Package ‘grafify’

Type     Package
Title    Easy Graphs for Data Visualisation and Linear Models for ANOVA
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Description  
Easily explore data by generating different kinds of graphs with few lines of code. Use these ggplot() wrappers to quickly draw graphs of scatter/dots with box-whiskers, violins or SD error bars, data distributions, before-after graphs, factorial ANOVA and more. Customise graphs and choose from 8 colour-blind friendly discreet or 1 continuous colour scheme. Simple code for ANOVA as ordinary (lm()) or mixed-effects linear models (lmer()), including randomised-block or repeated-measures designs. Carry out post-hoc comparisons on fitted models (via emmeans() wrappers). Also includes small datasets for practicing code and teaching basics before users move on to more complex designs. See vignettes for details on usage <https://grafify-vignettes.netlify.app/>. Citation: <doi:10.5281/zenodo.5136508>.

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colorRampPalette_d

Description

Variant of colorRampPalette for sequential use of colours for discrete scales. Thank you linog.
Called by other functions in grafify and not generally meant to be called by user.

Usage

colorRampPalette_d(colors, ...)

Arguments

colors  internal
...     internal

Value

This is a variant of colourRampPalette that generates sequential colours from chosen grafify palettes when called by graf_col_palette_d.
**Description**

Variant of colorRamp for sequential use of colours for discrete scales. Thank you linog. Called by other functions in grafify and not generally meant to be called by user.

**Usage**

```r
colorRamp_d(
  colors,
  n,
  bias = 1,
  space = c("rgb", "Lab"),
  interpolate = c("linear", "spline"),
  alpha = FALSE
)
```

**Arguments**

- `colors`: internal
- `n`: internal
- `bias`: internal
- `space`: internal
- `interpolate`: internal
- `alpha`: internal

**Value**

It generates the required number of sequential colours from chosen grafify palettes when called by `colorRampPalette_d`.

---

**Description**

*In vitro experiments measuring percentage cell death in three genotypes of cells.*

These data are from *in vitro* measurements of death of host cells (measured as percentage of total cells) after infection with three different strains of a pathogenic bacterium, from five independent experiments. The three strains are three levels within the fixed factor Genotype. The five independent experiments are levels within the random variable Experiment. These data can be analysed using linear mixed effects modeling. These data are from Goddard *et al*, Cell Rep, 2019, doi.org/10.1016/j.celrep.2019.03.100
data_2w_Festing

Usage

data_1w_death

Format

data.frame: 15 obs. of 3 variables.

  Experiment  Experiment - a random factor with 5 levels "Exp_1","Exp_2"...
  Genotype   Genotypes - a fixed factor with 3 levels: "WT","KO_1","KO_2".
  Death      Numerical dependent variable indicating percentage cell death.

Data from two-way ANOVA with randomised block design of treatments of strains of mice.

Description

Data from Festing, ILAR Journal (2014) 55, 472–476 <doi: 10.1093/ilar/ilu045>. These data are suitable for two-way linear mixed effects modelling. The activity of GST (numerical dependent variable) was measured in 4 strains of mice (levels with the fixed factor Strain) either treated or controls (levels within the fixed factor Treatment). Once mouse each was used in two randomised blocks, which is the random factor (Block).

Usage

data_2w_Festing

Format

data.frame: 16 obs. of 4 variables:

  Block     A random factor with 2 levels "A" and "B".
  Treatment A fixed factor with 2 levels: "Control" & "Treated"
  Strain    A fixed factor with 4 levels: "129Ola", "A/J", "NIH" & "BALB/C"
  GST       Numerical dependent variable indicating GST activity measurement
**data_2w_Tdeath**  
*In vitro measurement of percentage cell death - two-way ANOVA design with repeated measures, and randomised blocks.*

**Description**

These are measurements of death of infected host cells (as percentage of total cells) upon infection with two strains of bacteria, measured at two time points, in 6 independent experiments. These data repeated-measures data suitable for two-way linear mixed effects modeling with experiment and subjects as random factors.

**Usage**

`data_2w_Tdeath`

**Format**

data.frame: 24 obs. of 6 variables:

- **Experiment**  A random factor with 6 levels "e1", "e2"...
- **Time**  A fixed factor with 2 levels: "t100" & "t300".
- **Time2**  A numeric column that allows plotting data on a quantitative "Time" axis. The "Time" column has "factor" type values that should be used for the ANOVA.
- **Genotype**  A fixed factor with 2 levels that we want to compare "WT" & "KO".
- **Subject**  A random factor with 12 levels: "s1", "s2"... These are cell culture wells that were measured at two time points, and indicate "subjects" that underwent repeated-measures within each of 6 experiments. Subject IDs for WT and KO are unique and clearly indicate different wells.
- **PI**  Numerical dependent variable indicating propidium iodide dye uptake as a measure of cell death. These are percentage of dead cells out of total cells plated.

**data_cholesterol**  
*Hierarchical data from 25 subjects either treated or not at 5 hospitals - two-way ANOVA design with repeated measures.*

**Description**

An example dataset on measurements of blood cholesterol levels measured in 5 subjects measured before and after receiving a Drug. Five patients each were recruited at 5 hospitals (a-e), so that there are 25 different subjects (1-25) measured twice. Data are from Micro/Immuno Stats

**Usage**

`data_cholesterol`
**data_doubling_time**

**Format**

`tibble`: 30 obs. of 3 variables:

- **Hospital**  Factor with 5 levels (a-e), representing different hospitals where subjects were recruited.
- **Subject**  A factor with 25 levels denoting individuals on whom measurements were made twice.
- **Treatment**  A factor with 2 levels indicating when measurements were made, i.e. before and after drug.
- **Cholesterol**  Numerical dependent variable indicating measured doubling time in min.

**Description**

An example dataset showing measurements of *E. coli* doubling times (in min) measured by 10 different students in 3 independent experiments each. Note that Experiments are just called Exp1-Exp3 even though Exp1 of any of the students are not connected in anyway - this will confuse R! Data are from Micro/Immuno Stats

**Usage**

`data_doubling_time`

**Format**

`tibble`: 30 obs. of 3 variables:

- **Student**  Factor with 10 levels, representing different students.
- **Experiment**  A factor with 3 levels representing independent experiments.
- **Doubling_time**  Numerical dependent variable indicating measured doubling time in min.

**data_t_pdiff**  

**Description**

An example dataset for paired difference Student's *t* test. These are bodyweight (Mass) in grams of same mice left untreated or treated, which are two groups to be compared. The data are in a longtable format, and the two groups are levels within the factor "Condition". The Subject column lists ID of matched mice that were measured without and with treatment. These data are from Sanchez-Garrido *et al*, Sci Signal, 2018, DOI: 10.1126/scisignal.aat6903.
**Usage**

data_t_pdiff

**Format**

data.frame: 20 obs. of 3 variables:

- **Subject** Factor with 10 levels, denoted by capital letters, representing individuals or subjects.
- **Condition** A fixed factor with 2 levels: "Untreated" & "Treated".
- **Mass** Numerical dependent variable indicating body mass of mice

---

**data_t_pratio**  
*Matched data from two groups where ratio between them is consistent.*

**Description**

An example dataset for paired ratio Student’s t test. These are Cytokine measurements by ELISA (in ng/ml) from 33 independent in vitro experiments performed on two Genotypes that we want to compare. The data are in a longtable format, and the two groups are levels within the factor "Genotype". The Experiment column lists ID of matched experiments.

**Usage**

data_t_pratio

**Format**

data.frame: 66 obs. of 3 variables:

- **Genotype** Factor with 2 levels, representing genotypes to be compared ("WT" & "KO").
- **Experiment** A random factor with 33 levels representing independent experiments, denoted as "Exp_1", "Exp_2"...
- **Cytokine** Numerical dependent variable indicating cytokine measured by ELISA.
**get_graf_colours**

**get_graf_colours**  \hspace{1cm}  *Get graf internal*

---

**Description**

Function to make grafify colour scheme. Thank you Dr Simon.

**Usage**

```r
get_graf_colours(...)  
```

**Arguments**

```r
...  \hspace{1cm}  internal
```

**Details**

To visualise grafify colours use `plot_grafify_palette`.

**Value**

This function returns names and hexcodes of colours in grafify as a character vector.

---

**graf_colours**  \hspace{1cm}  *List of hexcodes of colours in grafify palettes*

---

**Description**

To visualise these colours use `plot_grafify_palette`.

**Usage**

```r
graf_colours
```

**Format**

An object of class character of length 64.

**Value**

This is a character vector with names and hexcodes of colours used by palette functions. It is used by `get_graf_colours` to generate palettes.
**graf_col_palette**  
*Call palettes for scale & fill*

### Description
Call palettes for scale & fill

### Usage

```
graf_col_palette(palette = "all_grafify", reverse = FALSE, ...)
```

### Arguments

- `palette` internal
- `reverse` internal
- `...` additional parameters

### Value
This generates required number of sequential colours from the chosen grafify palette when called by scale functions of ggplot2.

---

**graf_col_palette_default**  
*Call palettes for scale & fill with default colorRampPalette*

### Description
Call palettes for scale & fill with default colorRampPalette

### Usage

```
graf_col_palette_default(palette = "all_grafify", reverse = FALSE, ...)
```

### Arguments

- `palette` internal
- `reverse` internal
- `...` additional parameters

### Value
This generates required number of distant colours from the chosen grafify palette when called by scale functions of ggplot2.
**graf_palettes**

*List of palettes available in grafify package*

**Description**

To visualise these colours use `plot_grafify_palette`.

**Usage**

```r
graf_palettes
```

**Format**

An object of class list of length 10.

**Value**

This function returns a list of palettes in grafify with names and hexcodes of colours in those palettes.

**make_1way_data**

*Make one-way or two-way independent group or randomised block design data.*

**Description**

The `make_1way_data`, `make_1way_rb_data`, `make_2way_data` and `make_2way_rb_data` functions generate independent or randomised block (rb) design data of one-way or two-way designs.

**Usage**

```r
make_1way_data(Group_means, Num_obs, Residual_SD)
```

**Arguments**

- **Group_means**
  
a vector with means of each level of the first fixed factor (FixFac_X1) measured within Group 1.

- **Num_obs**
  
a single numeric value indicating the number of independent measurements, i.e. levels within the random factor Experiment.

- **Residual_SD**
  
a single numeric value indicating residual SD in the model.
make_1way_rb_data

Details

Random variates from the normal distribution based on user provided mean and SD provided are generated. For independent designs, the Residual_SD argument is used to set expected residual SD from the linear model. Exp_SD is used to set experiment-to-experiment SD, that will be assigned to the random factor for rb designs.

Num_exp sets the number of independent measurements per group.

For one-way designs, the user provides Group_means as a vector. Number of levels are recognised based on number of means. For two-way designs, two vectors are to be provided by the user containing means of levels of a second factor. Number of means in both vectors should be the same. These functions can only handle balanced designs, i.e. same number of observations in all groups.

The output is a data frame with one or two columns denoting the fixed factor with levels that match the number of means entered. For rb data, the column for RandFac denotes levels of the blocking factor. The quantitative response variables are in the numeric Values column.

Value

This function produces a data.frame object containing simulated data.

Examples

#Basic usage with three levels within Factor_X, #20 observations in each group, with residual SD 15
one_independent_tab <- make_1way_data(c(350, 250, 100), 15, 20)
str(one_independent_tab)
head(one_independent_tab)

make_1way_rb_data

Make one-way or two-way independent group or randomised block design data.

Description

The make_1way_data, make_1way_rb_data, make_2way_data and make_2way_rb_data functions generate independent or randomised block (rb) design data of one-way or two-way designs.

Usage

make_1way_rb_data(Group_means, Num_exp, Exp_SD, Residual_SD)
**make_2way_data**

**Arguments**

- **Group_means**: a vector with means of each level of the first fixed factor (FixFac_X1) measured within Group 1.
- **Num_exp**: a single numeric value indicating the number of independent measurements, i.e. levels within the random factor RandFac.
- **Exp_SD**: a single numeric value indicating the standard deviation (SD) between experiments, i.e. within RandFac.
- **Residual_SD**: a single numeric value indicating residual SD in the model.

**Details**

Random variates from the normal distribution based on user provided mean and SD provided are generated. For independent designs, the Residual_SD argument is used to set expected residual SD from the linear model. Exp_SD is used to set experiment-to-experiment SD, that will be assigned to the random factor for rb designs.

Num_exp sets the number of independent measurements per group.

For one-way designs, the user provides Group_means as a vector. Number of levels are recognised based on number of means. For two-way designs, two vectors are to be provided by the user containing means of levels of a second factor. Number of means in both vectors should be the same. These functions can only handle balanced designs, i.e. same number of observations in all groups.

The output is a data frame with one or two columns denoting the fixed factor with levels that match the number of means entered. For rb data, the column for RandFac denotes levels of the blocking factor. The quantitative response variables are in the numeric Values column.

**Value**

This function produces a data.frame object containing simulated data.

**Examples**

# Basic usage with two levels within FactorX2,
# 20 experiments with inter-experiment SD 20, and residual SD 15

two_rb_tab <- make_2way_rb_data(c(100, 20), c(200, 300), 20, 20, 15)

str(two_rb_tab)
head(two_rb_tab)

---

**Description**

The `make_1way_data`, `make_1way_rb_data`, `make_2way_data` and `make_2way_rb_data` functions generate independent or randomised block (rb) design data of one-way or two-way designs.
make_2way_data

Usage

make_2way_data(Group_1_means, Group_2_means, Num_obs, Residual_SD)

Arguments

Group_1_means a vector with means of each level of the first fixed factor (FixFac_X1) measured within Group 1.

Group_2_means only for make_2way_data and make_2way_rb_data: a vector with mean(s) of each level of FactorX2 measured within Group 2.

Num_obs a single numeric value indicating the number of independent measurements, i.e. levels within the random factor Experiment.

Residual_SD a single numeric value indicating residual SD in the model.

Details

Random variates from the normal distribution based on user provided mean and SD provided are generated. For independent designs, the Residual_SD argument is used to set expected residual SD from the linear model. Exp_SD is used to set experiment-to-experiment SD, that will be assigned to the random factor for rb designs.

Num_obs sets the number of independent measurements per group.

For one-way designs, the user provides Group_means as a vector. Number of levels are recognised based on number of means. For two-way designs, two vectors are to be provided by the user containing means of levels of a second factor. Number of means in both vectors should be the same. These functions can only handle balanced designs, i.e. same number of observations in all groups.

The output is a data frame with one or two columns denoting the fixed factor with levels that match the number of means entered. For rb data, the column for RandFac denotes levels of the blocking factor. The quantitative response variables are in the numeric Values column.

Value

This function produces a data.frame object containing simulated data.

Examples

#Basic usage with two levels within FactorX2, 20 observations in each group, with residual SD 15
two_independent_tab <- make_2way_data(c(100, 20), c(200, 300), 20, 15)

#Four levels with 5 observations and residual SD 5
two_independent_tab <- make_2way_data(c(100, 20, 1500, 20), c(150, 5, 1450, 25), 5, 5)
**Description**

The `make_1way_data`, `make_1way_rb_data`, `make_2way_data` and `make_2way_rb_data` functions generate independent or randomised block (rb) design data of one-way or two-way designs.

