer indicating the statistical layer rank in the ggplot (in the order added to the plot).

- **p.adjust.method**
  method for adjusting p values (see `p.adjust`). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Ignored when the specified method is "tukey_hsd" or "games_howell_test" because they come with internal p adjustment method. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don’t want to adjust the p value (not recommended), use `p.adjust.method = "none"`.

- **label**
  character string specifying label. Can be:
  - the column containing the label (e.g.: `label = "p"` or `label = "p.adjust"`), where `p` is the p-value. Other possible values are "p.signif", "p.adj.signif", "p.format", "p.adj.format".
  - an expression that can be formatted by the `glue()` package. For example, when specifying `label = "Wilcoxon, p = \{p\}"`, the expression `{p}` will be replaced by its value.
  - a combination of plotmath expressions and glue expressions. You may want some of the statistical parameter in italic; for example:`label = "Wilcoxon, italic(p)= \{p\}"`
hide.ns can be logical value (TRUE or FALSE) or a character vector ("p.adj" or "p").
symnum.args a list of arguments to pass to the function symnum for symbolic number coding of p-values. For example, symnum.args <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", ",", "ns")).
In other words, we use the following convention for symbols indicating statistical significance:

- ns: p > 0.05
- *: p <= 0.05
- **: p <= 0.01
- ***: p <= 0.001
- ****: p <= 0.0001

output character. Possible values are one of c("plot","stat_test"). Default is "plot".

Examples

# Data preparation
#:::::::::::::::::::::::::::::::::::::::
df <- ToothGrowth
df$dose <- as.factor(df$dose)
# Add a random grouping variable
df$group <- factor(rep(c("grp1", "grp2"), 30))
head(df, 3)

# Boxplot: Two groups by panel
#:::::::::::::::::::::::::::::::::::::::
# Create a box plot
bxp <- ggboxplot(
  df, x = "supp", y = "len", fill = "#00AFBB",
  facet.by = "dose"
)
# Make facet and add p-values
bxp <- bxp + geom_pwc(method = "t_test")
bxp
# Adjust all p-values together after
ggadjust_pvalue(
  bxp, p.adjust.method = "bonferroni",
  label = "(p.adj.format)(p.adj.signif)", hide.ns = TRUE
)

# Boxplot: Three groups by panel
#:::::::::::::::::::::::::::::::::::::::
# Create a box plot
bxp <- ggboxplot(
  df, x = "dose", y = "len", fill = "#00AFBB",
  facet.by = "supp"
)
# Make facet and add p-values
bxp <- bxp + geom_pwc(method = "t_test")
bxp

# Adjust all p-values together after
ggadjust_pvalue(
  bxp, p.adjust.method = "bonferroni",
  label = "\{(p.adj.format)(p.adj.signif)\"
)

---

### ggarrange

#### Arrange Multiple ggplots

Arrange multiple ggplots on the same page. Wrapper around `plot_grid()`. Can arrange multiple ggplots over multiple pages, compared to the standard `plot_grid()`. Can also create a common unique legend for multiple plots.

#### Usage

```r
ggarrange(
  ...
)
```

#### Arguments

- **...**: list of plots to be arranged into the grid. The plots can be either ggplot2 plot objects or arbitrary gtables.
- **plotlist**: (optional) list of plots to display.
- **ncol**: (optional) number of columns in the plot grid.
- **nrow**: (optional) number of rows in the plot grid.
- **labels**: (optional) list of labels to be added to the plots. You can also set labels="AUTO" to auto-generate upper-case labels or labels="auto" to auto-generate lower-case labels.
label.x (optional) Single value or vector of x positions for plot labels, relative to each subplot. Defaults to 0 for all labels. (Each label is placed all the way to the left of each plot.)

label.y (optional) Single value or vector of y positions for plot labels, relative to each subplot. Defaults to 1 for all labels. (Each label is placed all the way to the top of each plot.)

hjust Adjusts the horizontal position of each label. More negative values move the label further to the right on the plot canvas. Can be a single value (applied to all labels) or a vector of values (one for each label). Default is -0.5.

vjust Adjusts the vertical position of each label. More positive values move the label further down on the plot canvas. Can be a single value (applied to all labels) or a vector of values (one for each label). Default is 1.5.

font.label a list of arguments for customizing labels. Allowed values are the combination of the following elements: size (e.g.: 14), face (e.g.: "plain", "bold", "italic", "bold.italic"), color (e.g.: "red") and family. For example font.label = list(size = 14, face = "bold", color = "red").

align (optional) Specifies whether graphs in the grid should be horizontally ("h") or vertically ("v") aligned. Options are "none" (default), "hv" (align in both directions), "h", and "v".

widths (optional) numerical vector of relative columns widths. For example, in a two-column grid, widths = c(2, 1) would make the first column twice as wide as the second column.

heights same as widths but for column heights.

legend character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). To remove the legend use legend = "none".

common.legend logical value. Default is FALSE. If TRUE, a common unique legend will be created for arranged plots.

legend.grob a legend grob as returned by the function get_legend(). If provided, it will be used as the common legend.

Value
return an object of class ggarrange, which is a ggplot or a list of ggplot.

Author(s)
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See Also
annotate_figure()

Examples
data("ToothGrowth")
df <- ToothGrowth
df$dose <- as.factor(df$dose)

# Create some plots
# ::::::::::::::::::::::::::::::::::::::::::::::::::
# Box plot
bxp <- ggboxplot(df, x = "dose", y = "len",
                  color = "dose", palette = "jco")
# Dot plot
dp <- ggdotplot(df, x = "dose", y = "len",
                 color = "dose", palette = "jco")
# Density plot
dens <- ggdensity(df, x = "len", fill = "dose", palette = "jco")

# Arrange
# ::::::::::::::::::::::::::::::::::::::::::::::::::
ggarrange(bxp, dp, dens, ncol = 2, nrow = 2)
# Use a common legend for multiple plots
ggarrange(bxp, dp, common.legend = TRUE)

---

ggballoonplot | Balloon plot

Description
Plot a graphical matrix where each cell contains a dot whose size reflects the relative magnitude of the corresponding component. Useful to visualize contingency table formed by two categorical variables.

Usage
ggballoonplot(
  data,
  x = NULL,
  y = NULL,
  size = "value",
  facet.by = NULL,
  size.range = c(1, 10),
  shape = 21,
  color = "black",
  fill = "gray",
  show.label = FALSE,
  font.label = list(size = 12, color = "black"),
  rotate.x.text = TRUE,
  ggtheme = theme_minimal(),
  ...
)
Arguments

data a data frame. Can be:

- a standard contingency table formed by two categorical variables: a data frame with row names and column names. The categories of the first variable are columns and the categories of the second variable are rows.
- a stretched contingency table: a data frame containing at least three columns corresponding, respectively, to (1) the categories of the first variable, (2) the categories of the second variable, (3) the frequency value. In this case, you should specify the argument x and y in the function ggballoonplot().

x, y the column names specifying, respectively, the first and the second variable forming the contingency table. Required only when the data is a stretched contingency table.

size point size. By default, the points size reflects the relative magnitude of the value of the corresponding cell (size = "value"). Can be also numeric (size = 4).

facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

size.range a numeric vector of length 2 that specifies the minimum and maximum size of the plotting symbol. Default values are size.range = c(1, 10).

shape points shape. The default value is 21. Alternative values include 22, 23, 24, 25.

color point border line color.

fill point fill color. Default is "lightgray". Considered only for points 21 to 25.

show.label logical. If TRUE, show the data cell values as point labels.

font.label a vector of length 3 indicating respectively the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of point labels. For example font.label = c(14, "bold", "red"). To specify only the size and the style, use font.label = c(14, "plain").

rotate.x.text logical. If TRUE (default), rotate the x axis text.

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments passed to the function ggpar

Examples

# Define color palette
my_cols <- c("#0D0887FF", "#6A00A8FF", "#B12A90FF", "#E16462FF", "#F0F921FF")

# Standard contingency table
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::
# Read a contingency table: housetasks
# Repartition of 13 housetasks in the couple
data <- read.delim("..."


```r
# Basic balloon plot
ggballoonplot(data)

# Change color and fill
ggballoonplot(data, color = "#0073C2FF", fill = "#0073C2FF")

# Change color according to the value of table cells

ggballoonplot(data, fill = "value")+
scale_fill_gradientn(colors = my_cols)

# Change the plotting symbol shape

ggballoonplot(data, fill = "value", shape = 23)+
gradient_fill(c("blue", "white", "red"))

# Set points size to 8, but change fill color by values
# Sow labels

ggballoonplot(data, fill = "value", color = "lightgray",
size = 10, show.label = TRUE)+
gradient_fill(c("blue", "white", "red"))

# Stretched contingency table
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::

# Create an Example Data Frame Containing Car x Color data
carnames <- c("bmw","renault","mercedes","seat")
carcolors <- c("red","white","silver","green")
datavals <- round(rnorm(16, mean=100, sd=60),1)
car_data <- data.frame(Car = rep(carnames,4),
                       Color = rep(carcolors, c(4,4,4,4) ),
                       Value=datavals )

car_data

ggballoonplot(car_data, x = "Car", y = "Color",
size = "Value", fill = "Value")+
scale_fill_gradientn(colors = my_cols) +
guides(size = FALSE)

# Grouped frequency table
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::

data("Titanic")
dframe <- as.data.frame(Titanic)
head(dframe)
ggballoonplot(
  dframe, x = "Class", y = "Sex",
size = "Freq", fill = "Freq")
```

size = "Freq", fill = "Freq",
facet.by = c("Survived", "Age"),
ggtheme = theme_bw()
)+
scale_fill_gradientn(colors = my_cols)

# Hair and Eye Color of Statistics Students
data(HairEyeColor)
ggballoonplot( as.data.frame(HairEyeColor),
    x = "Hair", y = "Eye", size = "Freq",
ggtheme = theme_gray()) %>%
    facet("Sex")

---

**ggbarplot**

**Bar plot**

**Description**

Create a bar plot.

**Usage**

```r
ggbarplot(
data, 
x, 
y, 
combine = FALSE, 
merge = FALSE, 
color = "black", 
fill = "white", 
palette = NULL, 
size = NULL, 
width = NULL, 
title = NULL, 
xlab = NULL, 
ylab = NULL, 
facet.by = NULL, 
panel.labs = NULL, 
short.panel.labs = TRUE, 
select = NULL, 
remove = NULL, 
order = NULL, 
add = "none", 
add.params = list(), 
error.plot = "errorbar", 
label = FALSE,
```
Arguments

data a data frame
x, y x and y variables for drawing.
combine logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.

color, fill outline and fill colors.
palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggscale R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
size Numeric value (e.g.: size = 1). change the size of points and outlines.
width numeric value between 0 and 1 specifying box width.
title plot main title.
xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

panel.labs a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2")) .
short.panel.labs
logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

select
character vector specifying which items to display.

remove
character vector specifying which items to remove from the plot.

order
character vector specifying the order of items.

add
character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_q1q3", "median_range"; see ?desc.statby for more details.

add.params
parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

error.plot
plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where "}" = sd, se, ....

label
specify whether to add labels on the bar plot. Allowed values are:
• logical value: If TRUE, y values is added as labels on the bar plot
• character vector: Used as text labels; must be the same length as y.

lab.col, lab.size
text color and size for labels.

lab.pos
character specifying the position for labels. Allowed values are "out" (for outside) or "in" (for inside). Ignored when lab.vjust != NULL.

lab.vjust
numeric, vertical justification of labels. Provide negative value (e.g.: -0.4) to put labels outside the bars or positive value to put labels inside (e.g.: 2).

lab.hjust
numeric, horizontal justification of labels.

lab.nb.digits
integer indicating the number of decimal places (round) to be used.

sort.val
a string specifying whether the value should be sorted. Allowed values are "none" (no sorting), "asc" (for ascending) or "desc" (for descending).

sort.by.groups
logical value. If TRUE the data are sorted by groups. Used only when sort.val != "none".

top
a numeric value specifying the number of top elements to be shown.

position
Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

ggtheme
function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to be passed to ggpar().
**Details**

The plot can be easily customized using the function `ggpar()`. Read `?ggpar` for changing:

- main title and axis labels: `main`, `xlab`, `ylab`
- axis limits: `xlim`, `ylim` (e.g.: `ylim = c(0, 30)`)
- axis scales: `xscale`, `yscale` (e.g.: `yscale = "log2"`)
- color palettes: `palette = "Dark2"` or `palette = c("gray", "blue", "red")`
- legend title, labels and position: `legend = "right"`
- plot orientation: `orientation = c("vertical", "horizontal", "reverse")`

**See Also**

`ggpar`, `ggline`

**Examples**

```r
# Data
df <- data.frame(dose=c("D0.5", "D1", "D2"),
  len=c(4.2, 10, 29.5))
print(df)

# Basic plot with label outsite
# +++++++++++++++++++++++++++
ggbarplot(df, x = "dose", y = "len",
  label = TRUE, label.pos = "out")

# Change width
ggbarplot(df, x = "dose", y = "len", width = 0.5)

# Change the plot orientation: horizontal
ggbarplot(df, "dose", "len", orientation = "horiz")

# Change the default order of items
ggbarplot(df, "dose", "len",
  order = c("D2", "D1", "D0.5"))

# Change colors
# +++++++++++++++++++++++++++
# add labels inside bars
ggbarplot(df, "dose", "len",
  fill = "steelblue", color = "steelblue",
  label = TRUE, lab.pos = "in", lab.col = "white")

# Change colors by groups: dose
# Use custom color palette
ggbarplot(df, "dose", "len", color = "dose",
  palette = c("#00AFBB", "#E7B800", "#FC4E07"))
```
# Change fill and outline colors by groups
ggbarplot(df, "dose", "len",
    fill = "dose", color = "dose",
    palette = c("#00AFBB", "#E7B800", "#FC4E07"))

# Plot with multiple groups
# +++++++++++++++++++++
# Create some data
df2 <- data.frame(supp=rep(c("VC", "OJ"), each=3),
    dose=rep(c("D0.5", "D1", "D2"),2),
    len=c(6.8, 15, 33, 4.2, 10, 29.5))
print(df2)

# Plot "len" by "dose" and change color by a second group: "supp"
# Add labels inside bars
hhbarplot(df2, "dose", "len",
    fill = "supp", color = "supp", palette = "Paired",
    label = TRUE, lab.col = "white", lab.pos = "in")

# Change position: Interleaved (dodged) bar plot
hhbarplot(df2, "dose", "len",
    fill = "supp", color = "supp", palette = "Paired",
    label = TRUE,
    position = position_dodge(0.9))

# Add points and errors
# ++++++++++++++++++++++++++
# Data: ToothGrowth data set we'll be used.
df3 <- ToothGrowth
head(df3, 10)

# It can be seen that for each group we have
# different values
hhbarplot(df3, x = "dose", y = "len")

# Visualize the mean of each group
hhbarplot(df3, x = "dose", y = "len",
    add = "mean")

# Add error bars: mean_se
# (other values include: mean_sd, mean_ci, median_iqr, ....)
# Add labels
hhbarplot(df3, x = "dose", y = "len",
    add = "mean_se", label = TRUE, lab.vjust = -1.6)

# Use only "upper_errorbar"
hhbarplot(df3, x = "dose", y = "len",
    add = "mean_se", error.plot = "upper_errorbar")
# Change error.plot to "pointrange"
ggbarplot(df3, x = "dose", y = "len",
    add = "mean_se", error.plot = "pointrange")

# Add jitter points and errors (mean_se)
ggbarplot(df3, x = "dose", y = "len",
    add = c("mean_se", "jitter"))

# Add dot and errors (mean_se)
ggbarplot(df3, x = "dose", y = "len",
    add = c("mean_se", "dotplot"))

# Multiple groups with error bars and jitter point
ggbarplot(df3, x = "dose", y = "len", color = "supp",
    add = "mean_se", palette = c("#00AFBB", "#E7B800"),
    position = position_dodge())

---

**ggboxplot**  
*Box plot*

**Description**

Create a box plot with points. Box plots display a group of numerical data through their quartiles.

**Usage**

```r
ggboxplot(
    data,
    x,
    y,
    combine = FALSE,
    merge = FALSE,
    color = "black",
    fill = "white",
    palette = NULL,
    title = NULL,
    xlab = NULL,
    ylab = NULL,
    bxp.errorbar = FALSE,
    bxp.errorbar.width = 0.4,
    facet.by = NULL,
    panel.labs = NULL,
    short.panel.labs = TRUE,
    linetype = "solid",
    size = NULL,
    width = 0.7,
    ...)```
```
notch = FALSE,
outlier.shape = 19,
select = NULL,
remove = NULL,
order = NULL,
add = "none",
add.params = list(),
error.plot = "pointrange",
label = NULL,
font.label = list(size = 11, color = "black"),
label.select = NULL,
repel = FALSE,
label.rectangle = FALSE,
ggtheme = theme_pubr(),
...
)

Arguments

data a data frame
x character string containing the name of x variable.
y character vector containing one or more variables to plot
combine logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
color outline color.
fill fill color.
palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggscl R package, e.g.: "hpg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
title plot main title.
xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
bxp.errorbar logical value. If TRUE, shows error bars of box plots.
bxp.errorbar.width numeric value specifying the width of box plot error bars. Default is 0.4.
facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
panel.labs  a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2")).

short.panel.labs  logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

linetype  line types.

size  Numeric value (e.g.: size = 1). change the size of points and outlines.

width  numeric value between 0 and 1 specifying box width.

notch  If FALSE (default) make a standard box plot. If TRUE, make a notched box plot. Notches are used to compare groups; if the notches of two boxes do not overlap, this suggests that the medians are significantly different.

outlier.shape  point shape of outlier. Default is 19. To hide outlier, specify outlier.shape = NA. When jitter is added, then outliers will be automatically hidden.

select  character vector specifying which items to display.

remove  character vector specifying which items to remove from the plot.

order  character vector specifying the order of items.

add  character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see ?desc_statby for more details.

add.params  parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

error.plot  plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where "/" = sd, se, ....

label  the name of the column containing point labels. Can be also a character vector with length = nrow(data).

font.label  a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color = "red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").

label.select  can be of two formats:

- a character vector specifying some labels to show.
- a list containing one or the combination of the following components:
  - top.up and top.down: to display the labels of the top up/down points. For example, label.select = list(top.up = 10, top.down = 4).
- criteria: to filter, for example, by x and y variables values, use this:
  `label.select = list(criteria = "y > 2 & y < 5 & x %in% c('A', 'B')").

repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.

label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ...

... other arguments to be passed to `geom_boxplot`, `ggpar` and `facet`.

Details

The plot can be easily customized using the function `ggpar()`. Read ?ggpar for changing:

- main title and axis labels: `main`, `xlab`, `ylab`
- axis limits: `xlim`, `ylim` (e.g.: `ylim = c(0, 30)`)  
- axis scales: `xscale`, `yscale` (e.g.: `yscale = "log2"`) 
- color palettes: `palette = "Dark2"` or `palette = c("gray", "blue", "red")` 
- legend title, labels and position: `legend = "right"` 
- plot orientation: `orientation = c("vertical", "horizontal", "reverse")`

Suggestions for the argument "add"

Suggested values are one of c("dotplot", "jitter").

See Also

`ggpar`, `ggviolin`, `ggdotplot` and `ggstripchart`.

Examples

```r
# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
# +++++++++++++++++++++++++++
# width: change box plots width
ggboxplot(df, x = "dose", y = "len", width = 0.8)

# Change orientation: horizontal
ggboxplot(df, "dose", "len", orientation = "horizontal")

# Notched box plot
ggboxplot(df, x = "dose", y = "len",
notch = TRUE)
```
# Add dots
# +++++++++++++++++++++++++++
ggboxplot(df, x = "dose", y = "len",
         add = "dotplot")

# Add jitter points and change the shape by groups
ggboxplot(df, x = "dose", y = "len",
         add = "jitter", shape = "dose")

# Select and order items
# ++++++++++++++++++++++++++++++
# Select which items to display: "0.5" and "2"
ggboxplot(df, "dose", "len",
          select = c("0.5", "2"))

# Change the default order of items
ggboxplot(df, "dose", "len",
          order = c("2", "1", "0.5"))

# Change colors
# +++++++++++++++++++++++++++
# Change outline and fill colors
ggboxplot(df, "dose", "len",
          color = "black", fill = "gray")

# Change outline colors by groups: dose
# Use custom color palette
# Add jitter points and change the shape by groups
ggboxplot(df, "dose", "len",
          color = "dose", palette = c("#00AFBB", "#E7B800", "#FC4E07"),
          add = "jitter", shape = "dose")

# Change fill color by groups: dose
ggboxplot(df, "dose", "len",
          fill = "dose", palette = c("#00AFBB", "#E7B800", "#FC4E07"))

# Box plot with multiple groups
# +++++++++++++++++++++++++++++
# fill or color box plot by a second group : "supp"
ggboxplot(df, "dose", "len", color = "supp",
          palette = c("#00AFBB", "#E7B800"))
Description

Create a density plot.

Usage

ggdensity(
  data,
  x,
  y = "density",
  combine = FALSE,
  merge = FALSE,
  color = "black",
  fill = NA,
  palette = NULL,
  size = NULL,
  linetype = "solid",
  alpha = 0.5,
  title = NULL,
  xlab = NULL,
  ylab = NULL,
  facet.by = NULL,
  panel.labs = NULL,
  short.panel.labs = TRUE,
  add = c("none", "mean", "median"),
  add.params = list(linetype = "dashed"),
  rug = FALSE,
  label = NULL,
  font.label = list(size = 11, color = "black"),
  label.select = NULL,
  repel = FALSE,
  label.rectangle = FALSE,
  ggtheme = theme_pubr(),
  ...
)

Arguments

data a data frame
x variable to be drawn.
y one of "density" or "count".
combine logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
color, fill
density line color and fill color.

palette
the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

size
Numeric value (e.g.: size = 1). change the size of points and outlines.

linetype
line type. See show_line_types.

alpha
numeric value specifying fill color transparency. Value should be in [0, 1], where 0 is full transparency and 1 is no transparency.

title
plot main title.

xlab
character vector specifying x axis labels. Use xlab = FALSE to hide xlab.

ylab
character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

facet.by
character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

panel.labs
a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2").

short.panel.labs
logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

add
allowed values are one of "mean" or "median" (for adding mean or median line, respectively).

add.params
parameters (color, size, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

rug
logical value. If TRUE, add marginal rug.

label
the name of the column containing point labels. Can be also a character vector with length = nrow(data).

font.label
a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color ="red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").

label.select
can be of two formats:

• a character vector specifying some labels to show.
• a list containing one or the combination of the following components:
  - top. up and top. down: to display the labels of the top up/down points. For example, label.select = list(top.up = 10, top.down = 4).
  - criteria: to filter, for example, by x and y variables values, use this:
    label.select = list(criteria = "'y' > 2 & 'y' < 5 & 'x' %in% c('A', 'B'))").
repel

repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.

label.rectangle

label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.

ggtheme

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to geom_density and ggpar.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

• main title and axis labels: main, xlab, ylab
• axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
• axis scales: xscale,yscale (e.g.: yscale = "log2")
• color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
• legend title, labels and position: legend = "right"
• plot orientation : orientation = c("vertical", "horizontal", "reverse")

See Also

gghistogram and ggpar.

Examples

# Create some data format
set.seed(1234)
wdata = data.frame(
  sex = factor(rep(c("F", "M"), each=200)),
  weight = c(rnorm(200, 55), rnorm(200, 58)))

head(wdata, 4)

# Basic density plot
# Add mean line and marginal rug
ggdensity(wdata, x = "weight", fill = "lightgray",
      add = "mean", rug = TRUE)

# Change outline colors by groups ("sex")
# Use custom palette
ggdensity(wdata, x = "weight",
      add = "mean", rug = TRUE,
      color = "sex", palette = c("#00AFBB", "#E7B800"))

# Change outline and fill colors by groups ("sex")
# Use custom palette
ggdensity(wdata, x = "weight",
      add = "mean", rug = TRUE,
      color = "sex", palette = c("#00AFBB", "#E7B800"))
ggdonutchart

Donut chart

Description
Create a donut chart.

Usage

```
ggdonutchart(
  data,
  x, 
  label = x,  
  lab.pos = c("out", "in"),
  lab.adjust = 0,
  lab.font = c(4, "plain", "black"),
  font.family = "",
  color = "black",
  fill = "white",
  palette = NULL,
  size = NULL,
  ggtheme = theme_pubr(),
  ...
)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>a data frame</td>
</tr>
<tr>
<td>x</td>
<td>variable containing values for drawing.</td>
</tr>
<tr>
<td>label</td>
<td>variable specifying the label of each slice.</td>
</tr>
<tr>
<td>lab.pos</td>
<td>character specifying the position for labels. Allowed values are &quot;out&quot; (for outside) or &quot;in&quot; (for inside).</td>
</tr>
<tr>
<td>lab.adjust</td>
<td>numeric value, used to adjust label position when lab.pos = &quot;in&quot;. Increase or decrease this value to see the effect.</td>
</tr>
<tr>
<td>lab.font</td>
<td>a vector of length 3 indicating respectively the size (e.g.: 14), the style (e.g.: &quot;plain&quot;, &quot;bold&quot;, &quot;italic&quot;, &quot;bold.italic&quot;) and the color (e.g.: &quot;red&quot;) of label font. For example <code>lab.font = c(4, &quot;bold&quot;, &quot;red&quot;)</code>.</td>
</tr>
<tr>
<td>font.family</td>
<td>character vector specifying font family.</td>
</tr>
<tr>
<td>color, fill</td>
<td>outline and fill colors.</td>
</tr>
</tbody>
</table>
The color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "hpg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

Numeric value (e.g.: size = 1). change the size of points and outlines.

function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

other arguments to be passed to be passed to ggpar().

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation : orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar, ggpie

Examples

# Data: Create some data
# ++++++++++++++++++++++++++++++++
df <- data.frame(
  group = c("Male", "Female", "Child"),
  value = c(25, 25, 50))

head(df)

# Basic pie charts
# ++++++++++++++++++++++++++++++++
ggdonutchart(df, "value", label = "group")

# Change color
# ++++++++++++++++++++++++++++++++
# Change fill color by group
# set line color to white
# Use custom color palette
ggdonutchart(df, "value", label = "group",
    fill = "group", color = "white",
    palette = c("#00AFBB", "#E7B800", "#FC4E07")
)

# Change label
# ++++++++++++++++++++++++++++++++
# Show group names and value as labels
labs <- paste0(df$group, " (", df$value, ")")
ggdonutchart(df, "value", label = labs,
    fill = "group", color = "white",
    palette = c("#00AFBB", "#E7B800", "#FC4E07"))

# Change the position and font color of labels
ggdonutchart(df, "value", label = labs,
    lab.pos = "in", lab.font = "white",
    fill = "group", color = "white",
    palette = c("#00AFBB", "#E7B800", "#FC4E07"))

---

ggdotchart

**Cleveland’s Dot Plots**

**Description**

Draw a Cleveland dot plot.

**Usage**

```r
ggdotchart(
    data,
    x,
    y,
    group = NULL,
    combine = FALSE,
    color = "black",
    palette = NULL,
    shape = 19,
    size = NULL,
    dot.size = size,
    sorting = c("ascending", "descending", "none"),
    add = c("none", "segment"),
    add.params = list(),
    x.text.col = TRUE,
```
rotate = FALSE,
title = NULL,
xlab = NULL,
ylab = NULL,
facet.by = NULL,
panel.labs = NULL,
short.panel.labs = TRUE,
select = NULL,
remove = NULL,
order = NULL,
label = NULL,
font.label = list(size = 11, color = "black"),
label.select = NULL,
repel = FALSE,
label.rectangle = FALSE,
position = "identity",
ggtheme = theme_pubr(),
...)

theme_cleveland(rotate = TRUE)

Arguments

data | a data frame
x, y | x and y variables for drawing.
group | an optional column name indicating how the elements of x are grouped.
combine | logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
color, size | points color and size.
palette | the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
shape | point shape. See show_point_shapes.
dot.size | numeric value specifying the dot size.
sorting | a character vector for sorting into ascending or descending order. Allowed values are one of "descending", "ascending" and "none". Partial match are allowed (e.g. sorting = "desc" or "asc"). Default is "descending".
add | character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see ?desc_statby for more details.
add.params parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").
x.text.col logical. If TRUE (default), x axis texts are colored by groups.
rotate logical value. If TRUE, rotate the graph by setting the plot orientation to horizontal.
title plot main title.
xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
panel.labs a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2" )).
short.panel.labs logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.
select character vector specifying which items to display.
remove character vector specifying which items to remove from the plot.
order character vector specifying the order of items.
label the name of the column containing point labels.
font.label a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color = "red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").
label.select can be of two formats:
• a character vector specifying some labels to show.
• a list containing one or the combination of the following components:
  – top. up and top. down: to display the labels of the top up/down points.
    For example, label.select = list(top. up = 10, top. down = 4).
  – criteria: to filter, for example, by x and y variabes values, use this:
    label.select = list(criteria = "y > 2 & y < 5 & x %in% c('A', 'B')").
repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.
label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.
position Position adjustment, either as a string, or the result of a call to a position adjustment function.
ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....
... other arguments to be passed to geom_point and ggpar.
**Details**

The plot can be easily customized using the function `ggpar()`. Read ?ggpar for changing:

- main title and axis labels: `main`, `xlab`, `ylab`
- axis limits: `xlim`, `ylim` (e.g.: `ylim = c(0, 30)`)
- axis scales: `xscale`, `yscale` (e.g.: `yscale = 'log2'`)
- color palettes: `palette = 'Dark2'` or `palette = c('gray', 'blue', 'red')`
- legend title, labels and position: `legend = 'right'`
- plot orientation: `orientation = c('vertical', 'horizontal', 'reverse')`

**See Also**

`ggpar`

**Examples**

```r
# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)
df$name <- rownames(df)
head(df[, c("wt", "mpg", "cyl")], 3)

# Basic plot
ggdotchart(df, x = "name", y = "mpg",
gtheme = theme_bw())

# Change colors by group cyl
ggdotchart(df, x = "name", y = "mpg",
group = "cyl", color = "cyl",
palette = c("#999999", '#E69F00', '#56B4E9'),
rotate = TRUE,
sorting = "descending",
gtheme = theme_bw(),
y.text.col = TRUE)

# Plot with multiple groups
# +++++++++++++++++++++
# Create some data
df2 <- data.frame(supp=rep(c("VC", "OJ"), each=3),
dose=rep(c("D0.5", "D1", "D2"),2),
len=c(6.8, 15, 33, 4.2, 10, 29.5))
print(df2)

ggdotchart(df2, x = "dose", y = "len",
color = "supp", size = 3,
add = "segment",
add.params = list(color = "lightgray", size = 1.5),
position = position_dodge(0.3),
position = position_dodge(0.3),
```
ggdotplot

```
palette = "jco",
ggtheme = theme_pubclear()```

---

**Description**

Create a dot plot.

**Usage**

```r
ggdotplot(
  data,
  x,
  y,
  combine = FALSE,
  merge = FALSE,
  color = "black",
  fill = "lightgray",
  palette = NULL,
  title = NULL,
  xlab = NULL,
  ylab = NULL,
  facet.by = NULL,
  panel.labs = NULL,
  short.panel.labs = TRUE,
  size = NULL,
  binwidth = NULL,
  select = NULL,
  remove = NULL,
  order = NULL,
  add = "mean_se",
  add.params = list(),
  error.plot = "pointrange",
  label = NULL,
  font.label = list(size = 11, color = "black"),
  label.select = NULL,
  repel = FALSE,
  label.rectangle = FALSE,
  ggtheme = theme_pubr(),
  ...
)
```
ggdotplot

Arguments

data a data frame

x character string containing the name of x variable.

y character vector containing one or more variables to plot

combine logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.

merge logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.

color outline color.

fill fill color.

palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggscc R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

title plot main title.

xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.

ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

panel.labs a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2") ).

short.panel.labs logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

size Numeric value (e.g.: size = 1). change the size of points and outlines.

binwidth numeric value specifying bin width. use value between 0 and 1 when you have a strong dense dotplot. For example binwidth = 0.2.

select character vector specifying which items to display.

remove character vector specifying which items to remove from the plot.

order character vector specifying the order of items.

add character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "box-plot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_q1q2", "median_range": see ?desc_statby for more details.
add.params parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

error.plot plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_" or "med_" where "*" = sd, se, ....

label the name of the column containing point labels. Can be also a character vector with length = nrow(data).

font.label a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color = "red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").

label.select can be of two formats:
- a character vector specifying some labels to show.
- a list containing one or the combination of the following components:
  - top.up and top.down: to display the labels of the top up/down points. For example, label.select = list(top.up = 10, top.down = 4).
  - criteria: to filter, for example, by x and y variables values, use this: label.select = list(criteria = "y > 2 & y < 5 & x %in% c('A', 'B')").

repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.

label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to geom_dotplot, ggpar and facet.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:
- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation: orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar, ggviolin, ggbboxplot and ggstripchart.
Examples

```r
# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot with summary statistics: mean_sd
# +++++++++++++++++++++++++++
ggdotplot(df, x = "dose", y = "len",
         add = "mean_sd")

# Change error.plot to "crossbar"
ggdotplot(df, x = "dose", y = "len",
         add = "mean_sd", add.params = list(width = 0.5),
         error.plot = "crossbar")

# Add box plot
ggdotplot(df, x = "dose", y = "len",
         add = "boxplot")

# Add violin + mean_sd
ggdotplot(df, x = "dose", y = "len",
         add = c("violin", "mean_sd"))

# Change colors
# +++++++++++++++++++++++++++
# Change fill and outline colors by groups: dose
# Use custom color palette
ggdotplot(df, "dose", "len",
         add = "boxplot",
         color = "dose", fill = "dose",
         palette = c("#00AFBB", "#E7B800", "#FC4E07"))

# Plot with multiple groups
# +++++++++++++++++++++
# Change color by a second group: "supp"
ggdotplot(df, "dose", "len", fill = "supp", color = "supp",
         palette = c("#00AFBB", "#E7B800"))
```

ggecdf

Empirical cumulative density function

Description

Empirical Cumulative Density Function (ECDF).
Usage

ggecdf(
data, x,
combine = FALSE, merge = FALSE, color = "black", palette = NULL,
size = NULL, linetype = "solid", title = NULL, xlab = NULL, ylab = NULL,
facet.by = NULL, panel.labs = NULL, short.panel.labs = TRUE, ggtheme = theme_pubr(), ...
)

Arguments

data
  a data frame
x
  variable to be drawn.
combine
  logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge
  logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
color
  line and point color.
palette
  the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
size
  line and point size.
linetype
  line type. See show_line_types.
title
  plot main title.
xlab
  character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab
  character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
facet.by
  character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
panel.labs  a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2")).

short.panel.labs  logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

ggtheme  function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to stat_ecdf and ggpar.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

• main title and axis labels: main, xlab, ylab
• axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
• axis scales: xscale, yscale (e.g.: yscale = "log2")
• color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
• legend title, labels and position: legend = "right"
• plot orientation: orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar

Examples

# Create some data format
set.seed(1234)
wdata = data.frame(
    sex = factor(rep(c("F", "M"), each=200)),
    weight = c(rnorm(200, 55), rnorm(200, 58)))

head(wdata, 4)

# Basic ECDF plot
ggecdf(wdata, x = "weight")

# Change colors and linetype by groups ("sex")
# Use custom palette

# ggecdf(wdata, x = "weight",
# color = "sex", linetype = "sex",
# palette = c("#00AFBB", "#E7B880"))
ggerrorplot

Visualizing Error

Description

Visualizing error.

Usage

```r
ggerrorplot(
  data,
  x,
  y,
  desc_stat = "mean_se",
  numeric.x.axis = FALSE,
  combine = FALSE,
  merge = FALSE,
  color = "black",
  fill = "white",
  palette = NULL,
  size = NULL,
  width = NULL,
  title = NULL,
  xlab = NULL,
  ylab = NULL,
  facet.by = NULL,
  panel.labs = NULL,
  short.panel.labs = TRUE,
  select = NULL,
  remove = NULL,
  order = NULL,
  add = "none",
  add.params = list(),
  error.plot = "pointrange",
  ci = 0.95,
  position = position_dodge(),
  ggtheme = theme_pubr(),
  ...
)
```

Arguments

data a data frame
x, y x and y variables for drawing.
desc_stat descriptive statistics to be used for visualizing errors. Default value is "mean_se". Allowed values are one of, "mean", "mean_se", "mean_sd", "mean_ci", "mean_range",
"median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see desc_statby for more details.

**numeric.x.axis** logical. If TRUE, x axis will be treated as numeric. Default is FALSE.

**combine** logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.

**merge** logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.

**color, fill** outline and fill colors.

**palette** the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggscale R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

**size** Numeric value (e.g.: size = 1). change the size of points and outlines.

**width** numeric value between 0 and 1 specifying box width.

**title** plot main title.

**xlab** character vector specifying x axis labels. Use xlab = FALSE to hide xlab.

**ylab** character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

**facet.by** character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

**panel.labs** a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2" )).

**short.panel.labs** logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

**select** character vector specifying which items to display.

**remove** character vector specifying which items to remove from the plot.

**order** character vector specifying the order of items. Considered only when x axis is a factor variable.

**add** character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see ?desc_statby for more details.

**add.params** parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").
error.plot  plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerrange", "lower_linerrange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where "*" = sd, se, ....

ci  the percent range of the confidence interval (default is 0.95).

position  Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

ggtheme  function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

...  other arguments to be passed to be passed to ggpar().

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation: orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar, ggline

Examples

# Data: ToothGrowth data set we'll be used.
df<- ToothGrowth
head(df, 10)

# Plot mean_se
ggerrorplot(df, x = "dose", y = "len")

# Change desc_stat to mean_sd
# (other values include: mean_sd, mean_ci, median_iqr, ....)
# Add labels
ggerrorplot(df, x = "dose", y = "len",
    desc_stat = "mean_sd")

# Change error.plot to "errorbar" and add mean point
# Visualize the mean of each group
ggerrorplot(df, x = "dose", y = "len",
           add = "mean", error.plot = "errorbar")

# Horizontal plot
ggerrorplot(df, x = "dose", y = "len",
           add = "mean", error.plot = "errorbar",
           orientation = "horizontal")

# Change error.plot to "crossbar"
ggerrorplot(df, x = "dose", y = "len",
           error.plot = "crossbar", width = 0.5)

# Add jitter points and errors (mean_se)
ggerrorplot(df, x = "dose", y = "len",
           add = "jitter")

# Add dot and errors (mean_se)
ggerrorplot(df, x = "dose", y = "len",
           add = "dotplot")

# Multiple groups with error bars and jitter point
ggerrorplot(df, x = "dose", y = "len",
           color = "supp", palette = "Paired",
           error.plot = "pointrange",
           position = position_dodge(0.5))

---

**ggexport**

**Export ggplots**

**Description**

Export ggplots

**Usage**

```r
 ggexport(
   ..., 
   plotlist = NULL,
   filename = NULL,
   ncol = NULL,
   nrow = NULL,
   width = 480,
   height = 480,
   pointsize = 12,
   res = NA,
   verbose = TRUE
 )
```
Arguments

... list of plots to be arranged into the grid. The plots can be either ggplot2 plot objects, arbitrary gtables or an object of class `ggarrange`.

`plotlist` (optional) list of plots to display.

`filename` File name to create on disk.

`ncol` (optional) number of columns in the plot grid.

`nrow` (optional) number of rows in the plot grid.

`width, height` plot width and height, respectively (example, width = 800, height = 800). Applied only to raster plots: "png", "jpeg", "jpg", "bmp" and "tiff".

`pointsize` the default pointsize of plotted text (example, pointsize = 8). Used only for raster plots.

`res` the resolution in ppi (example, res = 250). Used only for raster plots.

`verbose` logical. If TRUE, show message.

Author(s)

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Examples

```r
## Not run:
require("magrittr")
# Load data
data("ToothGrowth")
df <- ToothGrowth
df$dose <- as.factor(df$dose)

# Box plot
bxp <- ggboxplot(df, x = "dose", y = "len",
                 color = "dose", palette = "jco")

# Dot plot
dp <- ggdotplot(df, x = "dose", y = "len",
                color = "dose", palette = "jco")

# Density plot
dens <- ggdensity(df, x = "len", fill = "dose", palette = "jco")

# Export to pdf
ggarrange(bxp, dp, dens, ncol = 2) %>%
ggexport(filename = "test.pdf")

# Export to png
ggarrange(bxp, dp, dens, ncol = 2) %>%
ggexport(filename = "test.png")

## End(Not run)
```
gghistogram  

Histogram plot

Description
Create a histogram plot.

Usage
gghistogram(
da
data,   
x,   
y = "count",   
combine = FALSE,   
merge = FALSE,   
weight = NULL,   
color = "black",   
fill = NA,   
palette = NULL,   
size = NULL,   
linetype = "solid",   
alpha = 0.5,   
bins = NULL,   
binwidth = NULL,   
title = NULL,   
xlab = NULL,   
ylab = NULL,   
facet.by = NULL,   
panel.labs = NULL,   
short.panel.labs = TRUE,   
add = c("none", "mean", "median"),   
add.params = list(linetype = "dashed"),   
rug = FALSE,   
add_density = FALSE,   
label = NULL,   
font.label = list(size = 11, color = "black"),   
label.select = NULL,   
repel = FALSE,   
label.rectangle = FALSE,   
position = position_identity(),   
ggtheme = theme_pubr(),   
...   
)

Arguments
data  
a data frame
x variable to be drawn.
y one of "density" or "count".
combine logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
weight a variable name available in the input data for creating a weighted histogram.
color, fill histogram line color and fill color.
palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
size Numeric value (e.g.: size = 1). change the size of points and outlines.
linetype line type. See show_line_types.
alpha numeric value specifying fill color transparency. Value should be in [0, 1], where 0 is full transparency and 1 is no transparency.
bins Number of bins. Defaults to 30.
binwidth numeric value specifying bin width. use value between 0 and 1 when you have a strong dense dotplot. For example binwidth = 0.2.
title plot main title.
xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
panel.labs a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2") ).
short.panel.labs logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.
add allowed values are one of "mean" or "median" (for adding mean or median line, respectively).
add.params parameters (color, size, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").
rug logical value. If TRUE, add marginal rug.
### add_density

Logical value. If TRUE, add density curves.

### label

The name of the column containing point labels. Can be also a character vector with length = nrow(data).

### font.label

A list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example, `font.label = list(size = 14, face = "bold", color = "red")`. To specify only the size and the style, use `font.label = list(size = 14, face = "plain")`.

### label.select

Can be of two formats:

- A character vector specifying some labels to show.
- A list containing one or the combination of the following components:
  - `top.up` and `top.down`: to display the labels of the top up/down points. For example, `label.select = list(top.up = 10, top.down = 4)`.
  - `criteria`: to filter, for example, by x and y variables values, use this: `label.select = list(criteria = "y > 2 & y < 5 & x %in% c('A', 'B')")`.

### repel

A logical value, whether to use ggrepel to avoid overplotting text labels or not.

### label.rectangle

A logical value. If TRUE, add rectangle underneath the text, making it easier to read.

### position

Position adjustment, either as a string, or the result of a call to a position adjustment function. Allowed values include "identity", "stack", "dodge".

### ggtheme

A function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... Other arguments to be passed to `geom_histogram` and `ggpar`.

### Details

The plot can be easily customized using the function `ggpar()`. Read `?ggpar` for changing:

- Main and axis labels: `main`, `xlab`, `ylab`
- Axis limits: `xlim`, `ylim` (e.g.: `ylim = c(0, 30)`)  
- Axis scales: `xscale`, `yscale` (e.g.: `yscale = "log2"`)  
- Color palettes: `palette = "Dark2"` or `palette = c("gray", "blue", "red")`  
- Legend title, labels and position: `legend = "right"`  
- Plot orientation: `orientation = c("vertical", "horizontal", "reverse")`

### See Also

- `ggdensity`
- `ggpar`
**Examples**

```r
# Create some data format
set.seed(1234)
wdata = data.frame(
  sex = factor(rep(c("F", "M"), each=200)),
  weight = c(rnorm(200, 55), rnorm(200, 58)))

head(wdata, 4)

# Basic density plot
# Add mean line and marginal rug
gghistogram(wdata, x = "weight", fill = "lightgray",
            add = "mean", rug = TRUE)

# Change outline colors by groups ("sex")
# Use custom color palette
gghistogram(wdata, x = "weight",
            add = "mean", rug = TRUE,
            color = "sex", palette = c("#00AFBB", "#E7B800"))

# Change outline and fill colors by groups ("sex")
# Use custom color palette
gghistogram(wdata, x = "weight",
            add = "mean", rug = TRUE,
            color = "sex", fill = "sex",
            palette = c("#00AFBB", "#E7B800"))

# Combine histogram and density plots
gghistogram(wdata, x = "weight",
            add = "mean", rug = TRUE,
            fill = "sex", palette = c("#00AFBB", "#E7B800"),
            add_density = TRUE)

# Weighted histogram
gghistogram(iris, x = "Sepal.Length", weight = "Petal.Length")
```

---

**ggline**

*Line plot*

**Description**

Create a line plot.