**Usage**

```r
make_2way_rb_data(Group1_means, Group2_means, Num_exp, Exp_SD, Residual_SD)
```

**Arguments**

- **Group1_means**: a vector with means of each level of the first fixed factor (FixFac_X1) measured within Group 1.
- **Group2_means**: only for `make_2way_data` and `make_2way_rb_data`: a vector with mean(s) of each level of FactorX2 measured within Group 2.
- **Num_exp**: a single numeric value indicating the number of independent measurements, i.e. levels within the random factor RandFac.
- **Exp_SD**: a single numeric value indicating the standard deviation (SD) between experiment, i.e. within RandFac.
- **Residual_SD**: a single numeric value indicating residual SD in the model.

**Details**

Random variates from the normal distribution based on user provided mean and SD provided are generated. For independent designs, the `Residual_SD` argument is used to set expected residual SD from the linear model. `Exp_SD` is used to set experiment-to-experiment SD, that will be assigned to the random factor RandFac for rb designs.

`Num_exp` sets the number of independent measurements per group.

For one-way designs, the user provides `Group_means` as a vector. Number of levels are recognised based on number of means. For two-way designs, two vectors are to be provided by the user containing means of levels of a second factor. Number of means in both vectors should be the same. These functions can only handle balanced designs, i.e. same number of observations in all groups.

The output is a data frame with one or two columns denoting the fixed factor with levels that match the number of means entered. For rb data, the column for RandFac denotes levels of the blocking factor. The quantitative response variables are in the numeric `Values` column.

**Value**

This function produces a data frame object containing simulated data.
Examples

```r
# Basic usage with two levels within FactorX2,
# 20 experiments with inter-experiment SD 20, and residual SD 15

two_rb_tab <- make_2way_rb_data(c(100, 20), c(200, 300), 20, 20, 15)

str(two_rb_tab)
head(two_rb_tab)
```

---

**mixed_anova**  
*ANOVA table from linear mixed effects analysis.*

Description

There are four related functions for mixed effects analyses: `mixed_model`, `mixed_anova`, `mixed_model_slopes`, and `mixed_anova_slopes`.

Usage

```r
mixed_anova(
  data,  
  Y_value,
  Fixed_Factor,
  Random_Factor,
  Df_method = "Kenward-Roger",
  SS_method = "II",
  ...
)
```

Arguments

- `data` : a data table object, e.g. data.frame or tibble.
- `Y_value` : name of column containing quantitative (dependent) variable, provided within "quotes".
- `Fixed_Factor` : name(s) of categorical fixed factors (independent variables) provided as a vector if more than one or within "quotes".
- `Random_Factor` : name(s) of random factors to allow random intercepts; to be provided as a vector when more than one or within "quotes".
- `Df_method` : method for calculating degrees of freedom. Default is Kenward-Roger, can be changed to "Satterthwaite".
- `SS_method` : type of sum of square, default is type II, can be changed to "I", "III", "1" or "2", or others.
- `...` : any additional arguments to pass on to `lmer` if required.
mixed_anova_slopes

Details

This function uses \texttt{lmer} to fit a linear mixed effect model and provides the model object, which could be used for post-hoc comparisons. The model object is converted to class \texttt{lmerModLmerTest} object by \texttt{as_lmerModLmerTest}. This is then passes on the model to \texttt{anova} and provides the ANOVA table with F and P values. It produces a type II sum of squares ANOVA table with Kenward-Roger approximation for degrees of freedom (as implemented in \texttt{lmerTest}) package. It requires a data table, one dependent variable (Y\_value), one or more independent variables (Fixed\_Factor), and at least one random factor (Random\_Factor). These should match names of variables in the long-format data table exactly. This function is related to \texttt{mixed_model}.

More than one fixed factors can be provided as a vector (e.g. c(“A”, “B”)). A full model with interaction term is fitted. This means when \texttt{Y\_value = Y, Fixed\_factor = c(“A”, “B”), Random\_factor = “R”} are entered as arguments, these are passed on as \texttt{Y ~ A*B + (1|R)} (which is equivalent to \texttt{Y ~ A + B + A:B + (1|R)}). For simplicity, only random intercepts are fitted ((1|R)).

Value

ANOVA table of class ”anova” and ”data.frame”.

Examples

#Usage with one fixed (Student) and random factor (Experiment)
mixed_anova(data = data_doubling_time, 
  Y_value = ”Doubling_time”, 
  Fixed_Factor = ”Student”, 
  Random_Factor = ”Experiment”)

#two fixed factors provided as a vector
mixed_anova(data = data_cholesterol, 
  Y_value = ”Cholesterol”, 
  Fixed_Factor = c(”Treatment”, ”Hospital”), 
  Random_Factor = ”Subject”)

mixed_anova_slopes

ANOVA table from linear mixed effects analysis.

Description

There are four related functions for mixed effects analyses: \texttt{mixed_model}, \texttt{mixed_anova}, \texttt{mixed_model_slopes}, and \texttt{mixed_anova_slopes}.

Usage

mixed_anova_slopes(
  data, 
  Y_value, 
  Fixed_Factor,
mixed_anova_slopes

Slopes_Factor,
Random_Factor,
Df_method = "Kenward-Roger",
SS_method = "II",
...
)

Arguments

data a data table object, e.g. data.frame or tibble.
Y_value name of column containing quantitative (dependent) variable, provided within "quotes".
Fixed_Factor name(s) of categorical fixed factors (independent variables) provided as a vector if more than one or within "quotes".
Slopes_Factor name of factor to allow varying slopes on.
Random_Factor name(s) of random factors to allow random intercepts; to be provided as a vector when more than one or within "quotes".
Df_method method for calculating degrees of freedom. Default is Kenward-Roger, can be changed to "Satterthwaite".
SS_method type of sum of square, default is type II, can be changed to "I", "III", "1" or "2", or others.
... any additional arguments to pass on to lmer if required.

Details

This function uses lmer to fit a linear mixed effect model and provides the model object, which could be used for post-hoc comparisons. The model object is converted to class lmerModLmerTest object by as.lmerModLmerTest.

It produces a type II sum of squares ANOVA table with Kenward-Roger approximation for degrees of freedom (as implemented in lmerTest package). It requires a data table, one dependent variable (Y_value), one or more independent variables (Fixed_Factor). Exactly one random factor (Random_Factor) and Slope_Factor should be provided. This function is related to mixed_model.

More than one fixed factors can be provided as a vector (e.g. c("A", "B")). A full model with interaction term is fitted with one term each for varying slopes and intercepts. This means when $Y_{value} = Y, Fixed\_factor = c("A", "B"), Slopes\_Factor = "S", Random\_factor = "R"$ are entered as arguments, these are passed on as $Y \sim A*B + (S|R)$ (which is equivalent to $Y \sim A + B + A:B + (S|R)$). In this experimental implementation, random slopes and intercepts are fitted ((Slopes\_Factor | Random\_Factor)). Only one term each is allowed for ~ and Random\_Factor.

Value

ANOVA table of class "anova" and "data.frame".
mixed_model

Examples

```r
mixed_anova_slopes(data = data_2w_Tdeath,
  Y_value = "PI",
  Fixed_Factor = c("Genotype", "Time"),
  Slopes_Factor = "Time",
  Random_Factor = "Experiment")
```

mixed_model  Model from a linear mixed effects model

Description

There are four related functions for mixed effects analyses: mixed_model, mixed_anova, mixed_model_slopes, and mixed_anova_slopes.

Usage

```r
mixed_model(data, Y_value, Fixed_Factor, Random_Factor, ...)
```

Arguments

data  a data table object, e.g. data.frame or tibble.
Y_value  name of column containing quantitative (dependent) variable, provided within "quotes".
Fixed_Factor  name(s) of categorical fixed factors (independent variables) provided as a vector if more than one or within "quotes".
Random_Factor  name(s) of random factors to allow random intercepts; to be provided as a vector when more than one or within "quotes".
...  any additional arguments to pass on to lmer if required.

Details

This function uses lmer to fit a linear mixed effect model and provides the model object, which could be used for post-hoc comparisons. The model object is converted to class lmerModLmerTest object by as_lmerModLmerTest.

It requires a data table, one dependent variable (Y_value), one or more independent variables (Fixed_Factor), and at least one random factor (Random_Factor). These should match names of variables in the long-format data table exactly. This function is related to mixed_anova. Output of this function can be used with posthoc_Pairwise, posthoc_Levelwise and posthoc_vsRef, or with emmeans.

More than one fixed factors can be provided as a vector (e.g. c("A","B")). A full model with interaction term is fitted. This means when Y_value = Y, Fixed_factor = c("A","B"), Random_factor = "R" are entered as arguments, these are passed on as Y ~ A*B + (1|R) (which is equivalent to Y ~ A + B + A:B + (1|R)). For simplicity, only random intercepts are fitted ((1|R)).

Also see mixed_anova_slopes and mixed_model_slopes for similar functions where variable slopes and intercept models are fit.
Value

This function returns an S4 object of class "lmerModLmerTest".

Examples

# one fixed factor and random factor
mixed_model(data = data_doubling_time,
    Y_value = "Doubling_time",
    Fixed_Factor = "Student",
    Random_Factor = "Experiment")

# two fixed factors as a vector, one random factor
mixed_model(data = data_cholesterol,
    Y_value = "Cholesterol",
    Fixed_Factor = c("Treatment", "Hospital"),
    Random_Factor = "Subject")

# save model
model <- mixed_model(data = data_doubling_time,
    Y_value = "Doubling_time",
    Fixed_Factor = "Student",
    Random_Factor = "Experiment")

# get model summary
summary(model)

mixed_model_slopes

Model from a linear mixed effects model with varying slopes

Description

There are four related functions for mixed effects analyses: mixed_model, mixed_anova, mixed_model_slopes, and mixed_anova_slopes.

Usage

mixed_model_slopes(
    data,
    Y_value,
    Fixed_Factor,
    Slopes_Factor,
    Random_Factor,
    ...
)
mixed_model_slopes

Arguments

- **data**: a data table object, e.g. data.frame or tibble.
- **Y_value**: name of column containing quantitative (dependent) variable, provided within "quotes".
- **Fixed_Factor**: name(s) of categorical fixed factors (independent variables) provided as a vector if more than one or within "quotes".
- **Slopes_Factor**: name of factor to allow varying slopes on.
- **Random_Factor**: name(s) of random factors to allow random intercepts; to be provided as a vector when more than one or within "quotes".
- **...**: any additional arguments to pass on to `lmer` if required.

Details

This function uses `lmer` to fit a linear mixed effect model and provides the model object, which could be used for post-hoc comparisons. The model object is converted to class `lmerModLmerTest` object by `as_lmerModLmerTest`. It requires a data table, one dependent variable (`Y_value`), one or more independent variables (`Fixed_Factor`). Exactly one random factor (`Random_Factor`) and Slope_Factor should be provided. This function is related to `mixed_anova_slopes`. Output of this function can be used with `posthoc_Pairwise`, `posthoc_Levelwise` and `posthoc_vsRef`, or with `emmeans`.

More than one fixed factors can be provided as a vector (e.g. c("A", "B")). A full model with interaction term is fitted with one term each for varying slopes and intercepts. This means when `Y_value = Y,Fixed_factor = c("A","B"),Slopes_Factor = "S",Random_factor = "R"` are entered as arguments, these are passed on as `Y ~ A*B + (S|R)` (which is equivalent to `Y ~ A + B + A:B + (S|R)`). In this experimental implementation, random slopes and intercepts are fitted `(Slopes_Factor|Random_Factor)`. Only one term each is allowed for Slopes_Factor and Random_Factor.

Value

This function returns an S4 object of class "lmerModLmerTest".

Examples

```r
#two fixed factors as a vector,
#exactly one slope factor and random factor
mod <- mixed_model_slopes(data = data_2w_Tdeath, 
Y_value = "PI", 
Fixed_Factor = c("Genotype", "Time"), 
Slopes_Factor = "Time", 
Random_Factor = "Experiment")
# get summary
summary(mod)
```
Plot a scatter graph with matched shapes on a bar plot using three variables.

Description

The functions `plot_3d_scatterbar`, `plot_3d_scatterbox`, `plot_4d_scatterbar` and `plot_4d_scatterbox` are useful for plotting one-way or two-way ANOVA designs with randomised blocks or repeated measures. The blocks or subjects can be mapped to the shapes argument in both functions (up to 25 levels can be mapped to shapes; there will be an error if this number is exceeded). The 3d versions use the categorical variable (xcol) for grouping (e.g. one-way ANOVA designs), and 4d versions take an additional grouping variable (e.g. two-way ANOVA designs) that is passed to either boxes or bars argument.

Usage

```r
plot_3d_scatterbar(
  data,
  xcol,
  ycol,
  shapes,
  ewid = 0.2,
  symsize = 2.5,
  symthick = 1,
  jitter = 0.1,
  fontsize = 20,
  b_alpha = 1,
  s_alpha = 1,
  ColSeq = TRUE,
  ColPal = "all_grafify",
  ColRev = FALSE,
  TextXAngle = 0
)
```

Arguments

data  a data table, e.g. data.frame or tibble.
xcol  name of the column with the categorical factor to be plotted on X axis.
ycol  name of the column with quantitative variable to plot on the Y axis.
shapes  name of the column with the second categorical factor, for example from a two-way ANOVA design.
ewid  width of error bars, default set to 0.2.
symsize  size of symbols, default set to 3.
symthick  size of outline of symbol lines (stroke = 1.5), default set to 1.5
plot_3d_scatterbar

jitter  extent of jitter (scatter) of symbols, default is 0.1. Increase to reduce symbol overlap, set to 0 for aligned symbols.

fontsize  parameter of base_size of fonts in theme_classic, default set to size 20.

b_alpha  fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).

s_alpha  fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).

ColSeq  logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.

ColPal  grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".

ColRev  whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).

TextXAngle  orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

Details

These functions rely on ggplot with geom_point and geom_bar (through stat_summary) or geom_boxplot geometries.

Variables other than the quantitative variable (ycol) will be automatically converted to categorical variables even if they are numeric in the data table.

Shapes are always plotted in black colour, and their opacity can be changed with the s_alpha argument and overlap can be reduced with the jitter argument. Other arguments are similar to other plot functions as briefly explained below.

Bars depict means using stat_summary with geom = "bar", fun = "mean", and bar width is set to 0.7 (cannot be changed). Error bar width can be changed with the ewid argument.

Boxplot geometry uses geom_boxplot with position = position_dodge(width = 0.9), width = 0.6. The thick line within the boxplot depicts the median, the box the IQR (interquantile range) and the whiskers show 1.5*IQR.

In 4d versions, the two grouping variables (i.e. xcol and either boxes or bars) are passed to geplot aesthetics through group = interaction{ xcol, shapes}.

Colours can be changed using ColPal, ColRev or ColSeq arguments. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

All four functions can be expanded further, for example with facet_grid or facet_wrap.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".
Examples

#3d version for 1-way data with blocking
plot_3d_scatterbox(data = data_1w_death,
xcol = Genotype, ycol = Death, shapes = Experiment)
#compare above graph to
plot_scatterbox(data = data_1w_death, xcol = Genotype, ycol = Death)

#4d version for 2-way data with blocking
plot_4d_scatterbox(data = data_2w_Tdeath,
xcol = Genotype,
ycol = PI,
boxes = Time,
shapes = Experiment)

plot_4d_scatterbar(data = data_2w_Festing,
xcol = Strain,
ycol = GST,
bars = Treatment,
shapes = Block)

---

plot_3d_scatterbox  *Plot a scatter and box plot with matched symbols.*

Description

The functions `plot_3d_scatterbar, plot_3d_scatterbox, plot_4d_scatterbar` and `plot_4d_scatterbox` are useful for plotting one-way or two-way ANOVA designs with randomised blocks or repeated measures. The blocks or subjects can be mapped to the `shapes` argument in both functions (up to 25 levels can be mapped to `shapes`; there will be an error if this number is exceeded). The 3d versions use the categorical variable (`xcol`) for grouping (e.g. one-way ANOVA designs), and 4d versions take an additional grouping variable (e.g. two-way ANOVA designs) that is passed to either `boxes` or `bars` argument.