**Usage**

```r
ggline(data,
       
```
Arguments

- **data**: a data frame
- **x, y**: x and y variables for drawing.
- **group**: grouping variable to connect points by line. Allowed values are 1 (for one line, one group) or a character vector specifying the name of the grouping variable (case of multiple lines).
- **numeric.x.axis**: logical. If TRUE, x axis will be treated as numeric. Default is FALSE.
- **combine**: logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot.
of y variables.

merge logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.

color line colors.

palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucesgb", "uchicago", "simpsons" and "rickandmorty".

linetype line type.

plot_type plot type. Allowed values are one of "b" for both line and point; "l" for line only; and "p" for point only. Default is "b".

size Numeric value (e.g.: size = 1). change the size of points and outlines.

shape point shapes.

stroke point stroke. Used only for shapes 21-24 to control the thickness of points border.

point.size point size.

point.color point color.

title plot main title.

xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.

ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

panel.labs a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2") )

short.panel.labs logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

select character vector specifying which items to display.

remove character vector specifying which items to remove from the plot.

order character vector specifying the order of items.

add character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_q1q3", "median_range": see ?desc_statby for more details.
add.params parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

error.plot plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where "*" = sd, se, ....

label the name of the column containing point labels. Can be also a character vector with length = nrow(data).

font.label a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color = "red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").

label.select can be of two formats:

• a character vector specifying some labels to show.
• a list containing one or the combination of the following components:
  – top.up and top.down: to display the labels of the top up/down points.
    For example, label.select = list(top.up = 10, top.down = 4).
  – criteria: to filter, for example, by x and y variables values, use this:
    label.select = list(criteria = "x > 2 & y < 5 & x %in% c('A', 'B')").

repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.

label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.

show.line.label logical value. If TRUE, shows line labels.

position Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

• main title and axis labels: main, xlab, ylab
• axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
• axis scales: xscale,yscale (e.g.: yscale = "log2")
• color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
• legend title, labels and position: legend = "right"
• plot orientation: orientation = c("vertical", "horizontal", "reverse")
**ggline**

**See Also**

`ggpar`, `ggbarplot`

**Examples**

```r
# Data
df <- data.frame(dose=c("D0.5", "D1", "D2"),
               len=c(4.2, 10, 29.5))
print(df)

# Basic plot
# +++++++++++++++++++++++++++
ggline(df, x = "dose", y = "len")

# Plot with multiple groups
# +++++++++++++++++++++++++

# Create some data
df2 <- data.frame(supp=rep(c("VC", "OJ"), each=3),
                  dose=rep(c("D0.5", "D1", "D2"),2),
                  len=c(6.8, 15, 33, 4.2, 10, 29.5))
print(df2)

# Plot "len" by "dose" and
# Change line types and point shapes by a second groups: "supp"
ggline(df2, "dose", "len",
       linetype = "supp", shape = "supp")

# Change colors
# +++++++++++++++++++++

# Change color by group: "supp"
# Use custom color palette
ggline(df2, "dose", "len",
       linetype = "supp", shape = "supp",
       color = "supp", palette = c("#00AFBB", "#E7B800"))

# Add points and errors
# +++++++++++++++++++++++

# Data: ToothGrowth data set we'll be used.
df3 <- ToothGrowth
head(df3, 10)

# It can be seen that for each group we have
# different values
ggline(df3, x = "dose", y = "len")

# Visualize the mean of each group
```
ggline(df3, x = "dose", y = "len",
       add = "mean")

# Add error bars: mean_se
# (other values include: mean_sd, mean_ci, median_iqr, ....)
# Add labels
ggline(df3, x = "dose", y = "len", add = "mean_se")

# Change error.plot to "pointrange"
ggline(df3, x = "dose", y = "len",
       add = "mean_se", error.plot = "pointrange")

# Add jitter points and errors (mean_se)
ggline(df3, x = "dose", y = "len",
       add = c("mean_se", "jitter"))

# Add dot and errors (mean_se)
ggline(df3, x = "dose", y = "len",
       add = c("mean_se", "dotplot"), color = "steelblue")

# Add violin and errors (mean_se)
ggline(df3, x = "dose", y = "len",
       add = c("mean_se", "violin"), color = "steelblue")

# Multiple groups with error bars
# ++++++++++++++++++++++

ggline(df3, x = "dose", y = "len", color = "supp",
       add = "mean_se", palette = c("#00AFBB", "#E7B800"))

# Add jitter
ggline(df3, x = "dose", y = "len", color = "supp",
       add = c("mean_se", "jitter"), palette = c("#00AFBB", "#E7B800"))

# Add dot plot
ggline(df3, x = "dose", y = "len", color = "supp",
       add = c("mean_se", "dotplot"), palette = c("#00AFBB", "#E7B800"))

---

**ggmaplot**

**MA-plot from means and log fold changes**

**Description**

Make MA-plot which is a scatter plot of log2 fold changes (M, on the y-axis) versus the average expression signal (A, on the x-axis). \( M = \log_2(x/y) \) and \( A = (\log_2(x) + \log_2(y))/2 = \log_2(xy)^{1/2} \), where x and y are respectively the mean of the two groups being compared.
Usage

```r
ggmaplot(
  data,
  fdr = 0.05,
  fc = 1.5,
  genenames = NULL,
  detection_call = NULL,
  size = NULL,
  alpha = 1,
  seed = 42,
  font.label = c(12, "plain", "black"),
  label.rectangle = FALSE,
  palette = c("#B31B21", "#1465AC", "darkgray"),
  top = 15,
  select.top.method = c("padj", "fc"),
  label.select = NULL,
  main = NULL,
  xlab = "Log2 mean expression",
  ylab = "Log2 fold change",
  ggtheme = theme_classic(),
  ...
)
```

Arguments

data an object of class DESeqResults, get_diff, DE_Results, matrix or data frame containing the columns baseMean (or baseMeanLog2), log2FoldChange, and padj. Rows are genes.

Two possible formats are accepted for the input data:

- 1/ baseMean | log2FoldChange | padj. This is a typical output from DESeq2 pipeline. Here, we’ll use log2(baseMean) as the x-axis variable.
- 2/ baseMeanLog2 | log2FoldChange | padj. Here, baseMeanLog2 is assumed to be the mean of logged values; so we’ll use it as the x-axis variable without any transformation. This is the real A in MA plot. In other words, it is the average of two log-scales values: A = (log2(x) + log2(y))/2 = log2(xy)*1/2

Terminology:

- baseMean: the mean expression of genes in the two groups.
- log2FoldChange: the log2 fold changes of group 2 compared to group 1
- padj: the adjusted p-value of the used statistical test.

fdr Accepted false discovery rate for considering genes as differentially expressed.

fc the fold change threshold. Only genes with a fold change >= fc and padj <= fdr are considered as significantly differentially expressed.

genenames a character vector of length nrow(data) specifying gene names corresponding to each row. Used for point labels.
`detection_call` a numeric vector with length = nrow(data), specifying if the genes is expressed (value = 1) or not (value = 0). For example `detection_call = c(1, 1, 0, 1, 0, 1)`. Default is `NULL`. If `detection_call` column is available in data, it will be used.

`size` points size.

`alpha` numeric value between 0 an 1 specifying point alpha for controlling transparency. For example, use `alpha = 0.5`.

`seed` Random seed passed to `set.seed`. If `NA`, `set.seed` will not be called. Default is 42 for reproducibility.

`font.label` a vector of length 3 indicating respectively the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of point labels. For example `font.label = c(14, "bold", "red")`.

`label.rectangle` logical value. If TRUE, add rectangle underneat the text, making it easier to read.

`palette` the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

`top` the number of top genes to be shown on the plot. Use `top = 0` to hide to gene labels.

`select.top.method` methods to be used for selecting top genes. Allowed values include "padj" and "fc" for selecting by adjusted p values or fold changes, respectively.

`label.select` character vector specifying some labels to show.

`main` plot main title.

`xlab` character vector specifying x axis labels. Use `xlab = FALSE` to hide xlab.

`ylab` character vector specifying y axis labels. Use `ylab = FALSE` to hide ylab.

`ggtheme` function, ggplot2 theme name. Default value is `theme_pubr()`. Allowed values include ggplot2 official themes: `theme_gray()`, `theme_bw()`, `theme_minimal()`, `theme_classic()`, `theme_void()`, ....

... other arguments to be passed to `ggpar`.

**Value**

returns a ggplot.

**Examples**

data(diff_express)

# Default plot
ggmaplot(diff_express, main = expression("Group 1" %->% "Group 2"),
  fdr = 0.05, fc = 2, size = 0.4,
  palette = c("#B31B21", "#1465AC", "darkgray"),
  genenames = as.vector(diff_express$name),
  ... other arguments to be passed to ggpar.
```r
legend = "top", top = 20,
font.label = c("bold", 11),
font.legend = "bold",
font.main = "bold",
ggtheme = ggplot2::theme_minimal()

# Add rectangle around labels
ggmaplot(diff_express, main = expression("Group 1" %->% "Group 2"),
fdr = 0.05, fc = 2, size = 0.4,
palette = c("#B31B21", "#1465AC", "darkgray"),
genenames = as.vector(diff_express$name),
legend = "top", top = 20,
font.label = c("bold", 11), label.rectangle = TRUE,
font.legend = "bold",
font.main = "bold",
ggtheme = ggplot2::theme_minimal())

# Select specific genes to show
# set top = 0, then specify genes using label.select argument
ggmaplot(diff_express, main = expression("Group 1" %->% "Group 2"),
fdr = 0.05, fc = 2, size = 0.4,
genenames = as.vector(diff_express$name),
ggtheme = ggplot2::theme_minimal(),
top = 0, label.select = c("BUB1", "CD83")
)
```

---

**ggpaired**

**Plot Paired Data**

**Description**

Plot paired data.

**Usage**

```r
ggpaired(
data,
cond1,
cond2,
x = NULL,
y = NULL,
id = NULL,
color = "black",
fill = "white",
palette = NULL,
width = 0.5,
point.size = 1.2,
line.size = 0.5,
```
line.color = "black",
linetype = "solid",
title = NULL,
xlab = "Condition",
ylab = "Value",
facet.by = NULL,
panel.labs = NULL,
short.panel.labs = TRUE,
label = NULL,
font.label = list(size = 11, color = "black"),
label.select = NULL,
repel = FALSE,
label.rectangle = FALSE,
ggtheme = theme_pubr(),
...
)

Arguments

data a data frame
cond1 variable name corresponding to the first condition.
cond2 variable name corresponding to the second condition.
x, y x and y variables, where x is a grouping variable and y contains values for each group. Considered only when cond1 and cond2 are missing.
id variable name corresponding to paired samples' id. Used to connect paired points with lines.
color points and box plot colors. To color by conditions, use color = "condition".
fill box plot fill color. To change fill color by conditions, use fill = "condition".
palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
width box plot width.
point.size, line.size point and line size, respectively.
line.color line color.
linetype line type.
title plot main title.
xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
panel.labs: a list of one or two character vectors to modify facet panel labels. For example, `panel.labs = list(sex = c("Male", "Female"))` specifies the labels for the "sex" variable. For two grouping variables, you can use for example `panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2"))`.

short.panel.labs: logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

label: the name of the column containing point labels. Can be also a character vector with length = nrow(data).

font.label: a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example `font.label = list(size = 14, face = "bold", color = "red")`. To specify only the size and the style, use `font.label = list(size = 14, face = "plain")`.

label.select: can be of two formats:
- a character vector specifying some labels to show.
- a list containing one or the combination of the following components:
  - `top.up` and `top.down`: to display the labels of the top up/down points. For example, `label.select = list(top.up = 10, top.down = 4)`.
  - `criteria`: to filter, for example, by x and y variables values, use this: `label.select = list(criteria = "y > 2 & y < 5 & x %in% c('A', 'B')")`.

repel: a logical value, whether to use ggrepel to avoid overplotting text labels or not.

label.rectangle: logical value. If TRUE, add rectangle underneath the text, making it easier to read.

ggtheme: function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to be passed to `ggpar()`.

Examples

# Example 1
```r
#::::::::::::::::::::::::::::::::::::::::::
before <-c(200.1, 190.9, 192.7, 213, 241.4, 196.9, 172.2, 185.5, 205.2, 193.7)
after <-c(392.9, 393.2, 345.1, 393, 434, 427.9, 422, 383.9, 392.3, 352.2)

d <- data.frame(before = before, after = after)
ggpaired(d, cond1 = "before", cond2 = "after",
         fill = "condition", palette = "jco")
```

# Example 2
```r
#::::::::::::::::::::::::::::::::::::::::::
ggpaired(ToothGrowth, x = "supp", y = "len",
         fill = "condition", palette = "jco")
```
ggpar

Graphical parameters

Description

Graphical parameters

Usage

```r
ggpar(
  p,
  palette = NULL,
  gradient.cols = NULL,
  main = NULL,
  submain = NULL,
  caption = NULL,
  xlab = NULL,
  ylab = NULL,
  title = NULL,
  subtitle = NULL,
  font.main = NULL,
  font.submain = NULL,
  font.x = NULL,
  font.y = NULL,
  font.caption = NULL,
  font.title = NULL,
  font.subtitle = NULL,
  font.family = "",
  xlim = NULL,
  ylim = NULL,
  xscale = c("none", "log2", "log10", "sqrt"),
  yscale = c("none", "log2", "log10", "sqrt"),
  format.scale = FALSE,
  legend = NULL,
  legend.title = NULL,
  font.legend = NULL,
  ticks = TRUE,
  tickslab = TRUE,
  font.tickslab = NULL,
  font.xtickslab = font.tickslab,
  font.ytickslab = font.tickslab,
  x.text.angle = NULL,
  y.text.angle = NULL,
  color = "supp", line.color = "gray", line.size = 0.4,
  palette = "npg")
```
Arguments

- **p** an object of class ggplot or a list of ggplots
- **palette** the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty". Can be also a numeric vector of length(groups); in this case a basic color palette is created using the function palette.
- **gradient.cols** vector of colors to use for n-colour gradient. Allowed values include brewer and ggsci color palettes.
- **main** plot main title.
- **submain, subtitle** plot subtitle.
- **caption** plot caption.
- **xlab** character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
- **ylab** character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
- **title** plot main title.
- **font.main, font.submain, font.caption, font.x, font.y** a vector of length 3 indicating respectively the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of main title, subtitle, caption, xlab and ylab, respectively. For example font.x = c(14, "bold", "red"). Use font.x = 14, to change only font size; or use font.x = "bold", to change only font face.
- **font.title, font.subtitle** alias of font.submain and font.submain, respectively.
- **font.family** character vector specifying font family.
- **xlim, ylim** a numeric vector of length 2, specifying x and y axis limits (minimum and maximum), respectively. e.g.: ylim = c(0, 50).
- **xscale, yscale** x and y axis scale, respectively. Allowed values are one of c("none", "log2", "log10", "sqrt"); e.g.: yscale="log2".
- **format.scale** logical value. If TRUE, axis tick mark labels will be formatted when xscale or yscale = "log2" or "log10".
legend character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). To remove the legend use legend = "none". Legend position can be also specified using a numeric vector c(x, y); see details section.

legend.title legend title, e.g.: legend.title = "Species". Can be also a list, legend.title = list(color = "Species", linetype = "Species", shape = "Species").

font.legend legend text font style; e.g.: font.legend = c(10, "plain", "black").

ticks logical value. Default is TRUE. If FALSE, hide axis tick marks.

tickslab logical value. Default is TRUE. If FALSE, hide axis tick labels.

font.tickslab, font.xtickslab, font.ytickslab Font style (size, face, color) for tick labels, e.g.: c(14, "bold", "red").

x.text.angle, y.text.angle Numeric value specifying the rotation angle of x and y axis tick labels, respectively. Default value is NULL. For vertical x axis texts use x.text.angle = 90.

xtickslab.rt, ytickslab.rt Same as x.text.angle and y.text.angle, respectively. Will be deprecated in the near future.

xticks.by, yticks.by numeric value controlling x and y axis breaks, respectively. For example, if yticks.by = 5, a tick mark is shown on every 5. Default value is NULL.

rotate logical value. If TRUE, rotate the graph by setting the plot orientation to horizontal.

orientation change the orientation of the plot. Allowed values are one of c("vertical", "horizontal", "reverse"). Partial match is allowed.

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... not used

Examples

# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic box plot
# ++++++++++++++++++++++++++
p <- ggboxplot(df, x = "dose", y = "len")

# Change the plot orientation: horizontal
ggpar(p, orientation = "horiz")

# Change main title and axis labels
# +++++++++++++++++++++++++++
ggpar(p,  
  main = "Plot of length \n by dose",  
  xlab = "Dose (mg)", ylab = "Length")

# Title font styles: 'plain', 'italic', 'bold', 'bold.italic'
ggpar(p,  
  main = "Length by dose",  
  font.main = c(14,"bold.italic", "red"),  
  font.x = c(14, "bold", "#2E9FDF"),  
  font.y = c(14, "bold", "#E7B800"))

# Hide axis labels  
ggpar(p, xlab = FALSE, ylab = FALSE)

# Change colors  
# ++++++++++++++++++++++

# Change outline colors by groups: dose  
p2 <- ggboxplot(df, "dose", "len", color = "dose")  
p2

# Use custom color palette  
ggpar(p2, palette = c("#00AFBB", "#E7B800", "#FC4E07"))

# Use brewer palette  
ggpar(p2, palette = "Dark2")

# Use grey palette  
ggpar(p2, palette = "grey")

# Use scientific journal palette from ggsci package  
ggpar(p2, palette = "npg")  # nature

# Axis ticks, limits, scales  
# ++++++++++++++++++++++++++

# Axis ticks labels and rotation  

# Axis limits  

# Axis scale  
ggpar(p, yscale = "log2")

# Format axis scale  
ggpar(p, yscale = "log2", format.scale = TRUE)
# Legends
# ++++++++++++++++++
# Change legend position and title
ggpar(p2,
  legend = "right", legend.title = "Dose (mg)",
  font.legend = c(10, "bold", "red"))

---

**Draw a Paragraph of Text**

**Description**

Draw a paragraph of text. Splits a long text into multiple lines (by inserting line breaks) so that the output will fit within the current viewport.

**Usage**

```r
ggparagraph(
  text,
  color = NULL,
  size = NULL,
  face = NULL,
  family = NULL,
  lineheight = NULL
)
```

```r
## S3 method for class 'splitText'
drawDetails(x, recording)
```

**Arguments**

- `text`: the text to plot.
- `color`: font color, example: color = "black"
- `size`: font size, example: size = 12
- `face`: font face. Allowed values are one of "plain", "italic", "bold", "bold.italic".
- `family`: font family
- `x`: a grid grob
- `recording`: a logical value indicating whether a grob is being added to the display list or redrawn from the display list.

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**Examples**

# Density plot
density.p <- ggdensity(iris, x = "Sepal.Length",
                       fill = "Species", palette = "jco")

# Text plot
text <- paste("iris data set gives the measurements in cm",
               "of the variables sepal length and width",
               "and petal length and width, respectively.",
               "for 50 flowers from each of 3 species of iris.",
               "The species are Iris setosa, versicolor, and virginica.", sep = " ")
text.p <- ggparagraph(text, face = "italic", size = 12)

# Arrange the plots on the same page
ggarrange(density.p, text.p,
          ncol = 1, nrow = 2,
          heights = c(1, 0.3))

---

**ggpie**  
*Pie chart*

**Description**

Create a pie chart.

**Usage**

```r
ggpie(
data,  
x,  
label = x,  
lab.pos = c("out", "in"),  
lab.adjust = 0,  
lab.font = c(4, "plain", "black"),  
font.family = "",  
color = "black",  
fill = "white",  
palette = NULL,  
size = NULL,  
ggtheme = theme_pubr(),  
...  
)
```

**Arguments**

- `data`  
a data frame
- `x`  
variable containing values for drawing.
label  variable specifying the label of each slice.
lab.pos character specifying the position for labels. Allowed values are "out" (for outside) or "in" (for inside).
lab.adjust numeric value, used to adjust label position when lab.pos = "in". Increase or decrease this value to see the effect.
lab.font a vector of length 3 indicating respectively the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of label font. For example lab.font= c(4, "bold", "red").
font.family character vector specifying font family.
color, fill outline and fill colors.
palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
size Numeric value (e.g.: size = 1). change the size of points and outlines.
ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....
  ... other arguments to be passed to be passed to ggpar().

Details
The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation : orientation = c("vertical", "horizontal", "reverse")

See Also
ggpar, ggline

Examples

# Data: Create some data
# +++++++++++++++++++++++++++++++

df <- data.frame(
  group = c("Male", "Female", "Child"),
  value = c(25, 25, 50))
# Basic pie charts
# ++++++++++++++++++++++++++++++++
ggpie(df, "value", label = "group")

# Reducing margins around the pie chart
ggpie(df, "value", label = "group") +
  theme( plot.margin = unit(c(-.75,-.75,-.75,-.75),"cm"))

# Change color
# ++++++++++++++++++++++++++++++++

# Change fill color by group
# set line color to white
# Use custom color palette
ggpie(df, "value", label = "group",
      fill = "group", color = "white",
      palette = c("#00AFBB", "#E7B800", "#FC4E07") )

# Change label
# ++++++++++++++++++++++++++++++++

# Show group names and value as labels
labs <- paste0(df$group, " (", df$value, ")")
ggpie(df, "value", label = labs,
      fill = "group", color = "white",
      palette = c("#00AFBB", "#E7B800", "#FC4E07") )

# Change the position and font color of labels
ggpie(df, "value", label = labs,
      lab.pos = "in", lab.font = "white",
      fill = "group", color = "white",
      palette = c("#00AFBB", "#E7B800", "#FC4E07") )
Arguments

data | a data frame
x | character string containing the name of x variable.
y | character vector containing one or more variables to plot
combine | logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge | logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
color | outline color.
fill | fill color.
palette | the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsce R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorthy".
linetype | line types.
size | numeric value (e.g.: size = 1). change the size of points and outlines.
select | character vector specifying which items to display.
remove | character vector specifying which items to remove from the plot.
order | character vector specifying the order of items.
add | character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_mad", "median_range"; see ?desc_statby for more details.
add.params | parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").
error.plot | plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where "*" = sd, se, ....
font.label | a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color = "red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").
title | plot main title.
xlab | character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ggpubr_options

Description
Displays allowed global options in ggpubr.

Usage
ggpubr_options()

Examples
ggpubr_options()

ggqqplot

Description
Quantile-Quantile plot.