Usage

```r
plot_3d_scatterbox(
  data,
  xcol,
  ycol,
  shapes,
  symsize = 2.5,
  symthick = 1,
  jitter = 0.1,
  fontsize = 20,
  b_alpha = 1,
  s_alpha = 1,
  ColSeq = TRUE,
)```
plot_3d_scatterbox

ColPal = "all_grafify",
ColRev = FALSE,
TextXAngle = 0,
...
)

Arguments

data
- a data table, e.g. data.frame or tibble.
xcol
- name of the column with the categorical factor to be plotted on X axis. If your table has numeric X, enter xcol = factor(name of column).
ycol
- name of the column with quantitative variable to plot on the Y axis.
shapes
- name of the column with the second categorical factor in a two-way ANOVA design.
syssize
- size of symbols, default set to 3.
synthick
- size of outline of symbol lines (stroke = 1.0), default set to 1.0.
jitter
- extent of jitter (scatter) of symbols, default is 0.1. Increase to reduce symbol overlap, set to 0 for aligned symbols.
fontsize
- parameter of base_size of fonts in theme_classic, default set to size 20.
b_alpha
- fractional opacity of boxes, default set to 1 (i.e. maximum opacity & zero transparency).
s_alpha
- fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
ColSeq
- logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColPal
- grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColRev
- whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle
- orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
...
- any additional arguments to pass to ggplot2geom_boxplot.

Details

These functions rely on ggplot with geom_point and geom_bar (through stat_summary) or geom_boxplot geometries.

Variables other than the quantitative variable (ycol) will be automatically converted to categorical variables even if they are numeric in the data table.

Shapes are always plotted in black colour, and their opacity can be changed with the s_alpha argument and overlap can be reduced with the jitter argument. Other arguments are similar to other plot functions as briefly explained below.

Bars depict means using stat_summary with geom = "bar", fun = "mean", and bar width is set to 0.7 (cannot be changed). Error bar width can be changed with the ewid argument.
Boxplot geometry uses `geom_boxplot` with `position = position_dodge(width = 0.9), width = 0.6`. The thick line within the boxplot depicts the median, the box the IQR (interquantile range) and the whiskers show 1.5*IQR.

In 4d versions, the two grouping variables (i.e. `xcol` and either `boxes` or `bars`) are passed to `ggplot` aesthetics through `group = interaction(xcol, shapes)`.

Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq` (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

All four functions can be expanded further, for example with `facet_grid` or `facet_wrap`.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```r
# 3d version for 1-way data with blocking
plot_3d_scatterbox(data = data_1w_death,
xcol = Genotype, ycol = Death, shapes = Experiment)
# compare above graph to
plot_scatterbox(data = data_1w_death, xcol = Genotype, ycol = Death)

# 4d version for 2-way data with blocking
plot_4d_scatterbox(data = data_2w_Tdeath,
xcol = Genotype,
ycol = PI,
boxes = Time,
shapes = Experiment)

plot_4d_scatterbar(data = data_2w_Festing,
xcol = Strain,
ycol = GST,
bars = Treatment,
shapes = Block)
```

Description

The functions `plot_3d_scatterbar`, `plot_3d_scatterbox`, `plot_3d_scatterviolin`, `plot_4d_scatterbar`, `plot_4d_scatterbox` and `plot_4d_scatterviolin` are useful for plotting one-way or two-way ANOVA designs with randomised blocks or repeated measures. The blocks or subjects can be mapped to the `shapes` argument in both functions (up to 25 levels can be mapped to `shapes`; there will be an error if this number is exceeded). The 3d versions use the categorical variable (`xcol`)
for grouping (e.g. one-way ANOVA designs), and 4d versions take an additional grouping variable (e.g. two-way ANOVA designs) that is passed to either boxes or bars argument.

Usage

```r
plot_3d_scatterviolin(
  data,
  xcol,
  ycol,
  shapes,
  symsize = 2.5,
  s_alpha = 1,
  symthick = 1,
  v_alpha = 1,
  b_alpha = 1,
  bwid = 0.5,
  bvthick = 1,
  jitter = 0.1,
  fontsize = 20,
  ColSeq = TRUE,
  ColPal = "all_grafify",
  ColRev = FALSE,
  TextXAngle = 0,
  scale = "width",
  trim = TRUE,
  ...
)
```

Arguments

data a data table, e.g. data.frame or tibble.
xcol name of the column with the categorical factor to be plotted on X axis. If your table has numeric X, enter xcol = factor(name of column).
ycol name of the column with quantitative variable to plot on the Y axis.
shapes name of the column with the second categorical factor in a two-way ANOVA design.
symsize size of symbols, default set to 3.
s_alpha fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency). Set s_alpha = 0 to not show scatter plot.
symthick size of outline of symbol lines (stroke = 1.0), default set to 1.0.
v_alpha fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency).
b_alpha fractional opacity of boxplots, default set to 1 (i.e. maximum opacity & zero transparency). For white boxplots inside violins, set b_alpha = 0.
bwid width of boxplots; default 0.2
bvthick thickness of both violin and boxplot lines; default 1.
**plot_3d_scatterviolin**

- **jitter**: extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
- **fontsize**: parameter of base_size of fonts in theme_classic, default set to size 20.
- **ColSeq**: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
- **ColPal**: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted", "dark", "light".
- **ColRev**: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- **scale**: set to "area" by default, can be changed to "count" or "width".
- **trim**: set whether tips of violin plot should be trimmed at high/low data. Default trim = T, can be changed to F.
- **...**: any additional arguments to pass to `ggplot2geom_boxplot` or `ggplot2geom_violin`.

**Details**

These functions rely on `ggplot` with `geom_point` and `geom_bar` (through `stat_summary`), or `geom_violin` and `geom_boxplot` geometries.

Variables other than the quantitative variable (`ycol`) will be automatically converted to categorical variables even if they are numeric in the data table.

Shapes are always plotted in black colour, and their opacity can be changed with the `s_alpha` argument and overlap can be reduced with the `jitter` argument. Other arguments are similar to other plot functions as briefly explained below.

Bars depict means using `stat_summary` with `geom = "bar", fun = "mean"` , and bar width is set to 0.7 (cannot be changed). Error bar width can be changed with the `ewid` argument.

Boxplot geometry uses `geom_boxplot` with `position = position_dodge(width = 0.9),width = 0.6`. The thick line within the boxplot depicts the median, the box the IQR (interquantile range) and the whiskers show 1.5*IQR.

In 4d versions, the two grouping variables (i.e. `xcol` and either `boxes` or `bars`) are passed to `ggplot` aesthetics through `group = interaction(xcol,shapes)`.

Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq` (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

All four functions can be expanded further, for example with `facet_grid` or `facet_wrap`.

**Value**

This function returns a `ggplot2` object of class "gg" and "ggplot".
Examples

#3d version for 1-way data with blocking
plot_3d_scatterviolin(data = data_1w_death,
xcol = Genotype, ycol = Death, shapes = Experiment,
b_alpha = 0)
#compare above graph to
plot_scatterviolin(data = data_1w_death,
xcol = Genotype, ycol = Death)

#4d version for 2-way data with blocking
plot_4d_scatterviolin(data = data_2w_Tdeath,
xcol = Genotype,
ycol = PI,
boxes = Time,
shapes = Experiment, b_alpha = 0)

Description

The functions `plot_3d_scatterbar`, `plot_3d_scatterbox`, `plot_4d_scatterbar` and `plot_4d_scatterbox` are useful for plotting one-way or two-way ANOVA designs with randomised blocks or repeated measures. The blocks or subjects can be mapped to the shapes argument in both functions (up to 25 levels can be mapped to shapes; there will be an error if this number is exceeded). The 3d versions use the categorical variable (xcol) for grouping (e.g. one-way ANOVA designs), and 4d versions take an additional grouping variable (e.g. two-way ANOVA designs) that is passed to either boxes or bars argument.

Usage

plot_4d_scatterbar(
data,
xcol,
ycol,
bars,
shapes,
symsize = 2.5,
symthick = 1,
jitter = 0.1,
ewid = 0.2,
fontsize = 20,
b_alpha = 1,
s_alpha = 1,
ColPal = "all_grafify",
ColRev = FALSE,
ColSeq = TRUE,
TextXAngle = 0
)

**Arguments**

- **data**: a data table, e.g. data.frame or tibble.
- **xcol**: name of the column with the categorical factor to plot on X axis. If column is numeric, enter as `factor(col)`.
- **ycol**: name of the column to plot on quantitative variable on the Y axis.
- **bars**: name of the column containing grouping within the factor plotted on X axis. Can be categorical or numeric X. If your table has numeric X and you want to plot as factor, enter `xcol = factor(name of column)`.
- **shapes**: name of the column that contains matched observations, e.g. subject IDs, experiment ID.
- **symsize**: size of symbols, default set to 3.
- **symthick**: size of outline of symbol lines (`stroke = 1.0`), default set to 1.0
- **jitter**: extent of jitter (scatter) of symbols, default is 0.1. Increase to reduce symbol overlap, set to 0 for aligned symbols.
- **ewid**: width of error bars, default set to 0.2.
- **fontsize**: parameter of `base_size` of fonts in `theme_classic`, default set to size 20.
- **b_alpha**: fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
- **s_alpha**: fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
- **ColPal**: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
- **ColRev**: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
- **ColSeq**: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

**Details**

These functions rely on `ggplot` with `geom_point` and `geom_bar` (through `stat_summary`) or `geom_boxplot` geometries.

Variables other than the quantitative variable (`ycol`) will be automatically converted to categorical variables even if they are numeric in the data table.

Shapes are always plotted in black colour, and their opacity can be changed with the `s_alpha` argument and overlap can be reduced with the `jitter` argument. Other arguments are similar to other plot functions as briefly explained below.

Bars depict means using `stat_summary` with `geom = "bar", fun = "mean"`, and bar width is set to 0.7 (cannot be changed). Error bar bar width can be changed with the `ewid` argument.
Boxplot geometry uses `geom_boxplot` with `position = position_dodge(width = 0.9), width = 0.6`. The thick line within the boxplot depicts the median, the box the IQR (interquantile range) and the whiskers show 1.5*IQR.

In 4d versions, the two grouping variables (i.e. `xcol` and either `boxes` or `bars`) are passed to `ggplot` aesthetics through `group = interaction(xcol, shapes)`.

Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq` (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

All four functions can be expanded further, for example with `facet_grid` or `facet_wrap`.

**Value**

This function returns a `ggplot2` object of class "gg" and "ggplot".

**Examples**

#3d version for 1-way data with blocking
```r
plot_3d_scatterbox(data = data_1w_death, xcol = Genotype, ycol = Death, shapes = Experiment)
#compare above graph to
plot_scatterbox(data = data_1w_death, xcol = Genotype, ycol = Death)
```

#4d version for 2-way data with blocking
```r
plot_4d_scatterbox(data = data_2w_Tdeath, xcol = Genotype, ycol = PI, boxes = Time, shapes = Experiment)

plot_4d_scatterbar(data = data_2w_Festing, xcol = Strain, ycol = GST, bars = Treatment, shapes = Block)
```

**Description**

The functions `plot_3d_scatterbar`, `plot_3d_scatterbox`, `plot_4d_scatterbar` and `plot_4d_scatterbox` are useful for plotting one-way or two-way ANOVA designs with randomised blocks or repeated measures. The blocks or subjects can be mapped to the `shapes` argument in both functions (up to 25 levels can be mapped to `shapes`; there will be an error if this number is exceeded). The 3d versions use the categorical variable (`xcol`) for grouping (e.g. one-way ANOVA designs), and 4d versions...
take an additional grouping variable (e.g. two-way ANOVA designs) that is passed to either boxes or bar's argument.

Usage

```
plot_4d_scatterbox(
  data,
  xcol,
  ycol,
  boxes,
  shapes,
  symsize = 2.5,
  symthick = 1,
  jitter = 0.1,
  fontsize = 20,
  b_alpha = 1,
  s_alpha = 1,
  ColSeq = TRUE,
  ColPal = "all_grafify",
  ColRev = FALSE,
  TextXAngle = 0,
  ...
)
```

Arguments

data a data table, e.g. data.frame or tibble.

xcol name of the column with the categorical factor to plot on X axis. If column is numeric, enter as `factor(col)`.

ycol name of the column to plot on quantitative variable on the Y axis.

boxes name of the column containing grouping within the factor plotted on X axis. Can be categorical or numeric X. If your table has numeric X and you want to plot as factor, enter `xcol = factor(name of column)`.

shapes name of the column that contains matched observations, e.g. subject IDs, experiment number etc.

symsize size of symbols, default set to 3.

symthick size of outline of symbol lines (`stroke = 1.0`), default set to 1.0.

jitter extent of jitter (scatter) of symbols, default is 0.1. Increase to reduce symbol overlap, set to 0 for aligned symbols.

fontsize parameter of `base_size` of fonts in `theme_classic`, default set to size 20.

b_alpha fractional opacity of boxes, default set to 1 (i.e. maximum opacity & zero transparency).

s_alpha fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).

ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`. 
ColPal

grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".

ColRev

whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).

TextXAngle

orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

... any additional arguments to pass to ggplot2 geom_boxplot.

Details

These functions rely on ggplot with geom_point and geom_bar (through stat_summary) or geom_boxplot geometries.

Variables other than the quantitative variable (ycol) will be automatically converted to categorical variables even if they are numeric in the data table.

Shapes are always plotted in black colour, and their opacity can be changed with the s_alpha argument and overlap can be reduced with the jitter argument. Other arguments are similar to other plot functions as briefly explained below.

Bars depict means using stat_summary with geom = "bar", fun = "mean", and bar width is set to 0.7 (cannot be changed). Error bar width can be changed with the ewid argument.

Boxplot geometry uses geom_boxplot with position = position_dodge(width = 0.9), width = 0.6. The thick line within the boxplot depicts the median, the box the IQR (interquantile range) and the whiskers show 1.5*IQR.

In 4d versions, the two grouping variables (i.e. xcol and either boxes or bars) are passed to ggplot aesthetics through group = interaction(xcol,shapes).

Colours can be changed using ColPal, ColRev or ColSeq arguments. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first within the chosen palette. ColSeq (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

All four functions can be expanded further, for example with facet_grid or facet_wrap.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

#3d version for 1-way data with blocking
plot_3d_scatterbox(data = data_1w_death, xcol = Genotype, ycol = Death, shapes = Experiment)
#compare above graph to
plot_scatterbox(data = data_1w_death, xcol = Genotype, ycol = Death)

#4d version for 2-way data with blocking
plot_4d_scatterbox(data = data_2w_Tdeath, xcol = Genotype, ycol = PI,
plot_4d_scatterviolin

Plot a dot plot with matched shapes on a violin & box plot using four variables.

Description

The functions plot_3d_scatterbar, plot_3d_scatterbox, plot_3d_scatterviolin, plot_4d_scatterbar, plot_4d_scatterbox and plot_4d_scatterviolin are useful for plotting one-way or two-way ANOVA designs with randomised blocks or repeated measures. The blocks or subjects can be mapped to the shapes argument in both functions (up to 25 levels can be mapped to shapes; there will be an error if this number is exceeded). The 3d versions use the categorical variable (xcol) for grouping (e.g. one-way ANOVA designs), and 4d versions take an additional grouping variable (e.g. two-way ANOVA designs) that is passed to either boxes or bars argument.

Usage

plot_4d_scatterviolin(
  data,
  xcol,
  ycol,
  boxes,
  shapes,
  symsize = 2.5,
  s_alpha = 1,
  symthick = 1,
  jitter = 0.1,
  v_alpha = 1,
  b_alpha = 1,
  bvthick = 1,
  bwid = 0.2,
  ColSeq = TRUE,
  ColPal = "all_grafify",
  ColRev = FALSE,
  TextXAngle = 0,
  scale = "width",
  trim = TRUE,
  fontsize = 20,
  ...
)
**Arguments**

- **data**: a data table, e.g. data.frame or tibble.
- **xcol**: name of the column with the categorical factor to plot on X axis. If column is numeric, enter as `factor(col)`.
- **ycol**: name of the column to plot on quantitative variable on the Y axis.
- **boxes**: name of the column containing grouping within the factor plotted on X axis. Can be categorical or numeric X. If your table has numeric X and you want to plot as factor, enter `xcol = factor(name of column)`.
- **shapes**: name of the column that contains matched observations, e.g. subject IDs, experiment number etc.
- **symsize**: size of symbols, default set to 3.
- **s_alpha**: fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency). Set `s_alpha = 0` to not show scatter plot.
- **symthick**: size of outline of symbol lines \( (stroke = 1.0) \), default set to 1.0.
- **jitter**: extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
- **v_alpha**: fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency).
- **b_alpha**: fractional opacity of boxplots, default set to 1 (i.e. maximum opacity & zero transparency). For white boxplots inside violins, set `b_alpha = 0`.
- **bvthick**: thickness of both violin and boxplot lines; default 1.
- **bwid**: width of boxplots; default 0.2
- **ColSeq**: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
- **ColPal**: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
- **ColRev**: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- **scale**: set to "area" by default, can be changed to "count" or "width".
- **trim**: set whether tips of violin plot should be trimmed at high/low data. Default trim = T, can be changed to F.
- **fontsize**: parameter of base_size of fonts in theme_classic, default set to size 20.
- **...**: any additional arguments to pass to `ggplot2geom_boxplot` or `ggplot2geom_violin`.

**Details**

These functions rely on `ggplot` with `geom_point` and `geom_bar` (through `stat_summary`), or `geom_violin` and `geom_boxplot` geometries.

Variables other than the quantitative variable (ycol) will be automatically converted to categorical variables even if they are numeric in the data table.
Shapes are always plotted in black colour, and their opacity can be changed with the `s_alpha` argument and overlap can be reduced with the `jitter` argument. Other arguments are similar to other plot functions as briefly explained below.

Bars depict means using `stat_summary` with `geom = "bar", fun = "mean"`, and bar width is set to 0.7 (cannot be changed). Error bar width can be changed with the `ewid` argument.

Boxplot geometry uses `geom_boxplot` with `position = position_dodge(width = 0.9), width = 0.6`. The thick line within the boxplot depicts the median, the box the IQR (interquantile range) and the whiskers show 1.5*IQR.

In 4d versions, the two grouping variables (i.e. `xcol` and either `boxes` or `bars`) are passed to `ggplot` aesthetics through `group = interaction{ xcol, shapes}`.

Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq` (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

All four functions can be expanded further, for example with `facet_grid` or `facet_wrap`.

**Value**

This function returns a ggplot2 object of class "gg" and "ggplot".

**Examples**

```r
#3d version for 1-way data with blocking
plot_3d_scatterviolin(data = data_1w_death, 
  xcol = Genotype, ycol = Death, shapes = Experiment, 
  b_alpha = 0)
#compare above graph to
plot_scatterviolin(data = data_1w_death, 
  xcol = Genotype, ycol = Death)

#4d version for 2-way data with blocking
plot_4d_scatterviolin(data = data_2w_Tdeath, 
  xcol = Genotype, ycol = PI, 
  boxes = Time, 
  shapes = Experiment, b_alpha = 0)
```

---

**Description**

This function takes a data table, categorical X and numeric Y variables, and plots bars showing the mean with SD error bars. The X variable is mapped to the `fill` aesthetic of bars. The related `plot_bar_sd_sc` plots bars with a single colour.
Usage

plot_bar_sd(
  data,
  xcol,
  ycol,
  b_alpha = 1,
  bwid = 0.7,
  bthick = 1,
  ewid = 0.3,
  ColPal = "all_grafify",
  ColSeq = TRUE,
  ColRev = FALSE,
  TextXAngle = 0,
  fontsize = 20,
  ...
)

Arguments

data a data table object, e.g. a data.frame or tibble.
xcol name of the column to plot on X axis. This should be a categorical variable.
ycol name of the column to plot on the Y axis. This should be a quantitative variable.
b_alpha fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
bwid width of bars, default 0.7
bthick thickness of bar borders; default 1.
ewid width of error bars, default 0.3
ColPal grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColRev whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize parameter of base_size of fonts in theme_classic, default set to size 20.
... any additional arguments to pass to stat_summary.

Details

The function uses stat_summary with geom = "bar". Standard deviation (SD) is plotted through stat_summary calculated using mean_sdl from the ggplot2 package (get help with ?mean_sdl), and 1x SD is plotted (fun.arg = list(mult = 1)).

Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light".
"light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

You are instead encouraged to show all data using the following functions: plot_scatterbar_sd, plot_scatterbox, plot_dotbox, plot_dotbar_sd, plot_scatterviolin or plot_dotviolin.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

# Basic usage
plot_bar_sd(data = data_doubling_time, 
            xcol = Student, ycol = Doubling_time)

# apply distant colours in the default palette
plot_bar_sd(data = data_doubling_time, 
            xcol = Student, ycol = Doubling_time, 
            ColSeq = FALSE)

plot_bar_sd_sc

Plot a bar graph indicating mean with error bars (SD) using two variables.

Description

This function is related to plot_bar_sd, but this one maps a single or same colour, therefore _sc. The only new argument is colour, which can be any hexcode or name of colours in the all_grafify palette. The default colour is ok_orange. ColPal and ColRev arguments are not available. Colours available can be seen quickly with plot_grafify_palette.

Usage

plot_bar_sd_sc(
    data, 
    xcol, 
    ycol, 
    colour = "ok_orange", 
    b_alpha = 1, 
    bwid = 0.7, 
    bthick = 1, 
    ewid = 0.3, 
    TextXAngle = 0, 
    fontsize = 20, 
    ...
)
Arguments

- **data**: a data table object, e.g. a data.frame or tibble.
- **xcol**: name of the column to plot on X axis. This should be a categorical variable.
- **ycol**: name of the column to plot on the Y axis. This should be a quantitative variable.
- **colour**: colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is ok_orange.
- **b_alpha**: fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
- **bwid**: width of bars (default 0.7).
- **bthick**: thickness of bar borders; default 1.
- **ewid**: width of error bars, default 0.3.
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- **fontsize**: parameter of base_size of fonts in theme_classic, default set to size 20.
- **...**: any additional arguments to pass to **stat_summary**.

Details

You are instead encouraged to show all data using the following functions: **plot_scatterbar_sd**, **plot_scatterbox**, **plot_dotbox**, **plot_dotbar_sd**, **plot_scatterviolin** or **plot_dotviolin**.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```r
plot_bar_sd_sc(data = data_doubling_time,
               xcol = Student, ycol = Doubling_time,
               colour = "ok_grey")
```

Description

The **plot_befafter_colours**, **plot_befafter_colors** and **plot_befafter_shapes** are for plotting matched data joined by lines. These functions take X and Y variables along with a data column with matching information (e.g. matched subjects or experiments etc.) and plot symbols matched by colour or shape.
Usage

plot_befafter_colors(
  data,
  xcol,
  ycol,
  match,
  symsize = 3,
  symthick = 1,
  s_alpha = 1,
  ColPal = "all_grafify",
  ColSeq = TRUE,
  ColRev = FALSE,
  TextXAngle = 0,
  fontsize = 20,
  groups,
  ...
)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column containing the categorical variable to be plotted on the X axis.
ycol name of the column containing the quantitative Y values.
match name of the column with the grouping variable to pass on to geom_line.
symsize size of symbols, default set to 3.
symthick thickness of symbol border, default set to 1.
s_alpha fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
ColPal grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColRev whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize parameter of base_size of fonts in theme_classic, default set to size 20.
groups old argument name for match; retained for backward compatibility.
... any additional arguments to pass to ggpplot2geom_line.
Details

Note that only 25 shapes are available, and there will be errors with `plot_befafter_shapes` when there are fewer than 25 matched observations; instead use `plot_befafter_colours` instead.

Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq` decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

More complex designs can also be plotted when used with `facet_wrap` or `facet_grid`.

Value

This function returns a `ggplot2` object of class "gg" and "ggplot".

Examples

```r
#plot without legends if necessary
plot_befafter_colors(data = data_t_pdiff,
                      xcol = Condition, ycol = Mass,
                      match = Subject, s_alpha = .9, ColSeq = FALSE)+
guides(fill = "none",
       colour = "none") #remove guides

#2way ANOVA design with randomised blocks
plot_befafter_colors(data = data_2w_Tdeath,
                      xcol = Genotype, ycol = PI,
                      match = Experiment) + facet_wrap("Time")
```

plot_befafter_colours  Plot a before-after plot with lines joining colour-matched symbols.

Description

The `plot_befafter_colours`, `plot_befafter_colors` and `plot_befafter_shapes` are for plotting matched data joined by lines. These functions take X and Y variables along with a data column with matching information (e.g. matched subjects or experiments etc.) and plot symbols matched by colour or shape.

Usage

```r
plot_befafter_colours(
  data,
  xcol,
  ycol,
  match,
  symsize = 3,
  symthick = 1,
```

s_alpha = 1,
ColPal = "all_grafify",
ColSeq = TRUE,
ColRev = FALSE,
TextXAngle = 0,
fontsize = 20,
groups,
...)

Arguments

data  a data table object, e.g. data.frame or tibble.
xcol   name of the column containing the categorical variable to be plotted on the X axis.
ycol   name of the column containing the quantitative Y values.
match  name of the column with the matching variable to pass on to geom_line.
szize  size of symbols, default set to 3.
synthick thickness of symbol border, default set to 1.
s_alpha  fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
ColPal  grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted", "dark", "light".
ColSeq  logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColRev  whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle  orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize  parameter of base_size of fonts in theme_classic, default set to size 20.
groups  old argument name for match; retained for backward compatibility.
...  any additional arguments to pass to ggplot2geom_line.

Details

Note that only 25 shapes are available, and there will be errors with plot_befafter_shapes when there are fewer than 25 matched observations; instead use plot_befafter_colours instead.

Colours can be changed using ColPal, ColRev or ColSeq arguments. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

More complex designs can also be plotted when used with facet_wrap or facet_grid.
plot_befafter_shapes

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

#plot without legends if necessary
plot_befafter_colors(data = data_t_pdiff,
xcol = Condition, ycol = Mass,
match = Subject, s_alpha = .9, ColSeq = FALSE)+
guides(fill = "none",
colour = "none") #remove guides
#2way ANOVA design with randomised blocks
plot_befafter_colors(data = data_2w_Tdeath,
xcol = Genotype, ycol = PI,
match = Experiment) + facet_wrap("Time")

plot_befafter_shapes  Plot a before-after plot with lines joining shape-matched symbols.

Description

The plot_befafter_colours, plot_befafter_colors and plot_befafter_shapes are for plotting matched data joined by lines. These functions take X and Y variables along with a data column with matching information (e.g. matched subjects or experiments etc.) and plot symbols matched by colour or shape.

Usage

plot_befafter_shapes(
  data,
  xcol,
  ycol,
  match,
  symsize = 3,
  symthick = 1,
  s_alpha = 0.8,
  ColPal = "all_grafify",
  ColSeq = TRUE,
  ColRev = FALSE,
  TextXAngle = 0,
  fontsize = 20,
  groups,
  ...
)
plot_befafter_shapes

Arguments

data: a data table object, e.g. data.frame or tibble.
xcol: name of the column containing the categorical variable to be plotted on the X axis.
ycol: name of the column containing the quantitative Y values.
match: name of the column with the matching variable to pass on to geom_line.
symsize: size of symbols, default set to 3.
symthick: size of outline of symbol lines (stroke = 1), default set to 1.
s_alpha: fractional opacity of symbols, default set to 0.8 (i.e. 80% opacity).
ColPal: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted", "dark", "light".
ColSeq: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColRev: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize: parameter of base_size of fonts in theme_classic, default set to size 20.
groups: old argument name for match; retained for backward compatibility.
...: any additional arguments to pass to ggplot2geom_line.

Details

Note that only 25 shapes are available, and there will be errors with plot_befafter_shapes when there are fewer than 25 matched observations; instead use plot_befafter_colours instead.

Colours can be changed using ColPal, ColRev or ColSeq arguments. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

More complex designs can also be plotted when used with facet_wrap or facet_grid.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

#plot without legends if necessary
plot_befafter_colors(data = data_t_pdiff, xcol = Condition, ycol = Mass, match = Subject, s_alpha = .9, ColSeq = FALSE)+ guides(fill = "none", colour = "none") #remove guides
#2way ANOVA design with randomised blocks
plot_befafter_colors(data = data_2w_Tdeath, 
xcol = Genotype, ycol = PI,  
match = Experiment) + facet_wrap("Time")

### plot_density

Plot density distribution of data.

**Description**

This function takes a data table, `ycol` of quantitative variable and a categorical grouping variable (`group`), if available, and plots a density graph using `geom_density()`.

**Usage**

```r
plot_density(data, 
ycol, 
group, 
linethick = 1, 
c_alpha = 0.2, 
ColPal = "all_grafify", 
ColSeq = TRUE, 
ColRev = FALSE, 
TextXAngle = 0, 
fontsize = 20, 
Group, 
alpha, 
...)
```

**Arguments**

- `data`: a data table e.g. data.frame or tibble.
- `ycol`: name of the column containing the quantitative variable whose density distribution is to be plotted.
- `group`: name of the column containing a categorical grouping variable
- `linethick`: thickness of symbol border, default set to 1.
- `c_alpha`: fractional opacity of filled colours under the curve, default set to 0.2 (i.e. 20% opacity).
- `ColPal`: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
- `ColSeq`: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
- `ColRev`: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

fontsize parameter of base_size of fonts in theme_classic, default set to size 20.

Group deprecated old argument for group; retained for backward compatibility.

alpha deprecated old argument for c_alpha; retained for backward compatibility.

... any additional arguments to pass to ggplot2geom_density.

Details

Note that the function requires the quantitative Y variable first, and groups them based on an X variable. The group variable is mapped to the fill and colour aesthetics in geom_density. Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

plot_density(data = data_t_pratio, ycol = log(Cytokine), group = Genotype)

# with faceting
plot_density(data = data_cholesterol, ycol = Cholesterol, group = Treatment, fontsize = 10)+facet_wrap("Treatment")

plot_dotbar_sd

Plot a dotplot on a bar graph with SD error bars with two variables.

Description

This function takes a data table, X and Y variables, and plots a graph with a dotplot and bars using stat_summary with geom = "bar", and geom_dotplot geometries. Standard deviation (SD) is plotted through stat_summary calculated using mean_sdl from the ggplot2 package (get help with ?mean_sdl), and 1x SD is plotted (fun.arg = list(mult = 1)).
**plot_dotbar_sd**

**Usage**