Usage
ggqqplot(
data, x, 
combine = FALSE, 
merge = FALSE, 
color = "black", 
palette = NULL, 
size = NULL, 
shape = NULL, 
add = c("qqline", "none"), 
add.params = list(linetype = "solid"), 
conf.int = TRUE, 
conf.int.level = 0.95, 
title = NULL, 
xlab = NULL,
Arguments

data: a data frame

x: variable to be drawn.

combine: logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.

merge: logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.

color: point color.

palette: the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

size: point size.

shape: point shape.

add: character vector. Allowed values are one of "none" and "qqline" (for adding qqline).

add.params: parameters (color, size, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

conf.int: logical value. If TRUE, confidence interval is added.

conf.int.level: the confidence level. Default value is 0.95.

title: plot main title.

xlab: character vector specifying x axis labels. Use xlab = FALSE to hide xlab.

ylab: character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

facet.by: character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

panel.labs: a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2")) 

short.panel.labs

Logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

ggtheme

Function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to ggpar.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation: orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar

Examples

# Create some data format
set.seed(1234)
wdata = data.frame(
  sex = factor(rep(c("F", "M"), each=200)),
  weight = c(rnorm(200, 55), rnorm(200, 58)))
head(wdata, 4)

# Basic QQ plot
ggqqplot(wdata, x = "weight")

# Change colors and shape by groups ("sex")
# Use custom palette
ggqqplot(wdata, x = "weight",
  color = "sex", palette = c("#00AFBB", "#E7B800"))
**ggscatter**

*Scatter plot*

**Description**
Create a scatter plot.

**Usage**

```r
ggscatter(
  data, x, y, 
  combine = FALSE, merge = FALSE, color = "black", fill = "lightgray", palette = NULL, shape = 19, size = 2, point = TRUE, rug = FALSE, title = NULL, xlab = NULL, ylab = NULL, facet.by = NULL, panel.labs = NULL, short.panel.labs = TRUE, add = c("none", "reg.line", "loess"), add.params = list(), conf.int = FALSE, conf.int.level = 0.95, fullrange = FALSE, ellipse = FALSE, ellipse.level = 0.95, ellipse.type = "norm", ellipse.alpha = 0.1, ellipse.border.remove = FALSE, mean.point = FALSE, mean.point.size = ifelse(is.numeric(size), 2 * size, size), star.plot = FALSE, star.plot.lty = 1, star.plot.lwd = NULL, label = NULL, font.label = c(12, "plain"), font.family = "", label.select = NULL,
```

- `data`: Data frame.
- `x`: Independent variable.
- `y`: Dependent variable.
- `combine`: Combine points if equal.
- `merge`: Merge points if equal.
- `color`: Color of points.
- `fill`: Color of fill.
- `palette`: A palette.
- `shape`: Shape of points.
- `size`: Size of points.
- `point`: Draw points.
- `title`: Title.
- `xlab`: X-axis label.
- `ylab`: Y-axis label.
- `facet.by`: Facet by.
- `panel.labs`: Panel labels.
- `short.panel.labs`: Short panel labels.
- `add`: Add linear regression or loess.
- `add.params`: Parameters for `add`.
- `conf.int`: Confidence interval.
- `conf.int.level`: Level of confidence interval.
- `fullrange`: Full range.
- `ellipse.level`: Level of ellipse.
- `ellipse.type`: Type of ellipse.
- `ellipse.alpha`: Alpha of ellipse.
- `ellipse.border.remove`: Remove border.
- `mean.point`: Draw mean point.
- `mean.point.size`: Size of mean point.
- `star.plot`: Draw star plot.
- `star.plot.lty`: Line type of star plot.
- `star.plot.lwd`: Line width of star plot.
- `label`: Label.
- `font.label`: Font family.
- `font.family`: Font size.
- `label.select`: Select label.
Arguments

- **data**: a data frame
- **x**: x variables for drawing.
- **y**: y variables for drawing.
- **combine**: logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
- **merge**: logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
- **color, fill**: point colors.
- **palette**: the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
- **shape**: point shape. See show_point_shapes.
- **size**: Numeric value (e.g.: size = 1). change the size of points and outlines.
- **point**: logical value. If TRUE, show points.
- **rug**: logical value. If TRUE, add marginal rug.
- **title**: plot main title.
- **xlab**: character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
- **ylab**: character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
- **facet.by**: character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
- **panel.labs**: a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2")) .
short.panel.labs
logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

add
allowed values are one of "none", "reg.line" (for adding linear regression line) or "loess" (for adding local regression fitting).

add.params
parameters (color, size, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

conf.int
logical value. If TRUE, adds confidence interval.

conf.int.level
Level controlling confidence region. Default is 95%. Used only when add != "none" and conf.int = TRUE.

fullrange
should the fit span the full range of the plot, or just the data. Used only when add != "none".

ellipse
logical value. If TRUE, draws ellipses around points.

ellipse.level
the size of the concentration ellipse in normal probability.

ellipse.type
Character specifying frame type. Possible values are "convex", "confidence" or types supported by stat_ellipse() including one of c("t", "norm", "euclid") for plotting concentration ellipses.

• "convex": plot convex hull of a set of points.
• "confidence": plot confidence ellipses around group mean points as FactoMineR::coord.ellipse().
• "t": assumes a multivariate t-distribution.
• "norm": assumes a multivariate normal distribution.
• "euclid": draws a circle with the radius equal to level, representing the euclidean distance from the center. This ellipse probably won’t appear circular unless coord_fixed() is applied.

ellipse.alpha
Alpha for ellipse specifying the transparency level of fill color. Use alpha = 0 for no fill color.

ellipse.border.remove
logical value. If TRUE, remove ellipse border lines.

mean.point
logical value. If TRUE, group mean points are added to the plot.

mean.point.size
numeric value specifying the size of mean points.

star.plot
logical value. If TRUE, a star plot is generated.

star.plot.lty, star.plot.lwd
line type and line width (size) for star plot, respectively.

label
the name of the column containing point labels. Can be also a character vector with length = nrow(data).

font.label
a vector of length 3 indicating respectively the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of point labels. For example font.label = c(14, "bold", "red"). To specify only the size and the style, use font.label = c(14, "plain").

font.family
character vector specifying font family.

label.select
character vector specifying some labels to show.
repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.
label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.
parse If TRUE, the labels will be parsed into expressions and displayed as described in ?plotmath.
cor.coef logical value. If TRUE, correlation coefficient with the p-value will be added to the plot.
cor.coef.args a list of arguments to pass to the function stat_cor for customizing the displayed correlation coefficients. For example: cor.coef.args = list(method = "pearson", label.x.npc = "right", label.y.npc = "top").
cor.method method for computing correlation coefficient. Allowed values are one of "pearson", "kendall", or "spearman".
cor.coef.coord numeric vector, of length 2, specifying the x and y coordinates of the correlation coefficient. Default values are NULL.
cor.coef.size correlation coefficient text font size.
ggp a ggplot. If not NULL, points are added to an existing plot.
show.legend.text logical. Should text be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....
... other arguments to be passed to geom_point and ggpar.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

• main title and axis labels: main, xlab, ylab
• axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
• axis scales: xscale, yscale (e.g.: yscale = "log2")
• color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
• legend title, labels and position: legend = "right"
• plot orientation : orientation = c("vertical", "horizontal", "reverse")

See Also

stat_cor, stat_stars, stat_conf_ellipse and ggpar.
Examples

# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)
head(df[, c("wt", "mpg", "cyl")], 3)

# Basic plot
# +++++++++++++++++++++++++++
ggscatter(df, x = "wt", y = "mpg",
  color = "black", shape = 21, size = 3, # Points color, shape and size
  add = "reg.line", # Add regression line
  add.params = list(color = "blue", fill = "lightgray"), # Customize reg. line
  conf.int = TRUE, # Add confidence interval
  cor.coef = TRUE, # Add correlation coefficient. see ?stat_cor
  cor.coeff.args = list(method = "pearson", label.x = 3, label.sep = "\n")
)

# loess method: local regression fitting
ggscatter(df, x = "wt", y = "mpg",
  add = "loess", conf.int = TRUE)

# Control point size by continuous variable values ("qsec")
ggscatter(df, x = "wt", y = "mpg",
  color = "#00AFBB", size = "qsec")

# Change colors
# +++++++++++++++++++++++++++
# Use custom color palette
# Add marginal rug

ggscatter(df, x = "wt", y = "mpg",
  color = "#00AFBB", palette = c("#00AFBB", "#E7B800", "#FC4E07")
)

# Add group ellipses and mean points
# Add stars
# ++++++++++++++++++++++++++

ggscatter(df, x = "wt", y = "mpg",
  color = "cyl", shape = "cyl",
  ellipse = TRUE, mean.point = TRUE,
  star.plot = TRUE)

# Textual annotation
# +++++++++++++++
df$name <- rownames(df)
ggscatter(df, x = "wt", y = "mpg",
  text = df$name,
  text.aes = list(x = "wt", y = "mpg", size = "qsec")
)
ggscatterhist

Scatter Plot with Marginal Histograms

Description

Create a scatter plot with marginal histograms, density plots or box plots.

Usage

ggscatterhist(
  data,
  x,
  y,
  group = NULL,
  color = "black",
  fill = NA,
  palette = NULL,
  shape = 19,
  size = 2,
  linetype = "solid",
  bins = 30,
  margin.plot = c("density", "histogram", "boxplot"),
  margin.params = list(),
  margin.ggtheme = theme_void(),
  margin.space = FALSE,
  main.plot.size = 2,
  margin.plot.size = 1,
  title = NULL,
  xlab = NULL,
  ylab = NULL,
  legend = "top",
  ggtheme = theme_pubr(),
  print = TRUE,
  ...
)

## S3 method for class 'ggscatterhist'
print( x,
  margin.space = FALSE,
  main.plot.size = 2,
  margin.plot.size = 1,
### ggscatterhist

```r
title = NULL,
legend = "top",
...
)
```

#### Arguments

- **data**: a data frame
- **x**: an object of class `ggscatterhist`
- **y**: `y` variables for drawing.
- **group**: a grouping variable. Change points color and shape by groups if the options `color` and `shape` are missing. Should be also specified when you want to create a marginal box plot that is grouped.
- **color, fill**: point colors.
- **palette**: the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. `c("blue", "red")`; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
- **shape**: point shape. See `show_point_shapes`.
- **size**: Numeric value (e.g.: size = 1). change the size of points and outlines.
- **linetype**: line type ("solid", "dashed", ...)
- **bins**: Number of histogram bins. Defaults to 30. Pick a better value that fit to your data.
- **margin.plot**: the type of the marginal plot. Default is "hist".
- **margin.params**: parameters to be applied to the marginal plots.
- **margin.ggtheme**: the theme of the marginal plot. Default is `theme_void()`.
- **margin.space**: logical value. If TRUE, adds space between the main plot and the marginal plot.
- **main.plot.size**: the width of the main plot. Default is 2.
- **margin.plot.size**: the width of the marginal plot. Default is 1.
- **title**: plot main title.
- **xlab**: character vector specifying x axis labels. Use `xlab = FALSE` to hide xlab.
- **ylab**: character vector specifying y axis labels. Use `ylab = FALSE` to hide ylab.
- **legend**: specify the legend position. Allowed values include: "top", "bottom", "left", "right".
- **ggtheme**: the theme to be used for the scatter plot. Default is `theme_pubr()`.
- **print**: logical value. If TRUE (default), print the plot.
- **...**: other arguments passed to the function `ggscatter()`. 
ggstripchart

Value
an object of class ggscatterhist, which is list of ggplots, including the following elements:

• sp: main scatter plot;
• xplot: marginal x-axis plot;
• yplot: marginal y-axis plot.

User can modify each of plot before printing.

Examples

# Basic scatter plot with marginal density plot
ggscatterhist(iris, x = "Sepal.Length", y = "Sepal.Width",
        color = "#00AFBB",
        margin.params = list(fill = "lightgray"))

# Grouped data
ggscatterhist(
    iris, x = "Sepal.Length", y = "Sepal.Width",
    color = "Species", size = 3, alpha = 0.6,
    palette = c("#00AFBB", "#E7B800", "#FC4E07"),
    margin.params = list(fill = "Species", color = "black", size = 0.2)
)

# Use boxplot as marginal
ggscatterhist(
    iris, x = "Sepal.Length", y = "Sepal.Width",
    color = "Species", size = 3, alpha = 0.6,
    palette = c("#00AFBB", "#E7B800", "#FC4E07"),
    margin.plot = "boxplot",
    ggtheme = theme_bw()
)

# Add vertical and horizontal line to a ggscatterhist
plots <- ggscatterhist(iris, x = "Sepal.Length", y = "Sepal.Width", print = FALSE)
plots$sp <- plots$sp +
    geom_hline(yintercept = 3, linetype = "dashed", color = "blue") +
    geom_vline(xintercept = 6, linetype = "dashed", color = "red")
plots

---

Stripcharts

Description
Create a stripchart, also known as one dimensional scatter plots. These plots are suitable compared to box plots when sample sizes are small.
Usage

`ggstripchart`
```r
  ggstripchart(
    data,
    x,
    y,
    combine = FALSE,
    merge = FALSE,
    color = "black",
    fill = "white",
    palette = NULL,
    title = NULL,
    xlab = NULL,
    ylab = NULL,
    facet.by = NULL,
    panel.labs = NULL,
    short.panel.labs = TRUE,
    shape = 19,
    size = NULL,
    select = NULL,
    remove = NULL,
    order = NULL,
    add = "mean_se",
    add.params = list(),
    error.plot = "pointrange",
    label = NULL,
    font.label = list(size = 11, color = "black"),
    label.select = NULL,
    repel = FALSE,
    label.rectangle = FALSE,
    jitter = 0.2,
    position = position_jitter(jitter, seed = 123),
    ggtheme = theme_pubr(),
    ...
  )
```

Arguments

- **data**: a data frame
- **x**: character string containing the name of x variable.
- **y**: character vector containing one or more variables to plot
- **combine**: logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
- **merge**: logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip".
If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.

color outline color.

fill fill color.

palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".

title plot main title.

xlab character vector specifying x axis labels. Use xlab = FALSE to hide xlab.

ylab character vector specifying y axis labels. Use ylab = FALSE to hide ylab.

facet.by character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.

panel.labs a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2") )

short.panel.labs logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.

shape point shape

size Numeric value (e.g.: size = 1). change the size of points and outlines.

select character vector specifying which items to display.

remove character vector specifying which items to remove from the plot.

order character vector specifying the order of items.

add character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see ?desc_statby for more details.

add.params parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").

error.plot plot type used to visualize error. Allowed values are one of c("pointrange", "linerange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linerange", "lower_linerange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where ")" = sd, se, ...

label the name of the column containing point labels. Can be also a character vector with length = nrow(data).
font.label a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color = "red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").

label.select can be of two formats:
- a character vector specifying some labels to show.
- a list containing one or the combination of the following components:
  - top.up and top.down: to display the labels of the top up/down points.
    For example, label.select = list(top.up = 10, top.down = 4).
  - criteria: to filter, for example, by x and y variables values, use this:
    label.select = list(criteria = "x > 2 & y < 5 & x %in% c('A', 'B')").

repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.

label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.

jitter the amount of jitter.

position position adjustment, either as a string, or the result of a call to a position adjustment function. Used to adjust position for multiple groups.

ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to geom_jitter, ggpar and facet.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:
- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation: orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar, ggviolin, ggdotplot and ggbboxplot.

Examples

# Load data
data("ToothGrowth")
df <- ToothGrowth
# Basic plot with summary statistics: mean_se
# +++++++++++++++++++++++++++
# Change point shapes by groups: "dose"
ggstripchart(df, x = "dose", y = "len",
shape = "dose", size = 3,
add = "mean_se")

# Use mean_sd
# Change error.plot to "crossbar"
ggstripchart(df, x = "dose", y = "len",
shape = "dose", size = 3,
add = "mean_sd", add.params = list(width = 0.5),
error.plot = "crossbar")

# Add summary statistics
# ++++++++++++++++++++++++++
# Add box plot
ggstripchart(df, x = "dose", y = "len",
shape = "dose", add = "boxplot")

# Add violin + mean_sd
ggstripchart(df, x = "dose", y = "len",
shape = "dose", add = c("violin", "mean_sd"))

# Change colors
# ++++++++++++++++++++++++++
# Change colors by groups: dose
# Use custom color palette
ggstripchart(df, x = "dose", y = "len",
shape = "dose", color = "dose", palette = c("#00AFBB", "#E7B800", "#FC4E07"),
add = "mean_sd")

# Plot with multiple groups
# +++++++++++++++++++++
# Change shape and color by a second group : "supp"
ggstripchart(df, x = "dose", y = "len",
shape = "supp", color = "supp", palette = c("#00AFBB", "#E7B800"))

# Adjust point position
ggstripchart(df, x = "dose", y = "len",
shape = "supp", color = "supp", palette = c("#00AFBB", "#E7B800"),
position = position_dodge(0.8))

# You can also use position_jitterdodge()
# but fill aesthetic is required
ggstripchart(df, x = "dose", y = "len", shape = "supp", color = "supp",
ggsummarytable

**GGPLOT with Summary Stats Table Under the Plot**

**Description**

Create a ggplot with summary stats (n, median, mean, iqr) table under the plot. Read more: How to Create a Beautiful Plots in R with Summary Statistics Labels.

**Usage**

```r
ggsummarytable(
data, x, y, digits = 0, size = 3, color = "black", palette = NULL, facet.by = NULL, labeller = "label_value", position = "identity", ggtheme = theme_pubr(), ...
)
```

```r
ggsummarystats(
data, x, y, summaries = c("n", "median", "iqr"), gfunc = ggboxplot, color = "black", fill = "white", palette = NULL, facet.by = NULL, free.panels = FALSE, labeller = "label_value", heights = c(0.8, 0.2), digits = 0,
```
## S3 method for class 'ggsummarystats'
print(x, heights = c(0.8, 0.2), ...)

## S3 method for class 'ggsummarystats_list'
print(x, heights = c(0.8, 0.2), legend = NULL, ...)

### Arguments

- **data**: a data frame
- **x**: a list of ggsummarystats.
- **y**: character vector containing one or more variables to plot
- **digits**: integer indicating the number of decimal places (round) to be used.
- **size**: Numeric value (e.g.: size = 1). change the size of points and outlines.
- **color**: outline color.
- **palette**: the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggscc R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorte".
- **facet.by**: character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
- **labeller**: Character vector. An alternative to the argument short.panel.labs. Possible values are one of "label_both" (panel labelled by both grouping variable names and levels) and "label_value" (panel labelled with only grouping levels).
- **position**: Position adjustment, either as a string, or the result of a call to a position adjustment function.
- **ggtheme**: function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....
- **...**: other arguments passed to the function ggpar(), facet() or ggarrange() when printing the plot.
- **summaries**: summary stats to display in the table. Possible values are those returned by the function get_summary_stats(), including: "n", "min", "max", "median", "q1", "q2", "q3", "mad", "mean", "sd", "se", "ci".
- **ggfunc**: a ggpubr function, including: ggboxplot, ggviolin, ggdotplot, ggbarplot, ggline, etc. Can be any other ggplot function that accepts the following arguments data, x, color, fill, palette, ggtheme, facet.by.
- **fill**: fill color.
- **free.panels**: logical. If TRUE, create free plot panels when the argument facet.by is specified.
heights a numeric vector of length 2, specifying the heights of the main and the summary table, respectively.

table.font.size the summary table font size.

legend character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). To remove the legend use legend = "none".

Functions

• ggsummarytable(): Create a table of summary stats
• ggsummarystats(): Create a ggplot with a summary stat table under the plot.

Examples

# Data preparation
#::::::::::::::::::::::::::::::::::::::::::::::::
data("ToothGrowth")
df <- ToothGrowth
df$dose <- as.factor(df$dose)
# Add random QC column
set.seed(123)
qc <- rep(c("pass", "fail"), 30)
df$qc <- as.factor(sample(qc, 60))
# Inspect the data
head(df)

# Basic summary stats
#::::::::::::::::::::::::::::::::::::::::::::::::
# Compute summary statistics
summary.stats <- df %>%
  group_by(dose) %>%
  get_summary_stats(type = "common")
summary.stats

# Visualize summary table
ggsummarytable(
  summary.stats, x = "dose", y = c("n", "median", "iqr"),
  ggtheme = theme_bw()
)

# Create plots with summary table under the plot
#::::::::::::::::::::::::::::::::::::::::::::::::
# Basic plot
ggsummarystats(
  df, x = "dose", y = "len",
  ggfunc = ggboxplot, add = "jitter"
)

# Color by groups
Description

Add text to a plot.

Usage

ggettext(
data,
x = NULL,
y = NULL,
label = NULL,
color = "black",
palette = NULL,
size = 11,
face = "plain",
family = "",
show.legend = NA,
label.select = NULL,
repel = FALSE,
label.rectangle = FALSE,
parse = FALSE,
grouping.vars = NULL,
position = "identity",
ggp = NULL,
ggtheme = theme_pubr(),
...
)

Arguments

data a data frame
x, y x and y variables for drawing.
label the name of the column containing point labels. Can be also a character vector with length = nrow(data).
color text font color.
palette the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
size text font size.
face text font style. Allowed values are one of c("plain", "bold", "italic", "bold.italic").
family character vector specifying font family.
show.legend logical. Should text be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
label.select can be of two formats:
• a character vector specifying some labels to show.
• a list containing one or the combination of the following components:
  – top.up and top.down: to display the labels of the top up/down points. For example, label.select = list(top.up = 10, top.down = 4).
  – criteria: to filter, for example, by x and y variabes values, use this:
    label.select = list(criteria = "y > 2 & y < 5 & x %in% c('A', 'B')").
repel a logical value, whether to use ggrepel to avoid overplotting text labels or not.
label.rectangle logical value. If TRUE, add rectangle underneath the text, making it easier to read.
parse If TRUE, the labels will be parsed into expressions and displayed as described in ?plotmath.
grouping.vars grouping variables to sort the data by, when the user wants to display the top n up/down labels.
position Position adjustment, either as a string, or the result of a call to a position adjustment function.
ggp a ggplot. If not NULL, points are added to an existing plot.
ggtheme function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to ggpar.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale,yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"

See Also

ggpar

Examples

# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)
df$name <- rownames(df)
head(df[, c("wt", "mpg", "cyl")], 3)

# Textual annotation
# +++++++++++++++++
ggtext(df, x = "wt", y = "mpg",
       color = "cyl", palette = c("#00AFBB",="#E7B800","#FC4E07"),
       label = "name", repel = TRUE)

# Add rectangle around label
ggtext(df, x = "wt", y = "mpg",
       color = "cyl", palette = c("#00AFBB",="#E7B800","#FC4E07"),
       label = "name", repel = TRUE, label.rectangle = TRUE)
Description

Draw a textual table.

- ggtexttable(): draw a textual table.
- ttheme(): customize table theme.
- rownames_style(), colnames_style(), tbody_style(): helper functions to customize the table row names, column names and body.
- table_cell_font(): access to a table cell for changing the text font (size and face).
- table_cell_bg(): access to a table cell for changing the background (fill, color, linewidth).
- tab_cell_crossout(): cross out a table cell.
- tab_ncol(), tab_nrow(): returns, respectively, the number of columns and rows in a ggtexttable.
- tab_add_hline(): Creates horizontal lines or separators at the top or the bottom side of a given specified row.
- tab_add_vline(): Creates vertical lines or separators at the right or the left side of a given specified column.
- tab_add_border(), tbody_add_border(), thead_add_border(): Add borders to table; tbody is for table body and thead is for table head.
- tab_add_title(), tab_add_footnote(): Add title, subtitle and footnote to a table.

Usage

ggtexttable(
  x,
  rows = rownames(x),
  cols = colnames(x),
  vp = NULL,
  theme = ttheme(),
  ...
)

ttheme(
  base_style = "default",
  base_size = 11,
  base_colour = "black",
  padding = unit(c(4, 4), "mm"),
  colnames.style = colnames_style(size = base_size),
  rownames.style = rownames_style(size = base_size),
  tbody.style = tbody_style(size = base_size)
)

colnames_style(
  color = "black",
  face = "bold",
  size = 12,
fill = "grey80",
linewidth = 1,
linecolor = "white",
parse = FALSE,
...
)

rownames_style(
  color = "black",
  face = "italic",
  size = 12,
  fill = NA,
  linewidth = 1,
  linecolor = "white",
  parse = FALSE,
...
)

tbody_style(
  color = "black",
  face = "plain",
  size = 12,
  fill = c("grey95", "grey90"),
  linewidth = 1,
  linecolor = "white",
  parse = FALSE,
...
)

table_cell_font(tab, row, column, face = NULL, size = NULL, color = NULL)

table_cell_bg(
  tab,
  row,
  column,
  fill = NULL,
  color = NULL,
  linewidth = NULL,
  alpha = NULL
)

tab_cell_crossout(
  tab,
  row,
  column,
  linetype = 1,
  linewidth = 1,
  linecolor = "black",
reduce.size.by = 0
)

tab_ncol(tab)

tab_nrow(tab)

tab_add_hline(
    tab,
    at.row = 2:tab_nrow(tab),
    row.side = c("bottom", "top"),
    from.column = 1,
    to.column = tab_ncol(tab),
    linetype = 1,
    linewidth = 1,
    linecolor = "black"
)

tab_add_vline(
    tab,
    at.column = 2:tab_ncol(tab),
    column.side = c("left", "right"),
    from.row = 1,
    to.row = tab_nrow(tab),
    linetype = 1,
    linewidth = 1,
    linecolor = "black"
)

tab_add_border(
    tab,
    from.row = 2,
    to.row = tab_nrow(tab),
    from.column = 1,
    to.column = tab_ncol(tab),
    linetype = 1,
    linewidth = 1,
    linecolor = "black"
)

tbody_add_border(
    tab,
    from.row = 2,
    to.row = tab_nrow(tab),
    from.column = 1,
    to.column = tab_ncol(tab),
    linetype = 1,
    linewidth = 1,
linecolor = "black"
)
	head_add_border(
  tab,
  from.row = 1,
  to.row = 1,
  from.column = 1,
  to.column = tab_ncol(tab),
  linetype = 1,
  linewidth = 1,
  linecolor = "black"
)

tab_add_title(
  tab,
  text,
  face = NULL,
  size = NULL,
  color = NULL,
  family = NULL,
  padding = unit(1.5, "line"),
  just = "left",
  hjust = NULL,
  vjust = NULL
)

tab_add_footnote(
  tab,
  text,
  face = NULL,
  size = NULL,
  color = NULL,
  family = NULL,
  padding = unit(1.5, "line"),
  just = "right",
  hjust = NULL,
  vjust = NULL
)

Arguments

x
  a data.frame or matrix.
rows
  optional vector to specify row names
cols
  optional vector to specify column names
vp
  optional viewport
theme
  a list, as returned by the function ttheme(), defining the parameters of the table theme. Allowed values include one of ttheme() and ttheme_clean().
extra parameters for text justification, e.g.: hjust and x. Default is "centre" for the body and header, and "right" for the row names. Left justification: hjust = 0, x = 0.1. Right justification: hjust = 1, x = 0.9.

base_style character string the table style/theme. The available themes are illustrated in the ggtexttable-theme.pdf file. Allowed values include one of c("default", "blank", "classic", "minimal", "light", "lBlack", "lBlue", "lRed", "lGreen", "lViolet", "lCyan", "lOrange", "lBlackWhite", "lBlueWhite", "lRedWhite", "lGreenWhite", "lVioletWhite", "lCyanWhite", "lOrangeWhite", "mBlack", "mBlue", "mRed", "mGreen", "mViolet", "mCyan", "mOrange", "mBlackWhite", "mBlueWhite", "mRedWhite", "mGreenWhite", "mVioletWhite", "mCyanWhite", "mOrangeWhite"). Note that, l = "light"; m = "medium".

base_size default font size
base_colour default font colour
padding length-2 unit vector specifying the horizontal and vertical padding of text within each cell

colnames.style a list, as returned by the function colnames_style(), defining the style of the table column names. Considered only when base_size = "default".
rownames.style a list, as returned by the function rownames_style(), defining the style of the table row names. Considered only when base_size = "default".
tbody.style a list, as returned by the function tbody_style(), defining the style of the table body. Considered only when base_size = "default".

color, face, size
text font color, face and size, respectively. Allowed values for face include c("plain", "bold", "italic", "bold.italic").
fill background color.
linewdith, linecolor line width and color, respectively.
parse logical, default behaviour for parsing text as plotmath
tab an object from ggtexttable or from gridExtra::tableGrob().
row, column an integer specifying the row and the column numbers for the cell of interest.
alpha numeric value specifying fill color transparency. Value should be in [0, 1], where 0 is full transparency and 1 is no transparency.
linetype line type
reduce.size.by Numeric value in [0, 1] to reduce the size by.
at.row a numeric vector of row indexes; for example at.row = c(1, 2).
row.side row side to which the horinzotal line should be added. Can be one of c("bottom", "top").
from.column integer indicating the column from which to start drawing the horizontal line.
to.column integer indicating the column to which the horizontal line should end.
at.column a numeric vector of column indexes; for example at.column = c(1, 2).
column.side column side to which the vertical line should be added. Can be one of c("left", "right").
from.row  integer indicating the row from which to start drawing the horizontal line.
to.row    integer indicating the row to which the vertical line should end.
text      text to be added as title or footnote.
family    font family
just       The justification of the text relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". For numeric values, 0 means left (bottom) alignment and 1 means right (top) alignment.
hjust     A numeric vector specifying horizontal justification. If specified, overrides the just setting.
vjust     A numeric vector specifying vertical justification. If specified, overrides the just setting.

Value

an object of class ggplot.

Examples

# data
df <- head(iris)

# Default table
# Remove row names using rows = NULL
ggtexttable(df, rows = NULL)

# Text justification for individual cells/rows/columns (#335)
# First column is left justified i.e., hjust = 0, x = 0.1
# Remaining columns are right justified i.e., hjust = 1, x = 0.9
table_theme <- ttheme(
  tbody.style = tbody_style(
    hjust = as.vector(matrix(c(0, 1, 1, 1, 1), ncol = 5, nrow = nrow(df), byrow = TRUE)),
    x = as.vector(matrix(c(.1, .9, .9,.9, .9), ncol = 5, nrow = nrow(df), byrow = TRUE))
  )
)
ggtexttable(df, rows = NULL, theme = table_theme)

# Blank theme
ggtexttable(df, rows = NULL, theme = ttheme("blank"))

# light theme
ggtexttable(df, rows = NULL, theme = ttheme("light"))

# Column names border only
ggtexttable(df, rows = NULL, theme = ttheme("blank")) %>%
  tab_add_hline(at.row = 1:2, row.side = "top", linewidth = 2)

# classic theme
ggtexttable(df, rows = NULL, theme = ttheme("classic"))
# minimal theme
ggtexttable(df, rows = NULL, theme = ttheme("minimal"))

# Medium blue (mBlue) theme
ggtexttable(df, rows = NULL, theme = ttheme("mBlue"))

# Customize the table as you want
ggtexttable(df, rows = NULL,
    theme = ttheme(
        colnames.style = colnames_style(color = "white", fill = "#8cc257"),
        tbody.style = tbody_style(color = "black", fill = c("#e8f3de", 
            "#d3e8bb"))
    )
)

# Use RColorBrewer palette
# Provide as many fill color as there are rows in the table body, here nrow = 6
ggtexttable(df,
    theme = ttheme(
        colnames.style = colnames_style(fill = "white"),
        tbody.style = tbody_style(fill = get_palette("RdBu", 6))
    )
)

# Text justification
#:::::::::::::::::::::::::::::::::::::::::::::::::
# Default is "centre" for the body and header, and "right" for the row names.
# Left justification: hjust=0, x=0.1
# Right justification: hjust=1, x=0.9
tbody.style = tbody_style(color = "black",
    fill = c("#e8f3de", 
        
            
            
            "#d3e8bb"),
    hjust=1, x=0.9)
ggtexttable(head(iris), rows = NULL,
    theme = ttheme(
        colnames.style = colnames_style(color = "white", fill = "#8cc257"),
        tbody.style = tbody.style
    )
)

# Access and modify the font and
# the background of table cells
# :::::::::::::::::::::::::::::::::::::::::::
tab <- ggtexttable(head(iris), rows = NULL,
    theme = ttheme("classic"))
tab <- table_cell_font(tab, row = 3, column = 2,
    face = "bold")
tab <- table_cell_bg(tab, row = 4, column = 3, linewidth = 5,
    fill="darkolivegreen1", color = "darkolivegreen4")
tab

# Change table cells background and font for column 3,
# Spaning from row 2 to the last row in the data
tab <- ggtexttable(df, rows = NULL, theme = ttheme("classic"))
ggtexttable

```r
# Add separators and borders
# :::::::::::::::::::::::::::::::::::::::::::::::::::
# Table with blank theme
tab <- ggtexttable(df, theme = ttheme("blank"), rows = NULL)
# Add horizontal and vertical lines
# Cross out some cells
# Add titles andd footnote
# :::::::::::::::::::::::::::::::::::::::::::::::::::

tab <- ggtexttable(head(iris), theme = ttheme("light"))
tab_add_hline(at.row = c(1, 2), row.side = "top", linewidth = 3, linetype = 1) %>%
tab_add_hline(at.row = c(7), row.side = "bottom", linewidth = 3, linetype = 1) %>%
tab_add_vline(at.column = 2:tab_ncol(tab), column.side = "left", from.row = 2, linetype = 2)

tab <- ggtexttable(head(iris), theme = ttheme("light"))
tab <- ggtexttable(head(iris), theme = ttheme("light"))

# Add titles andd footnote
# :::::::::::::::::::::::::::::::::::::::::::::::::::
# Add titles and footnote
# Wrap subtitle into multiple lines using strwrap()
main.title <- "Edgar Anderson's Iris Data"
subtitle <- paste0(
  "This famous (Fisher's or Anderson's) iris data set gives the measurements",
  " in centimeters of the variables sepal length and width and petal length and width",
  ", respectively, for 50 flowers from each of 3 species of iris.",
  " The species are Iris setosa, versicolor, and virginica."
) %>%
strwrap(width = 80) %>%
paste(collapse = "\n")

main.title <- "Edgar Anderson's Iris Data"
subtitle <- paste0(
  "This famous (Fisher's or Anderson's) iris data set gives the measurements",
  " in centimeters of the variables sepal length and width and petal length and width",
  ", respectively, for 50 flowers from each of 3 species of iris.",
  " The species are Iris setosa, versicolor, and virginica."
) %>%
strwrap(width = 80) %>%
paste(collapse = "\n")

# Combine density plot and summary table
# :::::::::::::::::::::::::::::::::::::::::::::::::::
# Density plot of "Sepal.Length"
density.p <- ggdensity(iris, x = "Sepal.Length",
  fill = "Species", palette = "jco")

# Draw the summary table of Sepal.Length
# Descriptive statistics by groups
```
stable <- desc_statby(iris, measure.var = "Sepal.Length",
                        grps = "Species")
stable <- stable[, c("Species", "length", "mean", "sd")]
stable.p <- ggtxttable(stable, rows = NULL,
                          theme = ttheme("mOrange"))

# Arrange the plots on the same page
ggarrange(density.p, stable.p,
          ncol = 1, nrow = 2,
          heights = c(1, 0.5))

---

**ggviolin**

**Violin plot**

**Description**

Create a violin plot with error bars. Violin plots are similar to box plots, except that they also show the kernel probability density of the data at different values.

**Usage**

```r
ggviolin(
data, 
  x, 
  y, 
  combine = FALSE, 
  merge = FALSE, 
  color = "black", 
  fill = "white", 
  palette = NULL, 
  alpha = 1, 
  title = NULL, 
  xlab = NULL, 
  ylab = NULL, 
  facet.by = NULL, 
  panel.labs = NULL, 
  short.panel.labs = TRUE, 
  linetype = "solid", 
  trim = FALSE, 
  size = NULL, 
  width = 1, 
  draw_quantiles = NULL, 
  select = NULL, 
  remove = NULL, 
  order = NULL, 
  add = "mean_se", 
  add.params = list(),
)```
Arguments

data          a data frame
x             character string containing the name of x variable.
y             character vector containing one or more variables to plot
combine       logical value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, create a multi-panel plot by combining the plot of y variables.
merge         logical or character value. Default is FALSE. Used only when y is a vector containing multiple variables to plot. If TRUE, merge multiple y variables in the same plotting area. Allowed values include also "asis" (TRUE) and "flip". If merge = "flip", then y variables are used as x tick labels and the x variable is used as grouping variable.
color         outline color.
fill          fill color.
palette       the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. c("blue", "red"); and scientific journal palettes from ggspec R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
alpha         color transparency. Values should be between 0 and 1.
title         plot main title.
xlab          character vector specifying x axis labels. Use xlab = FALSE to hide xlab.
ylab          character vector specifying y axis labels. Use ylab = FALSE to hide ylab.
facet.by       character vector, of length 1 or 2, specifying grouping variables for faceting the plot into multiple panels. Should be in the data.
panel.labs     a list of one or two character vectors to modify facet panel labels. For example, panel.labs = list(sex = c("Male", "Female")) specifies the labels for the "sex" variable. For two grouping variables, you can use for example panel.labs = list(sex = c("Male", "Female"), rx = c("Obs", "Lev", "Lev2"))
short.panel.labs logical value. Default is TRUE. If TRUE, create short labels for panels by omitting variable names; in other words panels will be labelled only by variable grouping levels.
linetype  line types.
trim  If TRUE (default), trim the tails of the violins to the range of the data. If FALSE, don’t trim the tails.
size  Numeric value (e.g.: size = 1). change the size of points and outlines.
width  violin width.
draw_quantiles  If not (NULL) (default), draw horizontal lines at the given quantiles of the density estimate.
select  character vector specifying which items to display.
remove  character vector specifying which items to remove from the plot.
order  character vector specifying the order of items.
add  character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see ?desc_statby for more details.
add.params  parameters (color, shape, size, fill, linetype) for the argument ‘add’; e.g.: add.params = list(color = "red").
error.plot  plot type used to visualize error. Allowed values are one of c("pointrange", "linrange", "crossbar", "errorbar", "upper_errorbar", "lower_errorbar", "upper_pointrange", "lower_pointrange", "upper_linrange", "lower_linrange"). Default value is "pointrange" or "errorbar". Used only when add != "none" and add contains one "mean_*" or "med_*" where ":" = sd, se, ...
label  the name of the column containing point labels. Can be also a character vector with length = nrow(data).
font.label  a list which can contain the combination of the following elements: the size (e.g.: 14), the style (e.g.: "plain", "bold", "italic", "bold.italic") and the color (e.g.: "red") of labels. For example font.label = list(size = 14, face = "bold", color ="red"). To specify only the size and the style, use font.label = list(size = 14, face = "plain").
label.select  can be of two formats:
• a character vector specifying some labels to show.
• a list containing one or the combination of the following components:
  – top.up and top.down: to display the labels of the top up/down points. For example, label.select = list(top.up = 10, top.down = 4).
  – criteria: to filter, for example, by x and y variabes values, use this:
    label.select = list(critera = "\'y\' > 2 & \'y\' < 5 & \'x\' %in% c('A', 'B').")
repel  a logical value, whether to use ggrepel to avoid overplotting text labels or not.
label.rectangle  logical value. If TRUE, add rectangle underneath the text, making it easier to read.
position  Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.
ggviolin function, ggplot2 theme name. Default value is theme_pubr(). Allowed values include ggplot2 official themes: theme_gray(), theme_bw(), theme_minimal(), theme_classic(), theme_void(), ....

... other arguments to be passed to geom_violin, ggpar and facet.

Details

The plot can be easily customized using the function ggpar(). Read ?ggpar for changing:

- main title and axis labels: main, xlab, ylab
- axis limits: xlim, ylim (e.g.: ylim = c(0, 30))
- axis scales: xscale, yscale (e.g.: yscale = "log2")
- color palettes: palette = "Dark2" or palette = c("gray", "blue", "red")
- legend title, labels and position: legend = "right"
- plot orientation: orientation = c("vertical", "horizontal", "reverse")

See Also

ggpar

Examples

# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
# +++++++++++++++++++++++++++
ggviolin(df, x = "dose", y = "len")
# Change the plot orientation: horizontal
ggviolin(df, "dose", "len", orientation = "horiz")

# Add summary statistics
# ++++++++++++++++++++++++++
# Draw quantiles
ggviolin(df, "dose", "len", add = "none",
         draw_quantiles = 0.5)

# Add box plot
ggviolin(df, x = "dose", y = "len",
         add = "boxplot")

ggviolin(df, x = "dose", y = "len",
         add = "dotplot")

# Add jitter points and
# change point shape by groups ("dose")
ggviolin(df, x = "dose", y = "len",
         add = "jitter", shape = "dose")
# Add mean_sd + jittered points
ggviolin(df, x = "dose", y = "len",
        add = c("jitter", "mean_sd"))

# Change error.plot to "crossbar"
ggviolin(df, x = "dose", y = "len",
        add = "mean_sd", error.plot = "crossbar")

# Change colors
# +++++++++++++++++++++++++++
# Change outline and fill colors
ggviolin(df, "dose", "len",
        color = "black", fill = "gray")

# Change outline colors by groups: dose
# Use custom color palette and add boxplot
ggviolin(df, "dose", "len", color = "dose",
        palette = c("#00AFBB", "#E7B800", "#FC4E07"),
        add = "boxplot")

# Change fill color by groups: dose
# add boxplot with white fill color
ggviolin(df, "dose", "len", fill = "dose",
        palette = c("#00AFBB", "#E7B800", "#FC4E07"),
        add = "boxplot", add.params = list(fill = "white"))

# Plot with multiple groups
# +++++++++++++++++++++
# fill or color box plot by a second group : "supp"
ggviolin(df, "dose", "len", color = "supp",
        palette = c("#00AFBB", "#E7B800"), add = "boxplot")

---

**gradient_color**

**Set Gradient Color**

**Description**

Change gradient color.

- `gradient_color()`: Change gradient color.
- `gradient_fill()`: Change gradient fill.

**Usage**

```r
gradient_color(palette)

gradient_fill(palette)
```
grids

Add Grids to a ggplot

Description

Add grids to ggplot.

Usage

grids(axis = c("xy", "x", "y"), color = "grey92", size = NULL, linetype = NULL)
Arguments

axis  axis for which grid should be added. Allowed values include \texttt{c("xy", "x", "y")}.
color grid line color.
size numeric value specifying grid line size.
linetype line type. An integer (0:8), a name (blank, solid, dashed, dotted, dotdash, long-dash, twodash). See \textit{show_line_types}.

Examples

\begin{verbatim}
# Load data
data("ToothGrowth")

# Basic plot
p <- ggboxplot(ToothGrowth, x = "dose", y = "len")
p

# Add border
p + grids(linetype = "dashed")
\end{verbatim}

\begin{tabular}{ll}
\textbf{npc_to_data_coord} & \textit{Convert NPC to Data Coordinates} \\
\end{tabular}

Description

Convert NPC (Normalized Parent Coordinates) into data coordinates.

Usage

\texttt{npc_to_data_coord(npc, data.ranges)}

Arguments

\begin{itemize}
\item \texttt{npc} a numeric vector. Each value should be in \([0-1]\)
\item \texttt{data.ranges} a numeric vector of length 2 containing the data ranges (minimum and the maximum)
\end{itemize}

Value

a numeric vector representing data coordinates.

See Also

\texttt{as_npc, get_coord}. 
Examples

```r
cpyc_to_data_coord(npc = c(0.2, 0.95), data.ranges = c(1, 20))

as_npc(c("top", "right")) %>%
npc_to_data_coord(data.ranges = c(1, 20))
```

---

`rotate`  
*Rotate a ggplot Horizontally*

Description

Rotate a ggplot to create horizontal plots. Wrapper around `coord_flip`.

Usage

```r
rotate(...)
```

Arguments

...  
other arguments to pass to `coord_flip`.

Examples

```r
# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
p <- ggboxplot(df, x = "dose", y = "len",
               color = "dose", palette = "jco")
p
# Create horizontal plots
p + rotate()
```

---

`rotate_axis_text`  
*Rotate Axes Text*

Description

Rotate the x-axis text (tick mark labels).

- `rotate_x_text()`: Rotate x axis text.
- `rotate_y_text()`: Rotate y axis text.
Usage

rotate_x_text(angle = 90, hjust = NULL, vjust = NULL, ...)

rotate_y_text(angle = 90, hjust = NULL, vjust = NULL, ...)

Arguments

angle numeric value specifying the rotation angle. Default is 90 for vertical x-axis text.

hjust horizontal justification (in [0, 1]).

vjust vertical justification (in [0, 1]).

... other arguments to pass to the function `element_text`.

Examples

# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
p <- ggboxplot(df, x = "dose", y = "len")
p
# Vertical x axis text
p + rotate_x_text()
# Set rotation angle to 45
p + rotate_x_text(45)
p + rotate_y_text(45)

rremove

Remove a ggplot Component

Description

Remove a specific component from a ggplot.

Usage

rremove(object)

Arguments

object character string specifying the plot components. Allowed values include:

- "grid" for both x and y grids
- "x.grid" for x axis grids
- "y.grid" for y axis grids
set_palette

- "axis" for both x and y axes
- "x.axis" for x axis
- "y.axis" for y axis
- "xlab", or "x.title" for x axis label
- "ylab", or "y.title" for y axis label
- "xylab", "xy.title" or "axis.title" for both x and y axis labels
- "x.text" for x axis texts (x axis tick labels)
- "y.text" for y axis texts (y axis tick labels)
- "xy.text" or "axis.text" for both x and y axis texts
- "ticks" for both x and y ticks
- "x.ticks" for x ticks
- "y.ticks" for y ticks
- "legend.title" for the legend title
- "legend" for the legend

Examples

# Load data
data("ToothGrowth")

# Basic plot
p <- ggboxplot(ToothGrowth, x = "dose", y = "len",
               ggtheme = theme_gray())
p

# Remove all grids
p + rremove("grid")

# Remove only x grids
p + rremove("x.grid")

---

set_palette

Set Color Palette

Description

- change_palette(), set_palette(): Change both color and fill palettes.
- color_palette(): change color palette only.
- fill_palette(): change fill palette only.

Usage

set_palette(p, palette)

change_palette(p, palette)
color_palette(palette = NULL, ...)
fill_palette(palette = NULL, ...)

Arguments

p    a ggplot
palette    Color palette. Allowed values include:

• **Grey color palettes**: "grey" or "gray";
• **RColorBrewer palettes**, see `brewer.pal` and details section. Examples of palette names include: "RdBu", "Blues", "Dark2", "Set2", ...;
• **Custom color palettes**. For example, `palette = c("#00AFBB", "#E7B800", "#FC4E07")`;
• **ggsci scientific journal palettes**, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmörty".

...    other arguments passed to ggplot2 `scale_color_xxx()` and `scale_fill_xxx()` functions.

See Also

`get_palette`.

Examples