```r
plot_dotbar_sd(
  data,
  xcol,
  ycol,
  dotsize = 1.5,
  dotthick = 1,
  bwid = 0.7,
  ewid = 0.2,
  b_alpha = 1,
  d_alpha = 1,
  ColPal = "all_grafify",
  ColRev = FALSE,
  ColSeq = TRUE,
  TextXAngle = 0,
  fontsize = 20,
  ...
)
```

**Arguments**

- `data`: a data table object, e.g. data.frame or tibble.
- `xcol`: name of the column to plot on X axis. This should be a categorical variable.
- `ycol`: name of the column to plot on quantitative Y axis. This should be a quantitative variable.
- `dotsize`: size of dots relative to binwidth used by `geom_dotplot`. Default set to 1.5, increase/decrease as needed.
- `dotthick`: thickness of dot border (stroke parameter of `geom_dotplot`), default set to 1.
- `bwid`: width of bars, default set to 0.7
- `ewid`: width of error bars, default set to 0.2.
- `b_alpha`: fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
- `d_alpha`: fractional opacity of dots, default set to 1 (i.e. maximum opacity & zero transparency).
- `ColPal`: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
- `ColRev`: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
- `ColSeq`: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
- `TextXAngle`: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- `...`: any additional arguments to pass to `ggplot2geom_dotplot`. 
plot_dotbar_sd_sc

Details
The X variable is mapped to the fill aesthetic in both bar and dotplot.
Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette. The size of dots can be adjusted using the parameter, which is dotsize = 1 by default.
This function is related to plot_scatterbar_sd, plot_dotbar_sd and plot_dotviolin.

Value
This function returns a ggplot2 object of class "gg" and "ggplot".

Examples
plot_dotbar_sd(data = data_cholesterol, xcol = Treatment, ycol = Cholesterol)
plot_dotbar_sd(data = data_1w_death, xcol = Genotype, ycol = Death, ColPal = "pale", ColSeq = FALSE, ColRev = TRUE)

plot_dotbar_sd_sc(data, xcol, ycol, colour = "ok_orange", dotsize = 1.5, dotthick = 1, bwid = 0.7, ewid = 0.2, b_alpha = 1, d_alpha = 1)

plot_dotbar_sd_sc
Plot a dotplot on a bar graph with SD error bars with two variables.

Description
This function is related to plot_dotbar_sd, but this one maps a single or same colour, therefore _sc. The only new argument is colour, which can be any hexcode or name of colours in the all_grafify palette.

Usage
plot_dotbar_sd_sc(data, xcol, ycol, colour = "ok_orange", dotsize = 1.5, dotthick = 1, bwid = 0.7, ewid = 0.2, b_alpha = 1, d_alpha = 1)
plot_dotbar_sd_sc

TextXAngle = 0,
fontsize = 20,
...
)

Arguments

data          a data table object, e.g. data.frame or tibble.
xcol          name of the column to plot on X axis. This should be a categorical variable.
ycol          name of the column to plot on quantitative Y axis. This should be a quantitative variable.
colour        colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is ok_orange.
dotsize       size of dots relative to binwidth used by geom_dotplot. Default set to 1.5, increase/decrease as needed.
dotthick      thickness of dot border (stroke parameter of geom_dotplot), default set to 1.
bw         width of bars, default set to 0.7
ewid          width of error bars, default set to 0.2.
b_alpha       fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
d_alpha       fractional opacity of dots, default set to 1 (i.e. maximum opacity & zero transparency).
TextXAngle    orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize       parameter of base_size of fonts in theme_classic, default set to size 20.
...           any additional arguments to pass to ggplot2 geom_dotplot.

Details
The default colour is ok_orange. ColPal and ColRev arguments are not available. Colours available can be seen quickly with plot_grafify_palette.

Value
This function returns a ggplot2 object of class "gg" and "ggplot".

Examples
#default "okabe_ito" colour
plot_dotbar_sd_sc(data = data_doubling_time,
                   xcol = Student, ycol = Doubling_time)

# a different colour
plot_dotbar_sd_sc(data = data_doubling_time,
                   xcol = Student, ycol = Doubling_time,
                   colour = "#88ccee")
Description

This function takes a data table, X and Y variables, and plots a graph with a dotplot and boxplot using `geom_boxplot` and `geom_dotplot` geometries. Note that `geom_boxplot` option for outliers is set to `outlier.alpha = 0`.

Usage

```r
plot_dotbox(
  data, xcol, ycol,
  dotsize = 1.5, dotthick = 1, b_alpha = 1, d_alpha = 1,
  ColPal = "all_grafify", ColRev = FALSE, ColSeq = TRUE,
  TextXAngle = 0, fontsize = 20,
  ...
)
```

Arguments

data     a data table object, e.g. data.frame or tibble.
xcol     name of the column to plot on X axis. This should be a categorical variable.
ycol     name of the column to plot on quantitative Y axis. This should be a quantitative variable.
dotsize  size of dots relative to binwidth used by `geom_dotplot`. Default set to 1.5, increase/decrease as needed.
dotthick thickness of dot border (`stroke` parameter of `geom_dotplot`), default set to 1.
b_alpha  fractional opacity of boxes, default set to 1 (i.e. maximum opacity & zero transparency).
d_alpha  fractional opacity of dots, default set to 1 (i.e. maximum opacity & zero transparency).
ColPal   `grafify` colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColRev   whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.

TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

fontsize parameter of base_size of fonts in theme_classic, default set to size 20.

... any additional arguments to pass to ggplot2geom_boxplot or ggplot2geom_dotplot.

Details

The X variable is mapped to the fill aesthetic in both boxplot and dotplot. Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

The size of dots can be adjusted using the parameter, which is dotsize = 1 by default.

This function is related to plot_scatterbar_sd, plot_dotbar_sd and plot_dotviolin.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

plot_dotbox(data = data_1w_death, xcol = Genotype, ycol = Death)

plot_dotbox(data = data_1w_death, xcol = Genotype, ycol = Death, ColPal = "vibrant", b_alpha = 0.5)

plot_dotbox_sc

Plot a dotplot on a boxplot with two variables.

Description

This function is related to plot_dotbox which maps the X variable to different fill colours, but this one maps a single or same colour, therefore _sc. The only new argument is colour, which can be any hexcode or name of colours in the all_grafify palette.
plot_dotbox_sc

Usage

plot_dotbox_sc(
  data,
  xcol,
  ycol,
  colour = "ok_orange",
  dotsize = 1.5,
  dotthick = 1,
  b_alpha = 1,
  d_alpha = 1,
  TextXAngle = 0,
  fontsize = 20,
  ...
)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column to plot on X axis. This should be a categorical variable.
ycol name of the column to plot on quantitative Y axis. This should be a quantitative variable.
colour colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is ok_orange.
dotsize size of dots relative to binwidth used by geom_dotplot. Default set to 1.5, increase/decrease as needed.
dotthick thickness of dot border (stroke parameter of geom_dotplot), default set to 1.
b_alpha fractional opacity of boxes, default set to 1 (i.e. maximum opacity & zero transparency).
d_alpha fractional opacity of dots, default set to 1 (i.e. maximum opacity & zero transparency).
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize parameter of base_size of fonts in theme_classic, default set to size 20.
... any additional arguments to pass to ggplot2geom_boxplot or ggplot2geom_dotplot.

Details

The default colour is ok_orange. ColPal and ColRev arguments are not available. Colours available can be seen quickly with plot_grafify_palette.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".
plot_dotviolin

Examples

# with default colour ("okabe_ito")
plot_dotbox_sc(data = data_doubling_time,
xcol = Student, ycol = Doubling_time)
# set a different colour
plot_dotbox_sc(data = data_doubling_time,
xcol = Student, ycol = Doubling_time,
colour = "pale_blue")

plot_dotviolin  Plot a dotplot on a violin plot with two variables.

Description

This function takes a data table, X and Y variables, and plots a graph with a dotplot, box-whiskers and violinplot using geom_violin, geom_boxplot geom_dotplot geometries.

Usage

plot_dotviolin(
  data,
  xcol,
  ycol,
  dotsize = 1.5,
  dotthick = 1,
  bvthick = 1,
  bwid = 0.2,
  trim = TRUE,
  scale = "width",
  b_alpha = 1,
  v_alpha = 1,
  d_alpha = 1,
  ColPal = "all_grafify",
  ColRev = FALSE,
  ColSeq = TRUE,
  TextXAngle = 0,
  fontsize = 20,
  ...
)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column to plot on X axis. This should be a categorical variable.
**plot_dotviolin**

- **ycol**
  - Name of the column to plot on quantitative Y axis. This should be a quantitative variable.

- **dotsize**
  - Size of dots relative to binwidth used by geom_dotplot. Default set to 1.5, increase/decrease as needed.

- **dotthick**
  - Thickness of dot border (stroke parameter of geom_dotplot), default set to 1.

- **bvthick**
  - Thickness of violin and boxplot lines; default 1.

- **bwid**
  - Width of boxplots; default 0.2

- **trim**
  - Set whether tips of violin plot should be trimmed at high/low data. Default trim = T, can be changed to F.

- **scale**
  - Set to "area" by default, can be changed to "count" or "width".

- **b_alpha**
  - Fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency). For white boxplots inside violins, set b_alpha = 0.

- **v_alpha**
  - Fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency).

- **d_alpha**
  - Fractional opacity of dots, default set to 1 (i.e. maximum opacity & zero transparency).

- **ColPal**
  - Grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".

- **ColRev**
  - Whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).

- **ColSeq**
  - Logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.

- **TextXAngle**
  - Orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

- **fontsize**
  - Parameter of base_size of fonts in theme_classic, default set to size 20.

- **...**
  - Any additional arguments to pass to ggplot2 geom_boxplot, ggplot2 geom_dotplot or ggplot2 geom_violin.

### Details

Note that the **geom_violin** options are set as follows: scale = "width". The trim = T set by default can be changed when calling the function. The boxplot shows IQR, and whiskers show 1.5*IQR; the median is marked with a thicker horizontal line.

The X variable is mapped to the fill aesthetic in both violinplot and dotplot. Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with **plot_grafify_palette**. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using **colorRampPalette**.

The size of dots can be adjusted using the parameter, which is dotsize = 1 by default.

This function is related to **plot_scatterbar_sd**, **plot_dotbar_sd** and **plot_dotviolin**.
Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```r
#plot with trim = FALSE
plot_dotviolin(data = data_t_pdiff,
               xcol = Condition, ycol = Mass,
               dotsize = 2, trim = FALSE)

#white boxplots
plot_dotviolin(data = data_t_pdiff,
               xcol = Condition, ycol = Mass,
               trim = FALSE, b_alpha = 0,
               ColPal = "pale", ColSeq = FALSE)
```

---

**plot_dotviolin_sc**  
Plot a dotplot on a violin plot with two variables.

Description

This function is related to plot_dotviolin, but this one maps a single or same colour, therefore _sc. The only new argument is colour, which can be any hexcode or name of colours in the all_grafify palette. The default colour is ok_orange. ColPal and ColRev arguments are not available. Colours available can be seen quickly with plot_grafify_palette.

Usage

```r
plot_dotviolin_sc(
  data, xcol, ycol, 
  colour = "ok_orange",
  dotsize = 1.5, 
  dotthick = 1, 
  bvthick = 1, 
  bwid = 0.2, 
  b_alpha = 1, 
  d_alpha = 1, 
  v_alpha = 1, 
  trim = TRUE, 
  scale = "width", 
  TextXAngle = 0, 
  fontsize = 20, 
  ...
)
```

Arguments

- **data**: a data table object, e.g. data.frame or tibble.
- **xcol**: name of the column to plot on X axis. This should be a categorical variable.
- **ycol**: name of the column to plot on quantitative Y axis. This should be a quantitative variable.
- **colour**: colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is `ok_orange`.
- **dotsize**: size of dots relative to binwidth used by `geom_dotplot`. Default set to 1.5, increase/decrease as needed.
- **dotthick**: thickness of dot border (stroke parameter of `geom_dotplot`), default set to 1.
- **bvthick**: thickness of violin an boxplot lines; default 1.
- **bwid**: width of boxplots; default 0.2
- **b_alpha**: fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency). For white boxplots inside violins, set `b_alpha = 0`.
- **d_alpha**: fractional opacity of dots, default set to 1 (i.e. maximum opacity & zero transparency).
- **v_alpha**: fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency)
- **trim**: set whether tips of violin plot should be trimmed at high/low data. Default `trim = T`, can be changed to F.
- **scale**: set to "area" by default, can be changed to "count" or "width".
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- **fontsize**: parameter of base_size of fonts in theme_classic, default set to size 20.
- **...**: any additional arguments to pass to ggplot2 `geom_boxplot`, ggplot2 `geom_dotplot` or ggplot2 `geom_violin`.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```r
# plot with trim = FALSE
plot_dotviolin_sc(data = data_1w_death,
xcol = Genotype, ycol = Death,
scale = "width", trim = FALSE)

# white boxplots
plot_dotviolin_sc(data = data_1w_death,
xcol = Genotype, ycol = Death, colour = "light_orange",
scale = "width", trim = FALSE, b_alpha = 0)
```
plot_grafify_palette  See grafify colour palettes

Description

This simple function allows quick visualisation of colours in Grafify palettes and their hex codes. It uses plot_bar_sd and some arguments are similar and can be adjusted.

Usage

plot_grafify_palette(palette = "okabe_ito", bthick = 0, fontsize = 14, ...)

Arguments

- **palette**: name of Grafify palettes: "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", "contrast" or "all_grafify".
- **bthick**: thickness of bars; passed on plot_bar_sd.
- **fontsize**: font size.
- **...**: any additional parameters to pass to plot_bar_sd

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

plot_grafify_palette("pale")
plot_grafify_palette("contrast")

plot_histogram  Plot data distribution as histograms.

Description

This function takes a data table, a quantitative variable (ycol) and a Grouping variable (group), if available, and plots a histogram graph using geom_histogram.
Usage

```r
plot_histogram(
  data, 
  ycol, 
  group, 
  BinSize = 30, 
  linethick = 1, 
  c_alpha = 0.2, 
  ColPal = "all_grafify", 
  ColRev = FALSE, 
  ColSeq = TRUE, 
  TextXAngle = 0, 
  fontsize = 20, 
  Group, 
  alpha, 
  ...
)
```

Arguments

data a data table e.g. data.frame or tibble.
ycol name of the column containing the quantitative variable whose histogram distribution is to be plotted.
group name of the column containing a categorical grouping variable.
BinSize bins to use on X-axis, default set to 30.
linethick thickness of symbol border, default set to 1.
c_alpha fractional opacity of colour filled within histograms, default set to 0.2 (i.e. 20% opacity).
ColPal grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColRev whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize parameter of `base_size` of fonts in `theme_classic`, default set to size 20.
Group deprecated old argument for group; retained for backward compatibility.
alpha deprecated old argument for `c_alpha`; retained for backward compatibility.
... any additional arguments to pass to `ggplot2::geom_histogram`. 
plot_point_sd

Details

Note that the function requires the quantitative Y variable first, and groups them based on an X variable. The group variable is mapped to the fill and colour aesthetics in geom_histogram.

ColPal & ColRev options are applied to both fill and colour scales. Colours available can be seen quickly with `plot_grafify_palette`. Colours can be changed using ColPal, ColRev or ColSeq arguments. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```
# Basic usage
plot_histogram(data = data_t_pratio, 
ycol = Cytokine, group = Genotype, 
BinSize = 10)
# with log transformation
plot_histogram(data = data_t_pratio, 
ycol = log(Cytokine), group = Genotype, 
BinSize = 10)
```

plot_point_sd

Plot a point as mean with SD error bars using two variables.