```r
# Load data
data("ToothGrowth")
df <- ToothGrowth

# Basic plot
p <- ggboxplot(df, x = "dose", y = "len",
               color = "dose")
p

# Change the color palette
set_palette(p, "jco")
```

---

**show_line_types**

**Line types available in R**

Description

Show line types available in R.

Usage

`show_line_types()`
show_point_shapes

Value

a ggplot.

See Also

ggpar and ggline.

Examples

show_line_types()+
theme_minimal()

---

show_point_shapes   Point shapes available in R

Description

Show point shapes available in R.

Usage

show_point_shapes()

Value

a ggplot.

See Also

ggpar and ggline.

Examples

show_point_shapes()+
theme_minimal()
stat_anova_test

Add Anova Test P-values to a GGPlot

Description

Adds automatically one-way and two-way ANOVA test p-values to a ggplot, such as box blots, dot plots and stripcharts.

Usage

stat_anova_test(
  mapping = NULL, 
  data = NULL, 
  method = c("one_way", "one_way_repeated", "two_way", "two_way_repeated", 
    "two_way_mixed"),
  wid = NULL, 
  group.by = NULL, 
  type = NULL, 
  effect.size = "ges",
  error = NULL,
  correction = c("auto", "GG", "HF", "none"),
  label = "{method}, p = {p.format}",
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  step.increase = 0.1,
  p.adjust.method = "holm",
  significance = list(),
  geom = "text",
  position = "identity",
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE,
  parse = FALSE,
  ...
)

Arguments

mapping
Set of aesthetic mappings created by \texttt{aes()}. If specified and \texttt{inherit.aes = TRUE} (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.

data
The data to be displayed in this layer. There are three options: If \texttt{NULL}, the default, the data is inherited from the plot data as specified in the call to \texttt{ggplot()}.  

A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.

A function will be called with a single argument, the plot data. The return value must be a `data.frame`, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

**method**

ANOVA test methods. Possible values are one of c("one_way", "one_way_repeated", "two_way", "two_way_repeated", "two_way_mixed").

**wid**

(factor) column name containing individuals/subjects identifier. Should be unique per individual. Required only for repeated measure tests ("one_way_repeated", "two_way_repeated", "friedman_test", etc).

**group.by**

(optional) character vector specifying the grouping variable; it should be used only for grouped plots. Possible values are:

- "x.var": Group by the x-axis variable and perform the test between legend groups. In other words, the p-value is compute between legend groups at each x position
- "legend.var": Group by the legend variable and perform the test between x-axis groups. In other words, the test is performed between the x-groups for each legend level.

**type**

the type of sums of squares for ANOVA. Allowed values are either 1, 2 or 3. type = 2 is the default because this will yield identical ANOVA results as type = 1 when data are balanced but type = 2 will additionally yield various assumption tests where appropriate. When the data are unbalanced the type = 3 is used by popular commercial softwares including SPSS.

**effect.size**

the effect size to compute and to show in the ANOVA results. Allowed values can be either "ges" (generalized eta squared) or "pes" (partial eta squared) or both. Default is "ges".

**error**

(optional) for a linear model, an lm model object from which the overall error sum of squares and degrees of freedom are to be calculated. Read more in `Anova()` documentation.

**correction**

character. Used only in repeated measures ANOVA test to specify which correction of the degrees of freedom should be reported for the within-subject factors. Possible values are:

- "GG": applies Greenhouse-Geisser correction to all within-subjects factors even if the assumption of sphericity is met (i.e., Mauchly’s test is not significant, p > 0.05).
- "HF": applies Hyunh-Feldt correction to all within-subjects factors even if the assumption of sphericity is met,
- "none": returns the ANOVA table without any correction and
- "auto": apply automatically GG correction to only within-subjects factors violating the sphericity assumption (i.e., Mauchly’s test p-value is significant, p <= 0.05).

**label**

character string specifying label. Can be:

- the column containing the label (e.g.: `label = "p"` or `label = "p.adj"`), where `p` is the p-value. Other possible values are "p.signif", "p.adj.signif", "p.format", "p.adj.format".
• an expression that can be formatted by the `glue()` package. For example, when specifying `label = "Anova, p = \{p\}"`, the expression `{p}` will be replaced by its value.

• a combination of plotmath expressions and glue expressions. You may want some of the statistical parameter in italic; for example: `label = "Anova, italic(p) = \{p\}"`.

• a constant: `label = "as_italic": display statistical parameters in italic; label = "as_detailed": detailed plain text; label = "as_detailed_expression" or label = "as_detailed_italic": detailed plotmath expression. Statistical parameters will be displayed in italic.

`label.x.npc, label.y.npc` can be numeric or character vector of the same length as the number of groups and/or panels. If too short they will be recycled.

- If numeric, value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates".
- If character, allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c( 'bottom', 'top', 'center', 'centre', 'middle') for y-axis.

`label.x, label.y` numeric Coordinates (in data units) to be used for absolute positioning of the label. If too short they will be recycled.

`step.increase` numeric value in with the increase in fraction of total height for every additional comparison to minimize overlap. The step value can be negative to reverse the order of groups.

`p.adjust.method` method for adjusting p values (see `p.adjust`). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don’t want to adjust the p value (not recommended), use `p.adjust.method = "none"`.

`significance` a list of arguments specifying the significance cutpoints and symbols. For example, `significance <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", ",*", "ns")).`

In other words, we use the following convention for symbols indicating statistical significance:

- ns: p > 0.05
- *: p <= 0.05
- **: p <= 0.01
- ***: p <= 0.001
- ****: p <= 0.0001

`geom` The geometric object to use to display the data, either as a `ggproto` `Geom` subclass or as a string naming the geom stripped of the `geom_` prefix (e.g. “point” rather than “geom_point”)
**Computed variables**

- DFn: Degrees of Freedom in the numerator (i.e. DF effect).
- DFd: Degrees of Freedom in the denominator (i.e., DF error).
- ges: Generalized Eta-Squared measure of effect size. Computed only when the option `effect.size = "ges"`.
- pes: Partial Eta-Squared measure of effect size. Computed only when the option `effect.size = "pes"`.
- F: F-value.
- p: p-value.
- p.adj: Adjusted p-values.
- p.signif: P-value significance.
- p.adj.signif: Adjusted p-value significance.
- p.format: Formated p-value.
- p.adj.format: Formated adjusted p-value.
- n: number of samples.

**Examples**

```r
# Data preparation
# Transform `dose` into factor variable
df <- ToothGrowth
df$dose <- as.factor(df$dose)
# Add individuals id
df$id <- rep(1:10, 6)
# Add a random grouping variable
```
```r
set.seed(123)
df$group <- sample(factor(rep(c("grp1", "grp2", "grp3"), 20)))
df$len <- ifelse(df$group == "grp2", df$len + 2, df$len)
df$len <- ifelse(df$group == "grp3", df$len + 7, df$len)
head(df, 3)

# Basic boxplot
# Create a basic boxplot
# Add 5% and 10% space to the plot bottom and the top, respectively
bxp <- ggboxplot(df, x = "dose", y = "len") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))
# Add the p-value to the boxplot
bxp + stat_anova_test()

## Not run:
# Change the label position
# Using coordinates in data units
bxp + stat_anova_test(label.x = "1", label.y = 10, hjust = 0)
## End(Not run)

# Format the p-value differently
custom_p_format <- function(p) {
  rstatix::p_format(p, accuracy = 0.0001, digits = 3, leading.zero = FALSE)
}
bxp + stat_anova_test(
  label = "Anova, italic(p) = {custom_p_format(p)}{p.signif}"
)
# Show a detailed label in italic
bxp + stat_anova_test(label = "as_detailed_italic")

# Faceted plots
# Create a ggplot facet
bxp <- ggboxplot(df, x = "dose", y = "len", facet.by = "supp") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))
# Add p-values
bxp + stat_anova_test()

# Grouped plots
bxp2 <- ggboxplot(df, x = "group", y = "len", color = "dose", palette = "npg")

# For each x-position, computes tests between legend groups
bxp2 + stat_anova_test(aes(group = dose), label = "p = {p.format}{p.signif}")
# For each legend group, computes tests between x variable groups
```
```r
bxp2 + stat_anova_test(aes(group = dose, color = dose), group.by = "legend.var")

# Not run:
# Two-way ANOVA: Independent measures
# Visualization: box plots with p-values
# Two-way interaction p-values between x and legend (group) variables
bxp3 <- ggboxplot(
  df, x = "supp", y = "len",
  color = "dose", palette = "jco"
)
 bxp3 + stat_anova_test(aes(group = dose), method = "two_way")

# One-way repeated measures ANOVA
# df$id <- as.factor(c(rep(1:10, 3), rep(11:20, 3)))
ggboxplot(df, x = "dose", y = "len") +
  stat_anova_test(method = "one_way_repeated", wid = "id")

# Two-way repeated measures ANOVA
# df$id <- as.factor(rep(1:10, 6))
ggboxplot(df, x = "dose", y = "len", color = "supp", palette = "jco") +
  stat_anova_test(aes(group = supp), method = "two_way_repeated", wid = "id")

# Grouped one-way repeated measures ANOVA
ggboxplot(df, x = "dose", y = "len", color = "supp", palette = "jco") +
  stat_anova_test(aes(group = supp, color = supp),
  method = "one_way_repeated", wid = "id", group.by = "legend.var")
```

---

## Description

add brackets with label annotation to a ggplot. Helpers for adding p-value or significance levels to a plot.

## Usage

```r
stat_bracket(
  mapping = NULL,
  data = NULL,
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
)```

Arguments

mapping Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.
The data to be displayed in this layer. There are three options:

- If NULL, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
- A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
- A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use `position_jitter`), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

A logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. `borders()`.

A character vector with alternative label, if not null test is ignored.

The label type. Can be one of "text" and "expression" (for parsing `plotmath` expression).

A numeric vector with the y positions of the brackets.

A numeric vector with the positions of the left sides of the brackets.

A numeric vector with the positions of the right sides of the brackets.

A numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.

A variable name for grouping brackets before adding `step.increase`. Useful to group bracket by facet panel.

A numeric vector with the fraction of total height that the bar goes down to indicate the precise column.

Vertical adjustment to nudge brackets by. Useful to move up or move down the bracket. If positive value, brackets will be moved up; if negative value, brackets are moved down.

A small numeric value in [0-1] for shortening the width of the bracket.

Change the width of the lines of the bracket.

Change the size of the label text.

Change the font used for the text.

Move the text up or down relative to the bracket.
... other arguments passed on to layer. These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat.

stat The statistical transformation to use on the data for this layer, either as a ggproto Geom subclass or as a string naming the stat stripped of the stat_ prefix (e.g. "count" rather than "stat_count")

coord.flip logical. If TRUE, flip x and y coordinates so that horizontal becomes vertical, and vertical, horizontal. When adding the p-values to a horizontal ggplot (generated using coord_flip()), you need to specify the option coord.flip = TRUE.

Examples

def <- ToothGrowth
def$dose <- factor(df$dose)

# Add bracket with labels
ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    xmin = "0.5", xmax = "1", y.position = 30,
    label = "t-test, p < 0.05"
  )

# Customize bracket tip.length

ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    xmin = "0.5", xmax = "1", y.position = 30,
    label = "t-test, p < 0.05", tip.length = c(0.2, 0.02)
  )

# Using plotmath expression

ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    xmin = "0.5", xmax = "1", y.position = 30,
    label = "list(~italic(p)<=0.001)", type = "expression",
    tip.length = c(0.2, 0.02)
  )

# Specify multiple brackets manually

ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    xmin = c("0.5", "1"), xmax = c("1", "2"),
    y.position = c(30, 35), label = c("***", "**"),
    tip.length = 0.01
  )

# Compute statistical tests and add p-values
stat.test <- compare_means(len ~ dose, ToothGrowth, method = "t.test")
ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    aes(xmin = group1, xmax = group2, label = signif(p, 2)),
    data = stat.test, y.position = 35
  )
# Increase step length between brackets
ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    aes(xmin = group1, xmax = group2, label = signif(p, 2)),
    data = stat.test, y.position = 35, step.increase = 0.1
  )

# Or specify the positions of each comparison
ggboxplot(df, x = "dose", y = "len") +
  geom_bracket(
    aes(xmin = group1, xmax = group2, label = signif(p, 2)),
    data = stat.test, y.position = c(32, 35, 38)
  )

stat_central_tendency  Add Central Tendency Measures to a GGPlot

Description
Add central tendency measures (mean, median, mode) to density and histogram plots created using ggplots.

Note that, normally, the mode is used for categorical data where we wish to know which is the most common category. Therefore, we can have two or more values that share the highest frequency. This might be problematic for continuous variable.

For continuous variable, we can consider using mean or median as the measures of the central tendency.

Usage
stat_central_tendency(
  mapping = NULL,
  data = NULL,
  geom = c("line", "point"),
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  type = c("mean", "median", "mode"),
  ...
)

Arguments

mapping  Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
The data to be displayed in this layer. There are three options:
If NULL, the default, the data is inherited from the plot data as specified in the call to \texttt{ggplot()}
A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See \texttt{fortify()} for which variables will be created.
A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. \texttt{~ head(.x, 10)}).

The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_prefix (e.g. \texttt{"point"} rather than \texttt{"geom_point"})

Position adjustment, either as a string naming the adjustment (e.g. \texttt{"jitter"} to use \texttt{position_jitter}), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.
logical. Should this layer be included in the legends? \texttt{NA}, the default, includes if any aesthetics are mapped. \texttt{FALSE} never includes, and \texttt{TRUE} always includes. It can also be a named logical vector to finely select the aesthetics to display.
If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. \texttt{borders()}.
the type of central tendency measure to be used. Possible values include: \texttt{"mean"}, \texttt{"median"}, \texttt{"mode"}.
other arguments to pass to \texttt{geom_line}.

### Examples

```r
# Simple density plot
data(mtcars)
ggdensity(mtcars, x = "mpg", fill = "red") +
  scale_x_continuous(limits = c(-1, 50)) +
  stat_central_tendency(type = "mean", linetype = "dashed")

# Color by groups
data(iris)
ggdensity(iris, "Sepal.Length", color = "Species") +
  stat_central_tendency(aes(color = Species), type = "median", linetype = 2)

# Use geom = "point" for central tendency
data(iris)
ggdensity(iris, "Sepal.Length", color = "Species") +
  stat_central_tendency(
```

### See Also

\texttt{ggdensity}
# Facet

ggdensity(iris, "Sepal.Length", facet.by = "Species") +
stat_central_tendency(type = "mean", color = "red", linetype = 2) +
stat_central_tendency(type = "median", color = "blue", linetype = 2)

---

**Description**

Plot convex hull of a set of points.

**Usage**

```r
stat_chull(
  mapping = NULL,
  data = NULL,
  geom = "path",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

**Arguments**

- **mapping**: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.

- **data**: The data to be displayed in this layer. There are three options:
  - If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
  - A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
  - A function will be called with a single argument, the plot data. The return value must be a `data.frame`, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

- **geom**: The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_ prefix (e.g. "point" rather than "geom_point")
stat_compare_means

Description

Add mean comparison p-values to a ggplot, such as box blots, dot plots and stripcharts.

Usage

stat_compare_means(
  mapping = NULL,
  data = NULL,
  method = NULL,
...
```
paired = FALSE,
method.args = list(),
ref.group = NULL,
comparisons = NULL,
hide.ns = FALSE,
label.sep = "",
label = NULL,
label.x.npc = "left",
label.y.npc = "top",
label.x = NULL,
label.y = NULL,
vjust = 0,
tip.length = 0.03,
bracket.size = 0.3,
step.increase = 0,
symnum.args = list(),
geom = "text",
position = "identity",
na.rm = FALSE,
show.legend = NA,
inherit.aes = TRUE,
...
```

**Arguments**

**mapping**  
Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.

**data**  
The data to be displayed in this layer. There are three options:  
If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.  
A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.  
A function will be called with a single argument, the plot data. The return value must be a `data.frame`, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

**method**  
a character string indicating which method to be used for comparing means.

**paired**  
a logical indicating whether you want a paired test. Used only in `t.test` and in `wilcox.test`.

**method.args**  
a list of additional arguments used for the test method. For example one might use `method.args = list(alternative = "greater")` for wilcoxon test.

**ref.group**  
a character string specifying the reference group. If specified, for a given grouping variable, each of the group levels will be compared to the reference group (i.e. control group).
ref.group can be also ".all.". In this case, each of the grouping variable
levels is compared to all (i.e. basemean).

comparisons A list of length-2 vectors. The entries in the vector are either the names of 2
values on the x-axis or the 2 integers that correspond to the index of the groups
of interest, to be compared.

hide.ns logical value. If TRUE, hide ns symbol when displaying significance levels.

label.sep a character string to separate the terms. Default is ",", to separate the correlation
coefficient and the p.value.

label character string specifying label type. Allowed values include "p.signif" (shows
the significance levels), "p.format" (shows the formatted p value).

label.x.npc, label.y.npc can be numeric or character vector of the same length as the number of groups
and/or panels. If too short they will be recycled.

• If numeric, value should be between 0 and 1. Coordinates to be used for
positioning the label, expressed in "normalized parent coordinates".
• If character, allowed values include: i) one of c('right', 'left', 'center',
'centre', 'middle') for x-axis; ii) and one of c('bottom', 'top', 'center',
'centre', 'middle') for y-axis.

label.x, label.y numeric Coordinates (in data units) to be used for absolute positioning of the
label. If too short they will be recycled.

vjust move the text up or down relative to the bracket.

tip.length numeric vector with the fraction of total height that the bar goes down to indicate
the precise column. Default is 0.03. Can be of same length as the number of comparisons to adjust specifically the tip length of each comparison. For example
tip.length = c(0.01, 0.03).

If too short they will be recycled.

bracket.size Width of the lines of the bracket.

step.increase numeric vector with the increase in fraction of total height for every additional
comparison to minimize overlap.

symnum.args a list of arguments to pass to the function symnum for symbolic number coding
of p-values. For example, symnum.args <- list(cutpoints = c(0, 0.0001,
0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", "+", "ns").

In other words, we use the following convention for symbols indicating statistical significance:

• ns: p > 0.05
• *: p <= 0.05
• **: p <= 0.01
• ***: p <= 0.001
• ****: p <= 0.0001

gem The geometric object to use to display the data, either as a ggproto Geom sub-
class or as a string naming the geom stripped of the geom_ prefix (e.g. "point"
rather than "geom_point")
stat_compare_means

position

Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

na.rm

If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

show.legend

logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

See Also

compare_means

Examples

# Load data
data("ToothGrowth")
head(ToothGrowth)

# Two independent groups
#:::::::::::::::::::::::::::::::::::::::::::::::::
p <- ggboxplot(ToothGrowth, x = "supp", y = "len",
               color = "supp", palette = "npg", add = "jitter")

# Add p-value
p + stat_compare_means()
# Change method
p + stat_compare_means(method = "t.test")

# Paired samples
#:::::::::::::::::::::::::::::::::::::::::::::::::
ggpaired(ToothGrowth, x = "supp", y = "len",
         color = "supp", line.color = "gray", line.size = 0.4,
         palette = "npg")+
stat_compare_means(paired = TRUE)

# More than two groups
#:::::::::::::::::::::::::::::::::::::::::::::::::
# Pairwise comparisons: Specify the comparisons you want
my_comparisons <- list( c("0.5", "1"), c("1", "2"), c("0.5", "2") )
ggboxplot(ToothGrowth, x = "dose", y = "len",
          color = "dose", palette = "npg")+
# Add pairwise comparisons p-value
stat_compare_means(comparisons = my_comparisons, label.y = c(29, 35, 40)) +
stat_compare_means(label.y = 45)  # Add global Anova p-value
# Multiple pairwise test against a reference group
ggboxplot(ToothGrowth, x = "dose", y = "len",
    color = "dose", palette = "npg")+
stat_compare_means(method = "anova", label.y = 40)+ # Add global p-value
stat_compare_means(aes(label = after_stat(p.signif)),
    method = "t.test", ref.group = "0.5")

# Multiple grouping variables
#:::::::::::::::::::::::::::::::::::::::::::::::::
# Box plot facetted by "dose"
p <- ggboxplot(ToothGrowth, x = "supp", y = "len",
    color = "supp", palette = "npg",
    add = "jitter",
    facet.by = "dose", short.panel.labs = FALSE)
# Use only p.format as label. Remove method name.
p + stat_compare_means(
    aes(label = paste0("p = ", after_stat(p.format))))

---

**stat_conf_ellipse**  
*Plot confidence ellipses.*

### Description
Plot confidence ellipses around barycenters. The method for computing confidence ellipses has been modified from FactoMineR::coord.ellipse().

### Usage
```r
stat_conf_ellipse(
  mapping = NULL,
  data = NULL,
  geom = "path",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  level = 0.95,
  npoint = 100,
  bary = TRUE,
  ...
)
```

### Arguments
- **mapping**: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
The data to be displayed in this layer. There are three options:
If NULL, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_ prefix (e.g. "point" rather than "geom_point")

Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.

logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. `borders()`

confidence level used to construct the ellipses. By default, 0.95.

number of points used to draw the ellipses.

logical value. If TRUE, the coordinates of the ellipse around the barycentre of individuals are calculated.

Other arguments passed on to `layer()`. These are often aesthetics, used to set an aesthetic to a fixed value, like `colour = "red"` or `size = 3`. They may also be parameters to the paired geom/stat.

See Also

`stat_conf_ellipse`

Examples

```r
# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)

# scatter plot with confidence ellipses
ggscatter(df, x = "wt", y = "mpg", color = "cyl")+
  stat_conf_ellipse(aes(color = cyl))

# scatter plot with confidence ellipses
ggscatter(df, x = "wt", y = "mpg", color = "cyl")+
  stat_conf_ellipse(aes(color = cyl))
```

Add Correlation Coefficients with P-values to a Scatter Plot

Description
Add correlation coefficients with p-values to a scatter plot. Can be also used to add ‘R2’.

Usage
```r
stat_cor(
  mapping = NULL,
  data = NULL,
  method = "pearson",
  alternative = "two.sided",
  cor.coef.name = c("R", "rho", "tau"),
  label.sep = ", ",
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  output.type = "expression",
  digits = 2,
  r.digits = digits,
  p.digits = digits,
  r.accuracy = NULL,
  p.accuracy = NULL,
  geom = "text",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments
- `mapping`: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.
- `data`: The data to be displayed in this layer. There are three options:
  - If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
  - If `data = NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
A data frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.