Description

This function takes a data table, categorical X and numeric Y variables, and plots a point showing the mean with SD error bars. The X variable is mapped to the fill aesthetic of symbols. The related `plot_point_sd_sc` plots bars with a single colour.

Usage

```
plot_point_sd( 
data, 
xcol, 
ycol, 
s_alpha = 1, 
symsize = 3.5, 
symthick = 1, 
ewid = 0.2, 
ColPal = "all_grafify", 
ColSeq = TRUE, 
ColRev = FALSE,
```
Arguments

- **data**: a data table object, e.g. data.frame or tibble.
- **xcol**: name of the column with a categorical X variable.
- **ycol**: name of the column with quantitative Y variable.
- **s_alpha**: fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
- **symsize**: size of point symbols, default set to 3.5.
- **symthick**: thickness of symbol border, default set to 1.
- **ewid**: width of error bars, default set to 0.2.
- **ColPal**: grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
- **ColSeq**: logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
- **ColRev**: whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- **fontsize**: parameter of base_size of fonts in theme_classic, default set to size 20.

Details

The function uses `stat_summary` with geom = "point" with size = 3. Standard deviation (SD) is plotted through `stat_summary` calculated using mean_sdl from the ggplot2 package (get help with ?mean_sdl), and 1x SD is plotted (fun.arg = list(mult = 1)).

Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

You are instead encouraged to show all data using the following functions: plot_scatterbar_sd, plot_scatterbox, plot_dotbox, plot_dotbar_sd, plot_scatterviolin or plot_dotviolin.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".
Examples

# Basic usage
plot_point_sd(data = data_doubling_time,
              xcol = Student, ycol = Doubling_time)

---

**plot_point_sd_sc**  
*Plot a point as mean with SD error bars using two variables.*

Description

This function is related to `plot_point_sd`, but this one maps a single or same colour, therefore `_sc`. The only new argument is `colour`, which can be any hexcode or name of colours in the [all_grafify palette](https://grafify.github.io/all_grafify_pal/). The default colour is `ok_orange`. ColPal and ColRev arguments are not available. Colours available can be seen quickly with `plot_grafify_palette`.

Usage

```r
plot_point_sd_sc(
    data,  
    xcol, 
    ycol, 
    colour = "ok_orange", 
    s_alpha = 1, 
    symsize = 3.5, 
    symthick = 1, 
    ewid = 0.2, 
    TextXAngle = 0, 
    fontsize = 20 
)
```

Arguments

- **data**: a data table object, e.g. data.frame or tibble.
- **xcol**: name of the column with a categorical X variable.
- **ycol**: name of the column with quantitative Y variable.
- **colour**: colour of boxes and dots; a number between 1-64, any hexcode or names from [grafify colour palettes](https://grafify.github.io/all_grafify_pal/). Default is `ok_orange`.
- **s_alpha**: fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
- **symsize**: size of point symbols, default set to 3.5.
- **symthick**: thickness of symbol border, default set to 1
- **ewid**: width of error bars, default set to 0.2.
- **TextXAngle**: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- **fontsize**: parameter of base_size of fonts in theme_classic, default set to size 20.
Details

You are instead encouraged to show all data using the following functions: `plot_scatterbar_sd`, `plot_scatterbox`, `plot_dotbox`, `plot_dotbar_sd`, `plot_scatterviolin` or `plot_dotviolin`.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

# Basic usage
plot_point_sd_sc(data = data_doubling_time, 
xcol = Student, ycol = Doubling_time)
plot_point_sd_sc(data = data_doubling_time, 
xcol = Student, ycol = Doubling_time, 
colour = "ok_grey")

plot_qqline

Plot quantile-quantile (QQ) graphs from data.

Description

This function takes a data table, a quantitative variable (ycol), and a categorical grouping variable (group), if available, and plots a QQ graph using `ggplot2[geom_qq]` and `ggplot2[geom_qq_line]`.

Usage

```r
plot_qqline(
  data, 
  ycol, 
  group, 
  symsize = 3, 
  symthick = 1, 
  s_alpha = 1, 
  ColPal = "all_grafify", 
  ColSeq = TRUE, 
  ColRev = FALSE, 
  TextXAngle = 0, 
  fontsize = 20, 
  Group, 
  ...
)
```
Arguments

- **data**
  a data table e.g. data.frame or tibble.

- **ycol**
  name of the column containing the quantitative variable whose distribution is to be plotted.

- **group**
  name of the column containing a categorical grouping variable.

- **symsize**
  size of symbols, default set to 3.

- **symthick**
  thickness of symbol border, default set to 1.

- **s_alpha**
  fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).

- **ColPal**
  grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".

- **ColSeq**
  logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.

- **ColRev**
  whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).

- **TextXAngle**
  orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

- **fontsize**
  parameter of base_size of fonts in theme_classic, default set to size 20.

- **Group**
  deprecated old argument for group; retained for backward compatibility.

- **...**
  any additional arguments to pass to ggplot2[geom_qq] or ggplot2[geom_qq_line].

Details

Note that the function requires the quantitative Y variable first, and can be passed on a grouping variable as group if required. The graph plots sample quantiles on Y axis & theoretical quantiles on X axis. The X variable is mapped to the fill aesthetic in stat_qq and colour aesthetic for the stat_qq_line.

Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. When only one level is present within group, symbols will receive "ok_orange" colour. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```r
plot_qqline(data = data_cholesterol,
            ycol = Cholesterol, group = Treatment)
```

#with faceting
plot_qqmodel(data = data_cholesterol, ycol = Cholesterol, group = Treatment, fontsize = 10)+facet_wrap("Treatment")

plot_qqmodel  
Plot quantile-quantile (QQ) graphs from residuals of linear models.

Description

This function takes a linear model (simple or mixed effects) and plots a QQ graph after running rstudent from rstudent to generate a table of studentized model residuals on an ordinary (simple_model), mixed model (mixed_model or mixed_model_slopes). The graph plots studentized residuals from the model (sample) on Y axis & Theoretical quantiles on X axis.

Usage

plot_qqmodel(Model, symsize = 2.5, symthick = 1, s_alpha = 1, fontsize = 20)

Arguments

- **Model**  name of a saved model generated by simple_model or mixed_model.
- **symsize** size of symbols, default set to 3.
- **symthick** thickness of symbol border, default set to 1.
- **s_alpha** fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
- **fontsize** parameter of base_size of fonts in theme_classic, default set to size 20.

Details

The function uses ggplot2[geom_qq] and ggplot2[geom_qq_line] geometries. Symbols receive "ok_orange" colour by default.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

# Basic usage
m1 <- simple_model(data = data_2w_Festing, Y_value = "GST", Fixed_Factor = c("Treatment", "Strain"))
plot_qqmodel(m1)
**plot_scatterbar_sd**

Plot scatter dots on a bar graph with SD error bars with two variables.

**Description**

This function takes a data table, categorical X and numeric Y variables, and plots a graph with a jitterplot or scatterplot and bars showing means with SD error bars. It uses `stat_summary` with `geom = "bar"`, and `geom_point` with `position = position_jitter(width = 0.05).

**Usage**

```r
plot_scatterbar_sd(
  data, xcol, ycol, symsize = 2.5, symthick = 1, bwid = 0.7, ewid = 0.3, jitter = 0, b_alpha = 1, s_alpha = 1, ColPal = "all_grafify", ColSeq = TRUE, ColRev = FALSE, TextXAngle = 0, fontsize = 20
)
```

**Arguments**

- **data** a data table object, e.g. data.frame or tibble.
- **xcol** name of the column to plot on X axis. This should be a categorical variable.
- **ycol** name of the column to plot on quantitative Y axis. This should be a quantitative variable.
- **symsize** size of point symbols, default set to 2.
- **symthick** thickness of symbol border, default set to 1.
- **bwid** width of bars, default set to 0.7.
- **ewid** width of error bars, default set to 0.3.
- **jitter** extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
- **b_alpha** fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
- **s_alpha** fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
plot_scatterbar_sd

ColPal  grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted", "dark", "light".

ColSeq  logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.

ColRev  whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).

TextXAngle  orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

fontsize  parameter of base_size of fonts in theme_classic, default set to size 20.

Details

Standard deviation (SD) is plotted through stat_summary calculated using mean_sdl from the ggplot2 package (get help with ?mean_sdl), and 1x SD is plotted (fun.arg = list(mult = 1). The X variable is mapped to the fill aesthetic in the bar geometry and colour aesthetic in geom_point.

Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

Three types of plots are available for scatter/jitter symbols and either bars+SD, boxplot or violin plots: plot_scatterbar_sd, plot_scatterbox and plot_scatterviolin. These are related to the three "dot" versions that use a different geometry for symbols: plot_dotbox, plot_dotbar_sd and plot_dotviolin.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

# with jitter
plot_scatterbar_sd(data = data_cholesterol,
xcol = Treatment, ycol = Cholesterol,
jitter = 0.1)

# white bars
plot_scatterbar_sd(data = data_cholesterol,
xcol = Treatment, ycol = Cholesterol,
b_alpha = 0)
plot_scatterbar_sd_sc  Plot scatter dots on a bar graph with SD error bars with two variables.

Description

This function is related to plot_scatterbar_sd, but this one maps a single or same colour, therefore _sc. The only new argument is colour, which can be any hexcode or name of colours in the all_grafify palette. The default colour is ok_orange. ColPal and ColRev arguments are not available. Colours available can be seen quickly with plot_grafify_palette.

Usage

plot_scatterbar_sd_sc(
  data,
  xcol,
  ycol,
  colour = "ok_orange",
  symsize = 2.5,
  symthick = 1,
  bwidth = 0.7,
  ewid = 0.3,
  jitter = 0,
  b_alpha = 1,
  s_alpha = 1,
  TextXAngle = 0,
  fontsize = 20
)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column to plot on X axis. This should be a categorical variable.
ycol name of the column to plot on quantitative Y axis. This should be a quantitative variable.
colour colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is ok_orange.
symsize size of point symbols, default set to 2.
symthick thickness of symbol border, default set to 1.
bwidth width of bars, default set to 0.7
ewid width of error bars, default set to 0.3.
jitter extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
b_alpha fractional opacity of bars, default set to 1 (i.e. maximum opacity & zero transparency).
plot_scatterbox

s_alpha fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).

TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

fontsize parameter of base_size of fonts in theme_classic, default set to size 20.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

plot_scatterbar_sd_sc(data = data_doubling_time, xcol = Student, ycol = Doubling_time)
plot_scatterbar_sd_sc(data = data_doubling_time, xcol = Student, ycol = Doubling_time, colour = "ok_grey")

plot_scatterbox Plot a scatter plot on a boxplot with two variables.

Description

This function takes a data table, X and Y variables, and plots a graph with a scatter plot and box and whiskers using geom_boxplot and geom_point geometries. The boxplot shows IQR and whiskers depict 1.5*IQR. Note that geom_boxplot option for outliers is set to outlier.alpha = 0. The X variable is mapped to the fill aesthetic in both boxplot and symbols, and its colour can be changed using ColPal option. Colours can be changed using ColPal, ColRev or ColSeq arguments. Colours available can be seen quickly with plot_grafify_palette. ColPal can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". ColRev (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. ColSeq (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

Usage

plot_scatterbox(
data,  
xcol,  
ycol,  
symsize = 2.5,  
symthick = 1,  
jitter = 0,  
b_alpha = 1,  
s_alpha = 1,  
ColPal = "all_grafify",  
)
plot_scatterbox

ColSeq = TRUE,
ColRev = FALSE,
TextXAngle = 0,
fontsize = 20,
...)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column to plot on X axis. This should be a categorical variable.
ycol name of the column to plot on quantitative Y axis. This should be a quantitative variable.
symsize size of symbols used by geom_point. Default set to 2.5, increase/decrease as needed.
symthick thickness of symbol border (stroke parameter of geom_point), default set to 1.
jitter extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
b_alpha fractional opacity of boxplot, default set to 1 (i.e. maximum opacity & zero transparency).
s_alpha fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
ColPal grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColRev whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize parameter of base_size of fonts in theme_classic, default set to size 20.
... any additional arguments to pass to ggplot2geom_boxplot.

Details

The size of symbols can be adjusted using symsize set to 1 by default. Transparency of boxplot and symbols can be set independently with b_alpha and s_alpha, respectively.

Three types of plots are available for scatter/jitter symbols and either bars+SD, boxplot or violin plots: plot_scatterbar_sd, plot_scatterbox and plot_scatterviolin. These are related to the three "dot" versions that use a different geometry for symbols: plot_scatterbox, plot_dotbar_sd and plot_dotviolin.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".
Examples

```r
plot_scatterbox(data = data_cholesterol, 
    xcol = Treatment, ycol = Cholesterol)

#with jitter
plot_scatterbox(data = data_cholesterol, 
    xcol = Treatment, ycol = Cholesterol, jitter = 0.1)
```

Description

This function is related to `plot_scatterbox_sd`, but this one maps a single or same colour, therefore _sc_. The only new argument is `colour`, which can be any hexcode or name of colours in the `all_grafify` palette. The default colour is `ok_orange`. ColPal and ColRev arguments are not available. Colours available can be seen quickly with `plot_grafify_palette`.

Usage

```r
plot_scatterbox_sc(
    data, 
    xcol, 
    ycol, 
    colour = "ok_orange", 
    symsize = 2.5, 
    symthick = 1, 
    jitter = 0, 
    b_alpha = 1, 
    s_alpha = 1, 
    TextXAngle = 0, 
    fontsize = 20, 
    ...
)
```

Arguments

- **data**: a data table object, e.g. data.frame or tibble.
- **xcol**: name of the column to plot on X axis. This should be a categorical variable.
- **ycol**: name of the column to plot on quantitative Y axis. This should be a quantitative variable.
- **colour**: colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is `ok_orange`.
- **symsize**: size of symbols used by `geom_point`. Default set to 2.5, increase/decrease as needed.
plot_scatterviolin

Plot a scatter plot on a violin plot with two variables.

Description
This function takes a data table, X and Y variables, and plots a graph with a scatter plot and violin-plot using ggplot.

Usage

```r
plot_scatterviolin(
  data,
  xcol,
  ycol,
  symsize = 2.5,
  symthick = 1,
  bwid = 0.1,
  bvthick = 1,
  b_alpha = 1,
  s_alpha = 1,
...```

Arguments

- `symthick`: thickness of symbol border (stroke parameter of geom_point), default set to 1.
- `jitter`: extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
- `b_alpha`: fractional opacity of boxplot, default set to 1 (i.e. maximum opacity & zero transparency).
- `s_alpha`: fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
- `TextXAngle`: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
- `...`: any additional arguments to pass to ggplot2::geom_boxplot.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

```r
# with jitter
plot_scatterbox_sc(data = data_cholesterol,
  xcol = Treatment, ycol = Cholesterol, jitter = 0.1)
# with "ok_grey" colour
plot_scatterbox_sc(data = data_cholesterol,
  xcol = Treatment, ycol = Cholesterol,
  colour = "ok_grey", jitter = 0.1)
```
v_alpha = 1,
ColPal = "all_grafify",
ColSeq = TRUE,
ColRev = FALSE,
jitter = 0,
trim = TRUE,
scale = "width",
TextXAngle = 0,
fontsize = 20,
...)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column to plot on X axis. This should be a categorical variable.
ycol name of the column to plot on quantitative Y axis. This should be a quantitative variable.
symsize size of dots relative to binwidth used by geom_point. Default set to 2.5, increase/decrease as needed.
sythick thickness of dot border (stroke parameter of geom_point), default set to 1.
bwthick width of boxplots; default 0.2
b_alpha fractional opacity of boxplots, default set to 1 (i.e. maximum opacity & zero transparency). For white boxplots inside violins, set b_alpha = 0.
s_alpha fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency). Set s_alpha = 0 to not show scatter plot.
v_alpha fractional opacity of violins, default set to 1 (i.e. maximum opacity & zero transparency).
ColPal grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
ColSeq logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using scale_fill_grafify2.
ColRev whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
jitter extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.
trim set whether tips of violin plot should be trimmed at high/low data. Default trim = T, can be changed to F.
scale set to "area" by default, can be changed to "count" or "width".
TextXAngle orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize parameter of base_size of fonts in theme_classic, default set to size 20.
... any additional arguments to pass to ggplot2geom_boxplot, ggplot2geom_point or ggplot2geom_violin.
Details

The function uses `geom_violin`, `geom_boxplot` and `geom_point` geometries. Note that the `geom_violin` options are set as follows: `scale = "width"`. The `trim = T` set by default can be changed when calling the function. The boxplot shows IQR and the median is marked with a thicker horizontal line, and whisker depicts 1.5*IQR. The X variable is mapped to the `fill` aesthetic in both violin and symbols, and its colour can be changed using `ColPal` option. Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. Colours available can be seen quickly with `plot_grafify_palette`. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq` (logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using `colorRampPalette`.