A function will be called with a single argument, the plot data. The return value must be a data frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

**method**

A character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman".

**alternative**

A character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.

**cor.coef.name**

Character. Can be one of "R" (pearson coef), "rho" (spearman coef) and "tau" (kendall coef). Uppercase and lowercase are allowed.

**label.sep**

A character string to separate the terms. Default is ",", to separate the correlation coefficient and the p.value.

**label.x.npc, label.y.npc**

Can be numeric or character vector of the same length as the number of groups and/or panels. If too short they will be recycled.

- If numeric, value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates".
- If character, allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c('bottom', 'top', 'center', 'centre', 'middle') for y-axis.

If too short they will be recycled.

**label.x, label.y**

Numeric Coordinates (in data units) to be used for absolute positioning of the label. If too short they will be recycled.

**output.type**

Character One of "expression", "latex", "tex" or "text".

**digits, r.digits, p.digits**

Integer indicating the number of decimal places (round) or significant digits (signif) to be used for the correlation coefficient and the p-value, respectively.

**r.accuracy**

A real value specifying the number of decimal places of precision for the correlation coefficient. Default is NULL. Use (e.g.) 0.01 to show 2 decimal places of precision. If specified, then r.digits is ignored.

**p.accuracy**

A real value specifying the number of decimal places of precision for the p-value. Default is NULL. Use (e.g.) 0.0001 to show 4 decimal places of precision. If specified, then p.digits is ignored.

**geom**

The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_prefix (e.g. "point" rather than "geom_point")

**position**

Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

**na.rm**

If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.
show.legend logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

Computed variables

r correlation coefficient

rr correlation coefficient squared

r.label formatted label for the correlation coefficient

rr.label formatted label for the squared correlation coefficient

p.label label for the p-value

label default label displayed by stat_cor()

See Also

ggscatter

Examples

# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)

# Scatter plot with correlation coefficient
#:::::::::::::::::::::::::::::::::::::::::::::::::
sp <- ggscatter(df, x = "wt", y = "mpg",
         add = "reg.line", # Add regression line
         add.params = list(color = "blue", fill = "lightgray"), # Customize reg. line
         conf.int = TRUE # Add confidence interval
)

# Add correlation coefficient
sp + stat_cor(method = "pearson", label.x = 3, label.y = 30)

# Specify the number of decimal places of precision for p and r
# Using 3 decimal places for the p-value and
# 2 decimal places for the correlation coefficient (r)
sp + stat_cor(p.accuracy = 0.001, r.accuracy = 0.01)

# Show only the r.label but not the p.label
sp + stat_cor(aes(label = ..r.label..), label.x = 3)

# Use R2 instead of R
ggscatter(df, x = "wt", y = "mpg", add = "reg.line") +
stat_cor(}
stat_friedman_test

```
aes(label = paste(..rr.label.., ..p.label.., sep = "-", "")),
  label.x = 3
}

# Color by groups and facet
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::
sp <- ggscatter(df, x = "wt", y = "mpg",
  color = "cyl", palette = "jco",
  add = "reg.line", conf.int = TRUE)
sp + stat_cor(aes(color = cyl), label.x = 3)
```

---

**stat_friedman_test**  
*Add Friedman Test P-values to a GGPlot*

**Description**

Add automatically Friedman test p-values to a ggplot, such as box blots, dot plots and stripcharts.

**Usage**

```
stat_friedman_test(
  mapping = NULL,
  data = NULL,
  wid = NULL,
  group.by = NULL,
  label = "{method}, p = {p.format}",
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  step.increase = 0.1,
  p.adjust.method = "holm",
  significance = list(),
  geom = "text",
  position = "identity",
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE,
  parse = FALSE,
  ...
)
```

**Arguments**

- **mapping**: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.
The data to be displayed in this layer. There are three options:
If NULL, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).
(factor) column name containing individuals/subjects identifier. Should be unique per individual. Required only for repeated measure tests ("one_way_repeated", "two_way_repeated", "friedman_test", etc).
(optional) character vector specifying the grouping variable; it should be used only for grouped plots. Possible values are:
- "x.var": Group by the x-axis variable and perform the test between legend groups. In other words, the p-value is compute between legend groups at each x position
- "legend.var": Group by the legend variable and perform the test between x-axis groups. In other words, the test is performed between the x-groups for each legend level.
the column containing the label (e.g.: `label = "p"` or `label = "p.adj"`), where p is the p-value. Can be also an expression that can be formatted by the `glue()` package. For example, when specifying `label = "t-test, p = {p}"`, the expression `{p}` will be replaced by its value.
can be numeric or character vector of the same length as the number of groups and/or panels. If too short they will be recycled.
- If numeric, value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates".
- If character, allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c('bottom', 'top', 'center', 'centre', 'middle') for y-axis.
numeric Coordinates (in data units) to be used for absolute positioning of the label. If too short they will be recycled.
numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.
method for adjusting p values (see `p.adjust`). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don't want to adjust the p value (not recommended), use `p.adjust.method = "none"`.
a list of arguments specifying the significance cutpoints and symbols. For example, `significance <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", ", ns"))`. 
In other words, we use the following convention for symbols indicating statistical significance:

- **ns**: \( p > 0.05 \)
- **``**: \( p \leq 0.05 \)
- ****: \( p \leq 0.01 \)
- ***: \( p \leq 0.001 \)
- ****: \( p \leq 0.0001 \)

**geom**  
The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_ prefix (e.g. “point” rather than “geom_point”)

**position**  
Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

**na.rm**  
If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

**show.legend**  
Logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

**inherit.aes**  
If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

**parse**  
If TRUE, the labels will be parsed into expressions and displayed as described in ?plotmath.

...  
other arguments passed to the function geom_bracket() or geom_text()

**Computed variables**

- statistic: the value of the test statistic (Chi-squared).
- df: the degrees of freedom of the approximate chi-squared distribution of the test statistic.
- p: p-value.
- p.adjust: Adjusted p-values.
- p.signif: P-value significance.
- p.adjust.signif: Adjusted p-value significance.
- p.format: Formated p-value.
- p.adjust.format: Formated adjusted p-value.
- n: number of samples.

**Examples**

```r
# Data preparation
#%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
# Transform `dose` into factor variable
df <- ToothGrowth
df$dose <- as.factor(df$dose)
```
df$id <- as.factor(c(rep(1:10, 3), rep(11:20, 3)))
# Add a random grouping variable
set.seed(123)
df$group <- sample(factor(rep(c("grp1", "grp2", "grp3"), 20)))
df$len <- ifelse(df$group == "grp2", df$len+2, df$len)
df$len <- ifelse(df$group == "grp3", df$len+7, df$len)
head(df, 3)

# Basic boxplot
#%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
# Create a basic boxplot
# Add 5% and 10% space to the plot bottom and the top, respectively
bxp <- ggboxplot(df, x = "dose", y = "len") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))
# Add the p-value to the boxplot
bxp + stat_friedman_test(aes(wid = id))
# Change the label position
# Using coordinates in data units
bxp + stat_friedman_test(aes(wid = id), label.x = "1", label.y = 10, hjust = 0)
# Format the p-value differently
custom_p_format <- function(p) {
  rstatix::p_format(p, accuracy = 0.0001, digits = 3, leading.zero = FALSE)
}
bxp + stat_friedman_test(
  aes(wid = id),
  label = "Friedman test, italic(p) = {custom_p_format(p)}{p.signif}")
# Show a detailed label in italic
bxp + stat_friedman_test(aes(wid = id), label = "as_detailed_italic")

# Faceted plots
#%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
# Create a ggplot facet
df$id <- rep(1:10,6)
bxp <- ggboxplot(df, x = "dose", y = "len", facet.by = "supp") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))
# Add p-values
bxp + stat_friedman_test(aes(wid = id))

# Grouped plots
#%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
df$id <- rep(1:10,6)
bxp <- ggboxplot(df, x = "dose", y = "len", color = "supp", palette = "jco")
# For each legend group, computes tests within x variable groups
bxp + stat_friedman_test(aes(wid = id, group = supp, color = supp), within = "x")
stat_kruskal_test

# For each x-position, computes tests within legend variable groups
bxp + stat_friedman_test(
  aes(wid = id, group = supp, color = supp),
  within = "group", label = "p = {p.format}"
)

---

stat_kruskal_test  Add Kruskal-Wallis Test P-values to a GGPlot

Description

Add Kruskal-Wallis test p-values to a ggplot, such as box blots, dot plots and stripcharts.

Usage

stat_kruskal_test(
  mapping = NULL,
  data = NULL,
  group.by = NULL,
  label = "{method}, p = {p.format}",
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  step.increase = 0.1,
  p.adjust.method = "holm",
  significance = list(),
  geom = "text",
  position = "identity",
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE,
  parse = FALSE,
...
)

Arguments

mapping  Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.

data  The data to be displayed in this layer. There are three options:
If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot().
A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.

A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

`group.by` (optional) character vector specifying the grouping variable; it should be used only for grouped plots. Possible values are:
- "x.var": Group by the x-axis variable and perform the test between legend groups. In other words, the p-value is computed between legend groups at each x position.
- "legend.var": Group by the legend variable and perform the test between x-axis groups. In other words, the test is performed between the x-groups for each legend level.

`label` the column containing the label (e.g.: `label = "p"` or `label = "p.adj"`), where p is the p-value. Can be an expression that can be formatted by the `glue()` package. For example, when specifying `label = "t-test, p = {p}"`, the expression `{p}` will be replaced by its value.

`label.x.npc, label.y.npc` can be numeric or character vector of the same length as the number of groups and/or panels. If too short, they will be recycled.
- If numeric, value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates".
- If character, allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c( 'bottom', 'top', 'center', 'centre', 'middle') for y-axis.

`label.x, label.y` numeric Coordinates (in data units) to be used for absolute positioning of the label. If too short, they will be recycled.

`step.increase` numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.

`p.adjust.method` method for adjusting p values (see `p.adjust`). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don’t want to adjust the p value (not recommended), use `p.adjust.method = "none"`.

`significance` a list of arguments specifying the significance cutpoints and symbols. For example, `significance <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", ",", "ns"))`. In other words, we use the following convention for symbols indicating statistical significance:
- ns: p > 0.05
- *: p <= 0.05
- **: p <= 0.01
• ***: p <= 0.001  
• ****: p <= 0.0001

geom  The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_ prefix (e.g. "point" rather than "geom_point")

position  Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

na.rm  If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

show.legend  logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes  If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

parse  If TRUE, the labels will be parsed into expressions and displayed as described in plotmath.

... other arguments passed to the function geom_bracket() or geom_text()

Computed variables

• statistic: the Kruskal-Wallis rank sum chi-squared statistic used to compute the p-value.
• p: p-value.
• p.adj: Adjusted p-values.
• p.signif: P-value significance.
• p.adj.signif: Adjusted p-value significance.
• p.format: Formated p-value.
• p.adj.format: Formated adjusted p-value.
• n: number of samples.

Examples

# Data preparation
# Transform `dose` into factor variable
df <- ToothGrowth
df$dose <- as.factor(df$dose)
# Add a random grouping variable
set.seed(123)
df$group <- sample(factor(rep(c("grp1", "grp2", "grp3"), 20)))
df$len <- ifelse(df$group == "grp2", df$len+2, df$len)
df$len <- ifelse(df$group == "grp3", df$len+7, df$len)
head(df, 3)
# Basic boxplot

```r
# Create a basic boxplot
# Add 5% and 10% space to the plot bottom and the top, respectively
bxp <- ggboxplot(df, x = "dose", y = "len") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))

# Add the p-value to the boxplot
bxp + stat_kruskal_test()

# Change the label position
# Using coordinates in data units
bxp + stat_kruskal_test(label.x = "1", label.y = 10, hjust = 0)

# Format the p-value differently
custom_p_format <- function(p) {
  rstatix::p_format(p, accuracy = 0.0001, digits = 3, leading.zero = FALSE)
}
bxp + stat_kruskal_test(
  label = "Kruskal-Wallis, italic(p) = {custom_p_format(p)}{p.signif}"
)

# Show a detailed label in italic
bxp + stat_kruskal_test(label = "as_detailed_italic")
```

# Faceted plots

```r
# Create a ggplot facet
bxp <- ggboxplot(df, x = "dose", y = "len", facet.by = "supp") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))

# Add p-values
bxp + stat_kruskal_test()
```

# Grouped plots

```r
bxp2 <- ggboxplot(df, x = "group", y = "len", color = "dose", palette = "npg")

# For each x-position, computes tests between legend groups
bxp2 + stat_kruskal_test(aes(group = dose), label = "p = {p.format}{p.signif}"
)

# For each legend group, computes tests between x variable groups
bxp2 + stat_kruskal_test(aes(group = dose, color = dose), group.by = "legend.var")
```

---

**stat_mean**

*Draw group mean points*
Description

Draw the mean point of each group.

Usage

```r
stat_mean(
  mapping = NULL,
  data = NULL,
  geom = "point",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments

- **mapping**: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.

- **data**: The data to be displayed in this layer. There are three options:
  - If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
  - A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
  - A function will be called with a single argument, the plot data. The return value must be a `data.frame`, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

- **geom**: The geometric object to use to display the data, either as a `ggproto` GeoM subclass or as a string naming the geom stripped of the `geom_` prefix (e.g. "point" rather than "geom_point")

- **position**: Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use `position_jitter`), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

- **na.rm**: If `FALSE` (the default), removes missing values with a warning. If `TRUE` silently removes missing values.

- **show.legend**: Logical. Should this layer be included in the legends? `NA`, the default, includes if any aesthetics are mapped. `FALSE` never includes, and `TRUE` always includes. It can also be a named logical vector to finely select the aesthetics to display.

- **inherit.aes**: If `FALSE`, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. `borders()`.

- **...**: other arguments to pass to `geom_point`.


stat_overlay_normal_density

Overlay Normal Density Plot

Description

Overlay normal density plot (with the same mean and SD) to the density distribution of 'x'. This is useful for visually inspecting the degree of deviance from normality.

Usage

stat_overlay_normal_density(
  mapping = NULL,
  data = NULL,
  geom = "line",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)

Arguments

mapping Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.

data The data to be displayed in this layer. There are three options:

Examples

# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)

# Scatter plot with ellipses and group mean points
ggscatter(df, x = "wt", y = "mpg",
    color = "cyl", shape = "cyl", ellipse = TRUE)+
  stat_mean(aes(color = cyl, shape = cyl), size = 4)
stat_overlay_normal_density

A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.

A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

geom

The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_prefix (e.g. "point" rather than "geom_point")

position

Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

na.rm

If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

show.legend

logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

...

other arguments to pass to geom_line.

See Also

ggdensity

Examples

# Simple density plot
data("mtcars")
ggdensity(mtcars, x = "mpg", fill = "red") +
  scale_x_continuous(limits = c(-1, 50)) +
  stat_overlay_normal_density(color = "red", linetype = "dashed")

# Color by groups
data(iris)
ggdensity(iris, "Sepal.Length", color = "Species") +
  stat_overlay_normal_density(aes(color = Species), linetype = "dashed")

# Facet
ggdensity(iris, "Sepal.Length", facet.by = "Species") +
  stat_overlay_normal_density(color = "red", linetype = "dashed")
Add manually p-values to a ggplot, such as box blots, dot plots and stripcharts. Frequently asked questions are available on Datanovia ggpubr FAQ page, for example:

- How to Add P-Values onto Basic GGPlots
- How to Add Adjusted P-values to a Multi-Panel GGPlot
- How to Add P-values to GGPlot Facets
- How to Add P-Values Generated Elsewhere to a GGPlot
- How to Add P-Values onto a Grouped GGPlot using the GGPUBR R Package
- How to Create Stacked Bar Plots with Error Bars and P-values
- How to Add P-Values onto Horizontal GGPlots

Usage

```r
stat_pvalue_manual(
  data,
  label = NULL,
  y.position = "y.position",
  xmin = "group1",
  xmax = "group2",
  x = NULL,
  size = 3.88,
  label.size = size,
  bracket.size = 0.3,
  bracket.nudge.y = 0,
  bracket.shorthen = 0,
  color = "black",
  linetype = 1,
  tip.length = 0.03,
  remove.bracket = FALSE,
  step.increase = 0,
  step.group.by = NULL,
  hide.ns = FALSE,
  vjust = 0,
  coord.flip = FALSE,
  position = "identity",
  ...
)
```
Arguments

data  a data frame containing statistical test results. The expected default format should contain the following columns: group1 | group2 | p | y.position | etc. group1 and group2 are the groups that have been compared. p is the resulting p-value. y.position is the y coordinates of the p-values in the plot.

label  the column containing the label (e.g.: label = "p" or label = "p.adj"), where p is the p-value. Can be also an expression that can be formatted by the glue() package. For example, when specifying label = "t-test, p = {p}"; the expression {p} will be replaced by its value.

y.position  column containing the coordinates (in data units) to be used for absolute positioning of the label. Default value is "y.position". Can be also a numeric vector.

xmin  column containing the position of the left sides of the brackets. Default value is "group1".

xmax  (optional) column containing the position of the right sides of the brackets. Default value is "group2". If NULL, the p-values are plotted as a simple text.

x  x position of the p-value. Should be used only when you want plot the p-value as text (without brackets).

size, label.size  size of label text.

bracket.size  Width of the lines of the bracket.

bracket.nudge.y  Vertical adjustment to nudge brackets by. Useful to move up or move down the bracket. If positive value, brackets will be moved up; if negative value, brackets are moved down.

bracket.shorten  a small numeric value in [0-1] for shortening the width of bracket.

color  text and line color. Can be variable name in the data for coloring by groups.

linetype  linetype. Can be variable name in the data for changing linetype by groups.

tip.length  numeric vector with the fraction of total height that the bar goes down to indicate the precise column. Default is 0.03.

remove.bracket  logical, if TRUE, brackets are removed from the plot. Considered only in the situation, where comparisons are performed against reference group or against "all".

step.increase  numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.

step.group.by  a variable name for grouping brackets before adding step.increase. Useful to group bracket by facet panel.

hide.ns  can be logical value or a character vector.

  • Case when logical value. If TRUE, hide ns symbol when displaying significance levels. Filter is done by checking the column p.adj.signif, p.signif, p.adj and p.
  • Case when character value. Possible values are "p" or "p.adj", for filtering out non significant.
vjust
move the text up or down relative to the bracket. Can be also a column name available in the data.

coord.flip
logical. If TRUE, flip x and y coordinates so that horizontal becomes vertical, and vertical, horizontal. When adding the p-values to a horizontal ggplot (generated using `coord_flip()`), you need to specify the option `coord.flip = TRUE`.

position
position adjustment, either as a string, or the result of a call to a position adjustment function.

... other arguments passed to the function `geom_bracket()` or `geom_text()`

See Also
`stat_compare_means`

Examples

```r
# T-test
stat.test <- compare_means(
  len ~ dose, data = ToothGrowth,
  method = "t.test"
)
stat.test

# Create a simple box plot
p <- ggboxplot(ToothGrowth, x = "dose", y = "len")
p

# Perform a t-test between groups
stat.test <- compare_means(
  len ~ dose, data = ToothGrowth,
  method = "t.test"
)
stat.test

# Add manually p-values from stat.test data
# First specify the y.position of each comparison
stat.test <- stat.test %>%
  mutate(y.position = c(29, 35, 39))
p + stat_pvalue_manual(stat.test, label = "p.adj")

# Customize the label with glue expression
# (https://github.com/tidyverse/glue)
p + stat_pvalue_manual(stat.test, label = "p = {p.adj}")

# Grouped bar plots
ToothGrowth$dose <- as.factor(ToothGrowth$dose)
# Comparisons against reference
stat.test <- compare_means(
  len ~ dose, data = ToothGrowth, group.by = "supp",
...```
method = "t.test", ref.group = "0.5"
)
stat.test
# Plot
bp <- ggbarplot(ToothGrowth, x = "supp", y = "len",
                 fill = "dose", palette = "jco",
                 add = "mean_sd", add.params = list(group = "dose"),
                 position = position_dodge(0.8))
bp + stat_pvalue_manual(
    stat.test, x = "supp", y.position = 33,
    label = "p.signif",
    position = position_dodge(0.8)
)

---

stat_pwc Add Pairwise Comparisons P-values to a GGPlot

**Description**

add pairwise comparison p-values to a ggplot such as box plots, dot plots and stripcharts.

**Usage**

```r
stat_pwc(
  mapping = NULL,
  data = NULL,
  method = "wilcox_test",
  method.args = list(),
  ref.group = NULL,
  label = "p.format",
  y.position = NULL,
  group.by = NULL,
  dodge = 0.8,
  bracket.nudge.y = 0.05,
  bracket.shorten = 0,
  bracket.group.by = c("x.var", "legend.var"),
  step.increase = 0.12,
  tip.length = 0.03,
  size = 0.3,
  label.size = 3.88,
  family = "",
  vjust = 0,
  hjust = 0.5,
  p.adjust.method = "holm",
  p.adjust.by = c("group", "panel"),
  symnum.args = list(),
  hide.ns = FALSE,
)```
arguments

remove.bracket = FALSE,
position = "identity",
na.rm = FALSE,
show.legend = NA,
inherit.aes = TRUE,
parse = FALSE,
...
)

geom_pwc(
  mapping = NULL,
  data = NULL,
  stat = "pwc",
  method = "wilcox_test",
  method.args = list(),
  ref.group = NULL,
  label = "p.format",
  y.position = NULL,
  group.by = NULL,
  dodge = 0.8,
  stack = FALSE,
  step.increase = 0.12,
  tip.length = 0.03,
  bracket.nudge.y = 0.05,
  bracket.shorten = 0,
  bracket.group.by = c("x.var", "legend.var"),
  size = 0.3,
  label.size = 3.88,
  family = "",
  vjust = 0,
  hjust = 0.5,
  p.adjust.method = "holm",
  p.adjust.by = c("group", "panel"),
  symnum.args = list(),
  hide.ns = FALSE,
  remove.bracket = FALSE,
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  parse = FALSE,
...
)

Arguments

mapping Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
The data to be displayed in this layer. There are three options:

- If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.

- A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.

- A function will be called with a single argument, the plot data. The return value must be a `data.frame`, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

A character string indicating which method to be used for pairwise comparisons. Default is "wilcox_test". Allowed methods include pairwise comparisons methods implemented in the `rstatix` R package. These methods are: "wilcox_test", "t_test", "sign_test", "dunn_test", "emmeans_test", "tukey_hsd", "games_howell_test".

A list of additional arguments used for the test method. For example one might use `method.args = list(alternative = "greater")` for wilcoxon test.

A character string or a numeric value specifying the reference group. If specified, for a given grouping variable, each of the group levels will be compared to the reference group (i.e. control group). `ref.group` can be also "all". In this case, each of the grouping variable levels is compared to all (i.e. basemean).

Allowed values can be:

- **numeric value**: specifying the rank of the reference group. For example, use `ref.group = 1` when the first group is the reference; use `ref.group = 2` when the second group is the reference, and so on. This works for all situations, including i) when comparisons are performed between x-axis groups and ii) when comparisons are performed between legend groups.

- **character value**: For example, you can use `ref.group = "ctrl"` instead of using the numeric rank value of the "ctrl" group.

- "all": In this case, each of the grouping variable levels is compared to all (i.e. basemean).

A character string specifying label. Can be:

- the column containing the label (e.g.: `label = "p"` or `label = "p.adj"`), where `p` is the p-value. Other possible values are "p.signif", "p.adj.signif", "p.format", "p.adj.format".

- an expression that can be formatted by the `glue()` package. For example, when specifying `label = "Wilcoxon, p = \{p\}"`, the expression `{p}` will be replaced by its value.

- a combination of plotmath expressions and glue expressions. You may want some of the statistical parameter in italic; for example: `label = "Wilcoxon, italic(p)=\{p\}"`.

A numeric vector with the y positions of the brackets.

(Optional) character vector specifying the grouping variable; it should be used only for grouped plots. Possible values are:
• "x.var": Group by the x-axis variable and perform the test between legend groups. In other words, the p-value is computed between legend groups at each x position.
• "legend.var": Group by the legend variable and perform the test between x-axis groups. In other words, the test is performed between the x-groups for each legend level.

dodge
dodge width for grouped ggplot2 test. Default is 0.8. It's used to dodge the brackets position when group.by = "legend.var".

bracket.nudge.y
Vertical adjustment to nudge brackets by (in fraction of the total height). Useful to move up or move down the bracket. If positive value, brackets will be moved up; if negative value, brackets are moved down.

bracket.shorten
a small numeric value in [0-1] for shortening the width of bracket.

bracket.group.by
(optional); a variable name for grouping brackets before adding step.increase. Useful for grouped plots. Possible values include "x.var" and "legend.var".

step.increase
numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.

tip.length
numeric vector with the fraction of total height that the bar goes down to indicate the precise column/size

label.size
change the size of the label text

family
change the font used for the text

vjust
move the text up or down relative to the bracket.

hjust
move the text left or right relative to the bracket.

p.adjust.method
method for adjusting p values (see p.adjust). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Ignored when the specified method is "tukey_hsd" or "games_howell_test" because they come with internal p adjustment method. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don’t want to adjust the p value (not recommended), use p.adjust.method = "none".

p.adjust.by
possible value is one of c("group", "panel"). Default is "group": for a grouped data, if pairwise test is performed, then the p-values are adjusted for each group level independently. P-values are adjusted by panel when p.adjust.by = "panel".

symnum.args
a list of arguments to pass to the function symnum for symbolic number coding of p-values. For example, symnum.args <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", ",", "ns"). In other words, we use the following convention for symbols indicating statistical significance:
  • ns: p > 0.05
• *: \( p \leq 0.05 \)
• **: \( p \leq 0.01 \)
• ***: \( p \leq 0.001 \)
• ****: \( p \leq 0.0001 \)

\( \text{hide.ns} \) can be logical value (TRUE or FALSE) or a character vector ("p.adj" or "p").

\( \text{remove.bracket} \) logical, if TRUE, brackets are removed from the plot.
• Case when logical value. If TRUE, hide ns symbol when displaying significance levels. Filter is done by checking the column p.adj.signif, p.signif, p.adj and p.
• Case when character value. Possible values are "p" or "p.adj", for filtering out non significant.

\( \text{position} \) Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

\( \text{na.rm} \) If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

\( \text{show.legend} \) logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

\( \text{inherit.aes} \) If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

\( \text{parse} \) logical for parsing plotmath expression.

... other arguments passed on to layer. These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat.

\( \text{stat} \) The statistical transformation to use on the data for this layer, either as a ggproto Geom subclass or as a string naming the stat stripped of the stat_ prefix (e.g. "count" rather than "stat_count")

\( \text{stack} \) logical value. Default is FALSE; should be set to TRUE for stacked bar plots or line plots. If TRUE, then the brackets are automatically removed and the dodge value is set to zero.

Details

Notes on adjusted p-values and facet. When using the ggplot facet functions, the p-values are computed and adjusted by panel, without taking into account the other panels. This is by design in ggplot2.

In this case, when there is only one computed p-value by panel, then using ‘label = "p"‘ or ‘label = "p.adj"‘ will give the same results using ‘geom_pwc()‘. Again, p-value computation and adjustment in a given facet panel is done independently to the other panels.

One might want to adjust the p-values of all the facet panels together. There are two solutions for that:

• Using \texttt{ggadjust_pvalue}(p) after creating the plot p
• or adding the adjusted p-value manually using `stat_pvalue_manual()`. Read more at:
  – How to Add P-values to GGPLOT Facets
  – Add P-values to GGPLOT Facets with Different Scales

See Also

`ggadjust_pvalue`

Examples

```r
df <- ToothGrowth
df$dose <- factor(df$dose)

# Data preparation
#%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
# Transform `dose` into factor variable
df <- ToothGrowth
df$dose <- as.factor(df$dose)
# Add a random grouping variable
df$group <- factor(rep(c("grp1", "grp2"), 30))
head(df, 3)

# Two groups by x position
#%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
# Create a box plot
# Add 10% spaces between the p-value labels and the plot border
bxp <- ggboxplot(
  df, x = "dose", y = "len",
  color = "supp", palette = c("#00AFBB", "#E7B800")
) +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.10)))

# Add p-values onto the box plots
# label can be "p.format" or "p.adj.format"
bxp + geom_pwc(
  aes(group = supp), tip.length = 0,
  method = "t_test", label = "p.format"
)

# Show adjusted p-values and significance levels
# Hide ns (non-significant)
bxp + geom_pwc(
  aes(group = supp), tip.length = 0,
  method = "t_test", label = "{p.adj.format}{p.adj.signif}",
  p.adjust.method = "bonferroni", p.adjust.by = "panel",
  hide.ns = TRUE
)

# Complex cases
```
# 1. Add p-values of OJ vs VC at each dose group
bxp.complex <- bxp +
  geom_pwc(
    aes(group = supp), tip.length = 0,
    method = "t_test", label = "p.adj.format",
    p.adjust.method = "bonferroni", p.adjust.by = "panel"
  )
# 2. Add pairwise comparisons between dose levels
# Nudge up the brackets by 20% of the total height
bxp.complex <- bxp.complex +
  geom_pwc(
    method = "t_test", label = "p.adj.format",
    p.adjust.method = "bonferroni",
    bracket.nudge.y = 0.2
  )
# 3. Display the plot
bxp.complex

# Three groups by x position

# Simple plots

# Box plots with p-values
bxp <- ggboxplot(
  df, x = "supp", y = "len", fill = "dose",
  palette = "npg"
)
bxp +
  geom_pwc(
    aes(group = dose), tip.length = 0,
    method = "t_test", label = "p.adj.format",
    bracket.nudge.y = -0.08
  ) +
  scale_y_continuous(expand = expansion(mult = c(0, 0.1)))

# Bar plots with p-values
bp <- ggbarchart(
  df, x = "supp", y = "len", fill = "dose",
  palette = "npg", add = "mean_sd",
  position = position_dodge(0.8)
)
bp +
  geom_pwc(
    aes(group = dose), tip.length = 0,
    method = "t_test", label = "p.adj.format",
    bracket.nudge.y = -0.08
  ) +
  scale_y_continuous(expand = expansion(mult = c(0, 0.1)))
**Description**

Add regression line equation and R^2 to a ggplot. Regression model is fitted using the function `lm`.

**Usage**

```r
stat_regline_equation(
  mapping = NULL,
  data = NULL,
  formula = y ~ x,
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  output.type = "expression",
  geom = "text",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

**Arguments**

- **mapping**: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.

- **data**: The data to be displayed in this layer. There are three options:
  - If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
  - A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
  - A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

- **formula**: a formula object

- **label.x.npc, label.y.npc**: can be numeric or character vector of the same length as the number of groups and/or panels. If too short they will be recycled.
  - If numeric, value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates".
stat_regline_equation

- If character, allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c( 'bottom', 'top', 'center', 'centre', 'middle') for y-axis.

If too short they will be recycled.

label.x, label.y
numeric Coordinates (in data units) to be used for absolute positioning of the label. If too short they will be recycled.

output.type character One of "expression", "latex" or "text".

geom The geometric object to use to display the data, either as a ggproto Geom sub-class or as a string naming the geom stripped of the geom_ prefix (e.g. "point" rather than "geom_point")

position Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

na.rm If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

show.legend logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

... other arguments to pass to geom_text or geom_label.

Computed variables

x x position for left edge
y y position near upper edge
eq.label equation for the fitted polynomial as a character string to be parsed
rr.label $R^2$ of the fitted model as a character string to be parsed
adj.rr.label Adjusted $R^2$ of the fitted model as a character string to be parsed
AIC.label AIC for the fitted model.
BIC.label BIC for the fitted model.
hjust Set to zero to override the default of the "text" geom.

References

the source code of the function stat_regline_equation() is inspired from the code of the function stat_poly_eq() in ggpmisc package.

See Also

ggscatter
Examples

```r
# Simple scatter plot with correlation coefficient and regression line
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::
ggscatter(mtcars, x = "wt", y = "mpg", add = "reg.line") +
  stat_cor(label.x = 3, label.y = 34) +
  stat_regline_equation(label.x = 3, label.y = 32)
```

```r
# Grouped scatter plot
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::

ggscatter(
  iris, x = "Sepal.Length", y = "Sepal.Width",
  color = "Species", palette = "jco",
  add = "reg.line"
) +
  facet_wrap(~Species) +
  stat_cor(label.y = 4.4) +
  stat_regline_equation(label.y = 4.2)
```

```r
# Polynomial equation
#:::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::::

# Demo data
set.seed(4321)
x <- 1:100
y <- (x + x^2 + x^3) + rnorm(length(x), mean = 0, sd = mean(x^3) / 4)
my.data <- data.frame(x, y, group = c("A", "B"),
y2 = y * c(0.5, 2), block = c("a", "a", "b", "b"))

# Fit polynomial regression line and add labels
formula <- y ~ poly(x, 3, raw = TRUE)
p <- ggplot(my.data, aes(x, y2, color = group)) +
  geom_point() +
  stat_smooth(aes(fill = group, color = group), method = "lm", formula = formula) +
  stat_regline_equation(
    aes(label = paste(..eq.label.., ..adj.rr.label.., sep = "~~~~")),
    formula = formula
  ) +
  theme_bw()
ggpar(p, palette = "jco")
```

---

**stat_stars**

*Add Stars to a Scatter Plot*

**Description**

Create a star plot by drawing segments from group centroid to each points.
Usage

```r
callout
stat_stars(
  mapping = NULL,
  data = NULL,
  geom = "segment",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
callout
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>mapping</td>
<td>Set of aesthetic mappings created by <code>aes()</code>. If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.</td>
</tr>
<tr>
<td>data</td>
<td>The data to be displayed in this layer. There are three options: If <code>NULL</code>, the default, the data is inherited from the plot data as specified in the call to <code>ggplot()</code>. A <code>data.frame</code>, or other object, will override the plot data. All objects will be fortified to produce a data frame. See <code>fortify()</code> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code>).</td>
</tr>
<tr>
<td>geom</td>
<td>The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_prefix (e.g. &quot;point&quot; rather than &quot;geom_point&quot;)</td>
</tr>
<tr>
<td>position</td>
<td>Position adjustment, either as a string naming the adjustment (e.g. &quot;jitter&quot; to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.</td>
</tr>
<tr>
<td>na.rm</td>
<td>If <code>FALSE</code> (the default), removes missing values with a warning. If <code>TRUE</code> silently removes missing values.</td>
</tr>
<tr>
<td>show.legend</td>
<td>logical. Should this layer be included in the legends? <code>NA</code>, the default, includes if any aesthetics are mapped. <code>FALSE</code> never includes, and <code>TRUE</code> always includes. It can also be a named logical vector to finely select the aesthetics to display.</td>
</tr>
<tr>
<td>inherit.aes</td>
<td>If <code>FALSE</code>, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. <code>borders()</code>.</td>
</tr>
</tbody>
</table>

See Also

`ggscatter`
Examples

```r
# Load data
data("mtcars")
df <- mtcars
df$cyl <- as.factor(df$cyl)

# Scatter plot with ellipses and group mean points
ggscatter(df, x = "wt", y = "mpg",
          color = "cyl", shape = "cyl",
          mean.point = TRUE, ellipse = TRUE)+
          stat_stars(aes(color = cyl))
```

---

**stat_welch_anova_test**  Add Welch One-Way ANOVA Test P-values to a GGPlot

Description

Add Welch one-way ANOVA test p-values to a ggplot, such as box plots, dot plots and stripcharts.

Usage

```r
stat_welch_anova_test(
  mapping = NULL,
  data = NULL,
  group.by = NULL,
  label = "{method}, p = {p.format}",
  label.x.npc = "left",
  label.y.npc = "top",
  label.x = NULL,
  label.y = NULL,
  step.increase = 0.1,
  p.adjust.method = "holm",
  significance = list(),
  geom = "text",
  position = "identity",
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE,
  parse = FALSE,
  ...)
```

Arguments

- `mapping` Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.
The data to be displayed in this layer. There are three options:

- If NULL, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
- A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
- A function will be called with a single argument, the plot data. The return value must be a data.frame, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

`group.by` (optional) character vector specifying the grouping variable; it should be used only for grouped plots. Possible values are:

- "x.var": Group by the x-axis variable and perform the test between legend groups. In other words, the p-value is computed between legend groups at each x position
- "legend.var": Group by the legend variable and perform the test between x-axis groups. In other words, the test is performed between the x-groups for each legend level.

`label` the column containing the label (e.g.: label = "p" or label = "p.adj"), where p is the p-value. Can be also an expression that can be formatted by the `glue()` package. For example, when specifying label = "t-test, p = {p}", the expression {p} will be replaced by its value.

`label.x.npc, label.y.npc` can be numeric or character vector of the same length as the number of groups and/or panels. If too short they will be recycled.

- If numeric, value should be between 0 and 1. Coordinates to be used for positioning the label, expressed in "normalized parent coordinates".
- If character, allowed values include: i) one of c('right', 'left', 'center', 'centre', 'middle') for x-axis; ii) and one of c('bottom', 'top', 'center', 'centre', 'middle') for y-axis.

`label.x, label.y` numeric Coordinates (in data units) to be used for absolute positioning of the label. If too short they will be recycled.

`step.increase` numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.

`p.adjust.method` method for adjusting p values (see `p.adjust`). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don’t want to adjust the p value (not recommended), use p.adjust.method = "none".

`significance` a list of arguments specifying the significance cutpoints and symbols. For example, `significance <- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", ", , "ns"))`.

In other words, we use the following convention for symbols indicating statistical significance:
• ns: \( p > 0.05 \)
• *: \( p \leq 0.05 \)
• **: \( p \leq 0.01 \)
• ***: \( p \leq 0.001 \)
• ****: \( p \leq 0.0001 \)

**geom**

The geometric object to use to display the data, either as a ggproto Geom subclass or as a string naming the geom stripped of the geom_ prefix (e.g. "point" rather than "geom_point")

**position**

Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use position_jitter), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

**na.rm**

If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

**show.legend**

Logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

**inherit.aes**

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn’t inherit behaviour from the default plot specification, e.g. borders().

**parse**

If TRUE, the labels will be parsed into expressions and displayed as described in ?plotmath.

... other arguments passed to the function geom_bracket() or geom_text()

**Computed variables**

- statistic: the value of the test statistic (F-value)
- DFn: Degrees of Freedom in the numerator (i.e. DF effect)
- DFd: Degrees of Freedom in the denominator (i.e., DF error)
- p: p-value.
- p.adj: Adjusted p-values.
- p.signif: P-value significance.
- p.adj.signif: Adjusted p-value significance.
- p.format: Formatted p-value.
- p.adj.format: Formatted adjusted p-value.
- n: number of samples.

**Examples**

```r
# Data preparation
# Transform `dose` into factor variable
df <- ToothGrowth
df$dose <- as.factor(df$dose)
# Add a random grouping variable
```
set.seed(123)
df$group <- sample(factor(rep(c("grp1", "grp2", "grp3"), 20)))
df$len <- ifelse(df$group == "grp2", df$len+2, df$len)
df$len <- ifelse(df$group == "grp3", df$len+7, df$len)
head(df, 3)

# Basic boxplot
# Create a basic boxplot
# Add 5% and 10% space to the plot bottom and the top, respectively
bxp <- ggboxplot(df, x = "dose", y = "len") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))
# Add the p-value to the boxplot
bxp + stat_welch_anova_test()
# Change the label position
# Using coordinates in data units
bxp + stat_welch_anova_test(label.x = "1", label.y = 10, hjust = 0)
# Format the p-value differently
custom_p_format <- function(p) {
  rstatix::p_format(p, accuracy = 0.0001, digits = 3, leading.zero = FALSE)
}
bxp + stat_welch_anova_test(label = "Welch Anova, italic(p) = {custom_p_format(p)}{p.signif}"
)
# Show a detailed label in italic
bxp + stat_welch_anova_test(label = "as_detailed_italic")

# Faceted plots
# Create a ggplot facet
bxp <- ggboxplot(df, x = "dose", y = "len", facet.by = "supp") +
  scale_y_continuous(expand = expansion(mult = c(0.05, 0.1)))
# Add p-values
bxp + stat_welch_anova_test()

# Grouped plots
bxp2 <- ggboxplot(df, x = "group", y = "len", color = "dose", palette = "npg")
# For each x-position, computes tests between legend groups
bxp2 + stat_welch_anova_test(aes(group = dose), label = "p = {p.format}{p.signif}%")
# For each legend group, computes tests between x variable groups
bxp2 + stat_welch_anova_test(aes(group = dose, color = dose), group.by = "legend.var")
Create easily a customized text grob (graphical object). Wrapper around textGrob.

Usage

text_grob(
  label,
  just = "centre",
  hjust = NULL,
  vjust = NULL,
  rot = 0,
  color = "black",
  face = "plain",
  size = NULL,
  lineheight = NULL,
  family = NULL,
  ...
)

Arguments

  label  A character or expression vector. Other objects are coerced by as.graphicsAnnot.
  just   The justification of the text relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". For numeric values, 0 means left (bottom) alignment and 1 means right (top) alignment.
  hjust A numeric vector specifying horizontal justification. If specified, overrides the just setting.
  vjust A numeric vector specifying vertical justification. If specified, overrides the just setting.
  rot   The angle to rotate the text.
  color text font color.
  face  font face. Allowed values include one of "plain", "bold", "italic", "bold.italic".
  size  font size (e.g.: size = 12)
  lineheight line height (e.g.: lineheight = 2).
  family font family.
  ... other arguments passed to textGrob.
Value

a text grob.

Examples

text <- paste("iris data set gives the measurements in cm",
"of the variables sepal length and width",
"and petal length and width, respectively,",
"for 50 flowers from each of 3 species of iris.",
"The species are Iris setosa, versicolor, and virginica.", sep = "\n")

# Create a text grob
tgrob <- text_grob(text, face = "italic", color = "steelblue")
# Draw the text
as_ggplot(tgrob)

theme_pubr          Publication ready theme

Description

• theme_pubr(): Create a publication ready theme
• theme_pubclean(): a clean theme without axis lines, to direct more attention to the data.
• labs_pubr(): Format only plot labels to a publication ready style
• theme_classic2(): Create a classic theme with axis lines.
• clean_theme(): Remove axis lines, ticks, texts and titles.
• clean_table_theme(): Clean the theme of a table, such as those created by ggsummarytable()

Usage

theme_pubr(
  base_size = 12,
  base_family = "",
  border = FALSE,
  margin = TRUE,
  legend = c("top", "bottom", "left", "right", "none"),
  x.text.angle = 0
)

theme_pubclean(base_size = 12, base_family = "", flip = FALSE)

labs_pubr(base_size = 14, base_family = "")

theme_classic2(base_size = 12, base_family = "")
clean_theme()

clean_table_theme()

Arguments

base_size    base font size
base_family  base font family
border       logical value. Default is FALSE. If TRUE, add panel border.
margin       logical value. Default is TRUE. If FALSE, reduce plot margin.
legend       character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). Default is "top" side position. To remove the legend use legend = "none". Legend position can be also specified using a numeric vector c(x, y). In this case it is possible to position the legend inside the plotting area. x and y are the coordinates of the legend box. Their values should be between 0 and 1. c(0,0) corresponds to the "bottom left" and c(1,1) corresponds to the "top right" position. For instance use legend = c(0.8, 0.2).
x.text.angle Rotation angle of x axis tick labels. Default value is 0. Use 90 for vertical text.
flip         logical. If TRUE, grid lines are added to y axis instead of x axis.

Examples

p <- ggplot(mtcars, aes(x = wt, y = mpg)) +
  geom_point(aes(color = gear))

# Default plot
p

# Use theme_pubr()
p + theme_pubr()

# Format labels
p + labs_pubr()

theme_transparent

Create a ggplot with Transparent Background

Description

Create a ggplot with transparent background.

Usage

theme_transparent(base_size = 12, base_family = "")
theme_transparent

Arguments

- base_size  base font size
- base_family base font family

See Also

- theme_pubr

Examples

# Create a scatter plot
sp <- ggscatter(iris, x = "Sepal.Length", y = "Sepal.Width",
               color = "Species", palette = "jco",
               size = 3, alpha = 0.6)
sp

# Transparent theme
sp + theme_transparent()
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