The size of symbols can be adjusted using `symsize` set to 1 by default. Transparency of violins and symbols can be set independently with `v_alpha` and `s_alpha`, respectively.

Three types of plots are available for scatter/jitter symbols and either bars+SD, boxplot or violin plots: `plot_scatterbar_sd`, `plot_scatterbox` and `plot_scatterviolin`. These are related to the three "dot" versions that use a different geometry for symbols: `plot_dotbox`, `plot_dotbar_sd` and `plot_dotviolin`.

Value

This function returns a `ggplot2` object of class "gg" and "ggplot".

Examples

```r
#plot without jitter
plot_scatterviolin(data = data_t_pdiff,
                   xcol = Condition, ycol = Mass,
                   symsize = 2, trim = FALSE)

#with jitter
plot_scatterviolin(data = data_t_pdiff,
                   xcol = Condition, ycol = Mass,
                   symsize = 2, trim = FALSE, jitter = 0.1)

#white boxplot and no symbols
plot_scatterviolin(data = data_t_pdiff,
                   xcol = Condition, ycol = Mass,
                   b_alpha = 0, s_alpha = 0,
                   symsize = 2, trim = FALSE, jitter = 0.1)
```
plot_scatterviolin_sc  Plot a scatter plot on a violin plot with two variables.

Description
This function is related to plot_scatterviolin, but this one maps a single or same colour, therefore _sc. The only new argument is colour, which can be any hexcode or name of colours in the all_grafify palette. The default colour is ok_orange. ColPal and ColRev arguments are not available. Colours available can be seen quickly with plot_grafify_palette.

Usage
plot_scatterviolin_sc(
  data,
  xcol,
  ycol,
  colour = "ok_orange",
  symsize = 2.5,
  symthick = 1,
  bwid = 0.2,
  bvthick = 1,
  b_alpha = 1,
  v_alpha = 1,
  s_alpha = 1,
  jitter = 0,
  trim = TRUE,
  scale = "width",
  TextXAngle = 0,
  fontsize = 20,
  ...
)

Arguments
  data  a data table object, e.g. data.frame or tibble.
  xcol  name of the column to plot on X axis. This should be a categorical variable.
  ycol  name of the column to plot on quantitative Y axis. This should be a quantitative variable.
  colour  colour of boxes and dots; a number between 1-64, any hexcode or names from grafify colour palettes. Default is ok_orange.
  symsize  size of dots relative to binwidth used by geom_point. Default set to 2.5, increase/decrease as needed.
  symthick  thickness of dot border (stroke parameter of geom_point), default set to 1.
  bwid  width of boxplots; default 0.2
  bvthick  thickness of both violin and box plot lines; default 1.
**plot_xy_CatGroup**

Plot points on a quantitative X - Y plot & a categorical grouping variable.

### Description

This function takes a data table, quantitative X and Y variables along with a categorical grouping variable, and a and plots a graph with using `geom_point`. The categorical CatGroup variable is mapped to the `fill` aesthetic of symbols.

### Examples

```r
plot_scatterviolin_sc(data = data_doubling_time, 
  xcol = Student, ycol = Doubling_time, 
  colour = "ok_grey", 
  symsize = 2, trim = FALSE, scale = "width")
```

```
#white boxplots and no symbols
plot_scatterviolin_sc(data = data_1w_death, 
  xcol = Genotype, ycol = Death, 
  colour = "pale_blue", b_alpha = 0, s_alpha = 0, 
  symsize = 2, trim = FALSE, scale = "width")
```

### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>b_alpha</td>
<td>fractional opacity of boxplots, default set to 1 (i.e. maximum opacity &amp; zero transparency). For white boxplots inside violins, set b_alpha = 0.</td>
</tr>
<tr>
<td>v_alpha</td>
<td>fractional opacity of violins, default set to 1 (i.e. maximum opacity &amp; zero transparency). Set s_alpha = 0 to not show scatter plot.</td>
</tr>
<tr>
<td>s_alpha</td>
<td>fractional opacity of symbols, default set to 1 (i.e. maximum opacity &amp; zero transparency).</td>
</tr>
<tr>
<td>jitter</td>
<td>extent of jitter (scatter) of symbols, default is 0 (i.e. aligned symbols). To reduce symbol overlap, try 0.1-0.3 or higher.</td>
</tr>
<tr>
<td>trim</td>
<td>set whether tips of violin plot should be trimmed at high/low data. Default trim = TRUE, can be changed to FALSE.</td>
</tr>
<tr>
<td>scale</td>
<td>set to &quot;area&quot; by default, can be changed to &quot;count&quot; or &quot;width&quot;.</td>
</tr>
<tr>
<td>TextXAngle</td>
<td>orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.</td>
</tr>
<tr>
<td>fontsize</td>
<td>parameter of base_size of fonts in theme_classic, default set to size 20.</td>
</tr>
<tr>
<td>...</td>
<td>any additional arguments to pass to ggplot2geom_boxplot, ggplot2geom_point or ggplot2geom_violin.</td>
</tr>
</tbody>
</table>

### Value

This function returns a ggplot2 object of class "gg" and "ggplot".
**Usage**

```r
plot_xy_CatGroup(
  data,
  xcol,
  ycol,
  CatGroup,
  symsize = 2.5,
  symthick = 1,
  s_alpha = 1,
  ColPal = "all_grafify",
  ColSeq = TRUE,
  ColRev = FALSE,
  TextXAngle = 0,
  fontsize = 20
)
```

**Arguments**

- `data` : a data table object, e.g. data.frame or tibble.
- `xcol` : name of the column with quantitative X variable.
- `ycol` : name of the column with quantitative Y variable.
- `CatGroup` : a categorical variable as grouping factor for colour of data points, should be a categorical variable for default colours to work. Will be converted to factor if your column is numeric.
- `symsize` : size of symbols used by `geom_point`. Default set to 2.5, increase/decrease as needed.
- `symthick` : thickness of symbol border, default set to 1.
- `s_alpha` : fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
- `ColPal` : grafify colour palette to apply, default "all_grafify"; alternatives: "okabe_ito", "bright", "pale", "vibrant", "contrast", "muted" "dark", "light".
- `ColSeq` : logical TRUE or FALSE. Default TRUE for sequential colours from chosen palette. Set to FALSE for distant colours, which will be applied using `scale_fill_grafify2`.
- `ColRev` : whether to reverse order of colour choice, default F (FALSE); can be set to T (TRUE).
- `TextXAngle` : orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.

**Details**

Colours can be changed using `ColPal`, `ColRev` or `ColSeq` arguments. Colours available can be seen quickly with `plot_grafify_palette`. `ColPal` can be one of the following: "okabe_ito", "dark", "light", "bright", "pale", "vibrant", "muted" or "contrast". `ColRev` (logical TRUE/FALSE) decides whether colours are chosen from first-to-last or last-to-first from within the chosen palette. `ColSeq`
(logical TRUE/FALSE) decides whether colours are picked by respecting the order in the palette or the most distant ones using colorRampPalette.

This plot is related to plot_xy_NumGroup which requires a numeric grouping factor. When summary statistics (mean/median) are required, use plot_3d_scatterbar, plot_3d_scatterbox or plot_4d_scatterbox.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

#The grouping factor cyl is automatically converted to categorical variable
plot_xy_CatGroup(data = mtcars,
    xcol = mpg, ycol = disp, CatGroup = cyl,
    ColPal = "vibrant", ColSeq = FALSE)

plot_xy_NumGroup(data = mtcars, xcol = mpg, ycol = disp, NumGroup = cyl)

Description

This function takes a data table, quantitative X and Y variables, and a numeric grouping variable, and plots a graph with using geom_point. The numerical NumGroup variable is mapped to the fill aesthetic of symbols, which receives the scale_fill_grafify_c default palette.

Usage

plot_xy_NumGroup(
    data, xcol, ycol, NumGroup,
    symsize = 2.5, symthick = 1,
    s_alpha = 1, TextXAngle = 0,
    fontsize = 20
)

Arguments

data a data table object, e.g. data.frame or tibble.
xcol name of the column with quantitative X variable.
ycol name of the column with quantitative Y variable.
NumGroup a numeric factor for fill aesthetic of data points.
posthoc_Levelwise

symsize: size of symbols used by geom_point. Default set to 2.5, increase/decrease as needed.
symthick: thickness of symbol border, default set to 1.
s_alpha: fractional opacity of symbols, default set to 1 (i.e. maximum opacity & zero transparency).
TextXAngle: orientation of text on X-axis; default 0 degrees. Change to 45 or 90 to remove overlapping text.
fontsize: parameter of base_size of fonts in theme_classic, default set to size 20.

Details

This plot is related to plot_xy_CatGroup which requires a categorical grouping factor. When summary statistics (mean/median) are required, use plot_3d_scatterbar, plot_3d_scatterbox or plot_4d_scatterbox.

Value

This function returns a ggplot2 object of class "gg" and "ggplot".

Examples

#The grouping factor gear is numeric
plot_xy_NumGroup(data = mtcars,
  xcol = mpg, ycol = disp, NumGroup = cyl,
  s_alpha = 0.8)

posthoc_Levelwise: Level-wise post-hoc comparisons from a linear or linear mixed effects model.

Description

This function is a wrapper based on emmeans, and needs an ordinary linear model produced by simple_model or a mixed effects model produced by mixed_model or mixed_model_slopes (or generated directly with lm, lme4 or lmerTest calls). It also needs to know the fixed factor(s), which should match those in the model and data table.

Usage

posthoc_Levelwise(Model, Fixed_Factor, P_Adj = "fdr", Factor, ...)
Arguments

Model: a model object fit using `simple_model` or `mixed_model` or related.

Fixed_Factor: one or more categorical variables, provided as a vector (see Examples), whose levels you wish to compare pairwise. Names of Fixed_Factor should match Fixed_Factor used to fit the model. When more than one factor is provided e.g. Fixed_factor = c("A","B"), this function passes this on as specs = A|B (note the vertical | between the two Fixed_Factor) to emmeans to produce comparisons between each level A with each other listed separately at each level of B.

P_Adj: method for correcting P values for multiple comparisons. Default is set to false discovery rate ("fdr"), can be changed to "none", "tukey", "bonferroni", "sidak". See Interaction analysis in emmeans in the manual for emmeans.

Factor: old argument name for Fixed_Factor; retained for backward compatibility.

...: additional arguments for emmeans such as lmer.df or others. See help for sophisticated models in emmeans.

Details

The function will generate level-wise comparisons (as described in Comparisons and contrasts in emmeans), i.e. comparison between of every level of one factor separately at each level of the other factor. By default, P values are corrected by the FDR method (which can be changed). If the model was fit by transforming the quantitative response variable using "log", "logit", "sqrt" etc., results will still be on the original scale, i.e. type = "response" is the default; data will be back-transformed (check results to confirm this), and for log or logit see Transformations and link functions in emmeans, ratios will be compared. The first part of the emmeans results has the estimated marginal means, SE and CI ($emmeans), which are generated from the fitted model, and not the original data table. The second part has the results of the comparisons ($contrasts).

Value

returns an "emm_list" object containing contrasts and emmeans through emmeans.

Examples

#make a linear model first
CholMod <- mixed_model(data = data_cholesterol, 
  Y_value = "Cholesterol", 
  Fixed_Factor = c("Hospital", "Treatment"), 
  Random_Factor = "Subject")

#note quotes used only for fixed Fixed_Factor
#to get comparisons between different hospitals separately for each level of Treatment
posthoc_Levelwise(Model = CholMod, 
  Fixed_Factor = c("Hospital", "Treatment"))

#get comparisons between treatments separately at each hospital
posthoc_Levelwise(Model = CholMod, 
  Fixed_Factor = c("Treatment", "Hospital"))
posthoc_Pairwise  
Pairwise post-hoc comparisons from a linear or linear mixed effects model.

Description
This function is a wrapper based on emmeans, and needs a ordinary linear model produced by simple_model or a mixed effects model produced by mixed_model or mixed_model_slopes (or generated directly with lm, lme4 or lmerTest calls). It also needs to know the fixed factor(s), which should match those in the model and data table.

Usage
posthoc_Pairwise(Model, Fixed_Factor, P_Adj = "fdr", Factor, ...)

Arguments
Model  
a model object fit using simple_model or mixed_model or related.

Fixed_Factor  
one or more categorical variables, provided as a vector (see Examples), whose levels you wish to compare pairwise. Names of Fixed_Factor should match Fixed_Factor used to fit the model. When more than one factor is provided e.g. Fixed_factor = c("A", "B"), this function passes this on as specs = A:B (note the colon between the two Fixed_Factor) to emmeans to produce pairwise comparisons.

P_Adj  
method for correcting P values for multiple comparisons. Default is set to false discovery rate ("fdr"), can be changed to "none", "tukey", "bonferroni", "sidak". See Interaction analysis in emmeans in the manual for emmeans.

Factor  
old argument name for Fixed_Factor; retained for backward compatibility.

...  
additional arguments for emmeans such as lmer.df or others. See help for sophisticated models in emmeans.

Details
The function will generate pairwise comparisons of every level of every factor (as described in Comparisons and contrasts in emmeans). Too many comparisons will be generated and only use this when necessary. By default, P values are corrected by the FDR method (which can be changed). If the model was fit by transforming the quantitative response variable using "log", "logit", "sqrt" etc., results will still be on the original scale, i.e. type = "response" is the default; data will be back-transformed (check results to confirm this), and for log or logit see Transformations and link functions in emmeans, ratios will be compared. The first part of the emmeans results has the estimated marginal means, SE and CI ($emmeans), which are generated from the fitted model, and not the original data table. The second part has the results of the comparisons ($contrasts).

Value
returns an "emm_list" object containing contrasts and emmeans through emmeans.
Examples

# make linear models first
DoublMod <- simple_model(data = data_doubling_time,
  Y_value = "Doubling_time", Fixed_Factor = "Student")
CholMod <- mixed_model(data = data_cholesterol,
  Y_value = "Cholesterol",
  Fixed_Factor = c("Hospital", "Treatment"),
  Random_Factor = "Subject")

posthoc_Pairwise(Model = DoublMod,
  Fixed_Factor = "Student")

# basic use with two Fixed_Factor provided as a vector
posthoc_Pairwise(Model = CholMod,
  Fixed_Factor = c("Treatment", "Hospital"))

# same call with "tukey" adjustment
posthoc_Pairwise(Model = CholMod,
  Fixed_Factor = c("Treatment", "Hospital"),
  P_adj = "tukey")

Description

This function is a wrapper based on emmeans, and needs a ordinary linear model produced by
simple_model or a mixed effects model produced by mixed_model or mixed_model_slopes (or
generated directly with lm, lme4 or lmerTest calls). At least one of the factors should be a numeric
 covariate whose slopes you wish to find. It also needs to know the fixed factor(s), which should
 match those in the model and data table.

Usage

posthoc_Trends_Levelwise(
  Model,
  Fixed_Factor,
  Trend_Factor,
  P_Adj = "sidak",
  ...
)

Arguments

Model a model object fit using simple_model or mixed_model (or lm or lmer).
Fixed_Factor

one or more categorical variables, provided as a vector (see Examples), whose levels you wish to compare pairwise. Names of Fixed_Factor should match Fixed_Factor used to fit the model. When more than one factor is provided e.g. Fixed_factor = c("A", "B"), this function passes this on as specs = A:B (note the colon between the two Fixed_Factor) to `emmeans` to produce pairwise comparisons.

Trend_Factor

a quantitative variable that interacts with a factor and whose slope (trend) is to be compared

P_Adj

method for correcting P values for multiple comparisons. Default is "sidak", can be changed to "bonferroni". See Interaction analysis in emmeans in the manual for emmeans.

... additional arguments for `emmeans` such as `lmer.df` or others. See help for sophisticated models in `emmeans`.

Details

Checkout the Interactions with covariates section in the `emmeans` vignette for more details. One of the independent variables should be a quantitative (e.g. time points) variable whose slope (trend) you want to find at levels of the other factor.

Value

returns an "emm_list" object containing slopes and their contrasts calculated through `emtrends`.

Examples

```r
# create an lm model
# Time2 is numeric (time points)
m1 <- simple_model(data = data_2w_Tdeath, Y_value = "PI", Fixed_Factor = c("Genotype", "Time2"))
posthoc_Trends_Levelwise(Model = m1, Fixed_Factor = "Genotype", Trend_Factor = "Time2")
```

Description

This function is a wrapper based on `emmeans`, and needs a ordinary linear model produced by `simple_model` or a mixed effects model produced by `mixed_model` or `mixed_model_slopes` (or generated directly with `lm`, `lme4` or `lmerTest` calls). At least one of the factors should be a numeric covariate whose slopes you wish to find. It also needs to know the fixed factor(s), which should match those in the model and data table.
Usage

posthoc_Trends_Pairwise(
  Model,
  Fixed_Factor,
  Trend_Factor,
  P_Adj = "sidak",
  ...
)

Arguments

Model a model object fit using simple_model or mixed_model (or lm or lmer).
Fixed_Factor one or more categorical variables, provided as a vector (see Examples), whose levels you wish to compare pairwise. Names of Fixed_Factor should match Fixed_Factor used to fit the model. When more than one factor is provided e.g. Fixed_factor = c("A","B"), this function passes this on as specs = A:B (note the colon between the two Fixed_Factor) to emmeans to produce pairwise comparisons.
Trend_Factor a quantitative variable that interacts with a factor and whose slope (trend) is to be compared
P_Adj method for correcting P values for multiple comparisons. Default is "sidak", can be changed to "bonferroni". See Interaction analysis in emmeans in the manual for emmeans.
... additional arguments for emmeans such as lmer.df or others. See help for sophisticated models in emmeans.

Details

Checkout the Interactions with covariates section in the emmeans vignette for more details. One of the independent variables should be a quantitative (e.g. time points) variable whose slope (trend) you want to find at levels of the other factor.

Value

returns an "emm_list" object containing slopes and their contrasts calculated through emtrends.

Examples

#create an lm model
#Time2 is numeric (time points)
m1 <- simple_model(data = data_2w_Tdeath,
  Y_value = "PI", Fixed_Factor = c("Genotype", "Time2"))
posthoc_Trends_Pairwise(Model = m1,
  Fixed_Factor = "Genotype",
  Trend_Factor = "Time2")
posthoc_Trends_vsRef Use emtrends to get level-wise comparison of slopes from a linear model.

Description

This function is a wrapper based on emmeans, and needs a ordinary linear model produced by simple_model or a mixed effects model produced by mixed_model or mixed_model_slopes (or generated directly with lm, lme4 or lmerTest calls). At least one of the factors should be a numeric covariate whose slopes you wish to find. It also needs to know the fixed factor(s), which should match those in the model and data table.

Usage

posthoc_Trends_vsRef(
  Model,
  Fixed_Factor,
  Trend_Factor,
  Ref_Level = 1,
  P_Adj = "sidak",
  ...
)

Arguments

Model a model object fit using simple_model or mixed_model (or lm or lmer).

Fixed_Factor one or more categorical variables, provided as a vector (see Examples), whose levels you wish to compare pairwise. Names of Fixed_Factor should match Fixed_Factor used to fit the model. When more than one factor is provided e.g. Fixed_factor = c("A","B"), this function passes this on as specs = A:B (note the colon between the two Fixed_Factor) to emmeans to produce pairwise comparisons.

Trend_Factor a quantitative variable that interacts with a factor and whose slope (trend) is to be compared.

Ref_Level the level within that factor to be considered the reference or control to compare other levels to (to be provided as a number - by default R orders levels alphabetically); default Ref_Level = 1.

P_Adj method for correcting P values for multiple comparisons. Default is "sidak", can be changed to "bonferroni". See Interaction analysis in emmeans in the manual for emmeans.

... additional arguments for emmeans such as lmer.df or others. See help for sophisticated models in emmeans.
Details

Checkout the Interactions with covariates section in the emmeans vignette for more details. One of the independent variables should be a quantitative (e.g. time points) variable whose slope (trend) you want to find at levels of the other factor.

Value

returns an "emm_list" object containing slopes and their contrasts calculated through emtrends.

Examples

# create an lm model
# Time2 is numeric (time points)
m1 <- simple_model(data = data_2w_Tdeath,
  Y_value = "PI", Fixed_Factor = c("Genotype", "Time2"))
posthoc_Trends_vsRef(Model = m1,
  Fixed_Factor = "Genotype",
  Trend_Factor = "Time2",
  Ref_Level = 2)

Arguments

Model a model object fit using simple_model or mixed_model or related.
Fixed_Factor Fixed_Factor one or more categorical variables, provided as a vector (see Examples), whose levels you wish to compare pairwise. Names of Fixed_Factor should match Fixed_Factor used to fit the model. When more than one factor is provided e.g. Fixed_Factor = c("A","B"), this function passes this on as specs = A|B (note the vertical | between the two Fixed_Factor) to emmeans. The specification internally is set to specs = trt.vs.ctrl,Ref_Level = 1 to compare each group in A to the reference first group in A, separately at each level of B.
Ref_Level  

the level within that factor to be considered the reference or control to compare other levels to (to be provided as a number - by default R orders levels alphabetically); default Ref_Level = 1.

P_Adj  

method for correcting P values for multiple comparisons. Default is set to false discovery rate ("fdr"), can be changed to "none", "tukey", "bonferroni", "sidak". See Interaction analysis in emmeans in the manual for emmeans.

Factor  

old argument name for Fixed_Factor; retained for backward compatibility.

...  

additional arguments for emmeans such as lmer.df or others. See help for sophisticated models in emmeans.

Details  

The function will generate treatment vs control type of comparisons (as described in Comparisons and contrasts in emmeans), i.e. comparison of each level of a factor to a reference level, which is set by default to the first level in the factor (Ref_Level = 1). By default, P values are corrected by the FDR method (which can be changed). If the model was fit by transforming the quantitative response variable using "log", "logit", "sqrt" etc., results will still be on the original scale, i.e. type = "response" is the default; data will be back-transformed (check results to confirm this), and for log or logit see Transformations and link functions in emmeans, ratios will be compared. The first part of the emmeans results has the estimated marginal means, SE and CI (\$emmeans), which are generated from the fitted model, and not the original data table. The second part has the results of the comparisons (\$contrasts).

Value  

returns an "emm_list" object containing contrasts and emmeans through emmeans.

Examples  

```r
#make linear models first
DoublMod <- simple_model(data = data_doubling_time,
    Y_value = "Doubling_time",
    Fixed_Factor = "Student")

CholMod <- mixed_model(data = data_cholesterol,
    Y_value = "Cholesterol",
    Fixed_Factor = c("Hospital", "Treatment"),
    Random_Factor = "Subject")

#to compare all students with student #9
posthoc_vsRef(Model = DoublMod,
    Fixed_Factor = "Student", Ref_Level = 9)

#for comparison between hospital_a to every other hospital, separately at levels of Treatment
posthoc_vsRef(Model = CholMod,
    Fixed_Factor = c("Hospital", "Treatment"), Ref_Level = 1)
```
Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that these only work for categorical variables. Use the brewer or viridis packages for numeric gradient scales.

Usage

scale_color_grafify(palette = "all_grafify", reverse = FALSE, ...)

Arguments

- **palette**: Name of the colour scheme. Default set to palette = "all_grafify". Provide names as above in quotes.
- **reverse**: Whether the colour order should be reversed.
- **...**: Additional parameters for scale_fill or scale_colour.

Details

The default for scale_fill_grafify(), scale_colour_grafify() or scale_color_grafify() is a list of 55 colours as part of palette = "all_grafify". Obviously, it is not recommended to use so many colours, but implementing this was easiest to prevent errors when using a lot of categorical variables.

Colours available can be seen quickly with plot_grafify_palette. There are eight palettes with 5-10 colours each, which are recommended. These can be called by naming the colour scheme using palette = argument. Additional options include "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", and "contrast". These are taken from Paul Taul, Mike Mol and Okabe Ito. scale_fill_grafify2 and scale_colour_grafify2 are identical except that when the number of categorical variables is fewer than the total number of colour shades in the palette (e.g. if you have 3 groups and the "okabe_ito" palette has 7 colours), these functions will pick the most 'distant' colours from the scheme than going sequentially. If you want colours assigned sequentially use scale_fill_grafify or scale_colour_grafify.

Value

ggplot scale_fill function for discrete colours.

Examples

```r
#add a colour scheme to a ggplot object
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration))+
  geom_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9, position = position_jitter(0.15)) +
  scale_color_grafify(palette = "bright") + facet_wrap("Sex")
```
```r
#reverse colour order
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
  geom_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
             position = position_jitter(0.1)) +
  scale_color_grafify(palette = "bright", reverse = TRUE) +
  facet_wrap("Sex")
```

---

### scale_color_grafify2

**Scale colour colour scheme**

**Description**

`grafify` internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that these **only** work for categorical variables. Use the brewer or viridis packages for numeric gradient scales.

**Usage**

```r
scale_color_grafify2(palette = "all_grafify", reverse = FALSE, ...)
```

**Arguments**

- **palette**
  - Name of the colour scheme. Default set to `palette = "all_grafify"`. Provide names as above in quotes.
- **reverse**
  - Whether the colour order should be reversed.
- **...**
  - Additional parameters for `scale_fill` or `scale_colour`.

**Details**

The default for `scale_fill_grafify()`, `scale_colour_grafify()` or `scale_color_grafify()` is a list of 55 colours as part of `palette = "all_grafify"`.

Obviously, it is not recommended to use so many colours, but implementing this was easiest to prevent errors when using a lot of categorical variables.

Colours available can be seen quickly with `plot_grafify_palette`. There are eight palettes with 5-10 colours each, which are recommended. These can be called by naming the colour scheme using `palette =` argument. Additional options include "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", and "contrast". These are taken from Paul Taul, Mike Mol and Okabe Ito. `scale_fill_grafify2` and `scale_colour_grafify2` are identical except that when the number of categorical variables is fewer than the total number of colour shades in the palette (e.g. if you have 3 groups and the "okabe_ito" palette has 7 colours), these functions will pick the most 'distant' colours from the scheme than going sequentially. If you want colours assigned sequentially use `scale_fill_grafify` or `scale_colour_grafify`.

**Value**

`ggplot` scale_fill function for discrete colours.
Examples

# add a colour scheme to a ggplot object
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
  geom_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
             position = position_jitter(0.15)) +
  scale_color_grafify2(palette = "bright") + facet_wrap("Sex")

# reverse colour order
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
  geom_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
             position = position_jitter(0.1)) +
  scale_color_grafify2(palette = "bright", reverse = TRUE) + facet_wrap("Sex")

scale_color_grafify_c  Scale_color continuous scheme

description

grafify internally includes color-blind compatible schemes for fill and color/color aesthetics. Note that this scheme is only for continuous variables and has one palette (yellow_conti) modified from the YlOrBr scheme from RColorBrewer.

Usage

scale_color_grafify_c(reverse = FALSE, ...)

Arguments

reverse  Whether the color order should be reversed.
...
additional parameters for scale_fill or scale_color.

details

Colour palettes available are as follows:
Colours available can be seen quickly with plot_grafify_palette.

Value

ggplot scale_fill function for continuous colours.

Examples

# basic usage on mtcars data with x and y quantitative axes
ggplot(mtcars, aes(x = mpg, y = disp)) +
  geom_point(aes(color = disp), size = 3) +
  scale_color_grafify_c()
scale_colour_grafify  Scale_colour colour scheme

Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that these only work for categorical variables. Use the brewer or viridis packages for numeric gradient scales.

Usage

scale_colour_grafify(palette = "all_grafify", reverse = FALSE, ...)

Arguments

palette  Name of the colour scheme. Default set to palette = "all_grafify". Provide names as above in quotes.
reverse  Whether the colour order should be reversed.
...  Additional parameters for scale_fill or scale_colour.

Details

The default for scale_fill_grafify(), scale_colour_grafify() or scale_color_grafify() is a list of 55 colours as part of palette = "all_grafify". Obviously, it is not recommended to use so many colours, but implementing this was easiest to prevent errors when using a lot of categorical variables.

Colours available can be seen quickly with plot_grafify_palette. There are eight palettes with 5-10 colours each, which are recommended. These can be called by naming the colour scheme using palette = argument. Additional options include "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", and "contrast". These are taken from Paul Taul, Mike Mol and Okabe Ito. scale_fill_grafify2 and scale_colour_grafify2 are identical except that when the number of categorical variables is fewer than the total number of colour shades in the palette (e.g. if you have 3 groups and the "okabe_ito" palette has 7 colours), these functions will pick the most ‘distant’ colours from the scheme than going sequentially. If you want colours assigned sequentially use scale_fill_grafify or scale_colour_grafify.

Value

ggplot scale_fill function for discrete colours.

Examples

#add a colour scheme to a ggplot object
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration))+
  geom_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
             position = position_jitter(0.15) )+
  scale_colour_grafify(palette = "bright")+facet_wrap("Sex")
#reverse colour order
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
geom_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
position = position_jitter(0.1)) +
scale_colour_grafify(palette = "bright", reverse = TRUE) + facet_wrap("Sex")

scale_colour_grafify2  Scale_colour colour scheme

Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that these only work for categorical variables. Use the brewer or viridis packages for numeric gradient scales.

Usage

scale_colour_grafify2(palette = "all_grafify", reverse = FALSE, ...)

Arguments

palette  Name of the colour scheme. Default set to palette = "all_grafify". Provide names as above in quotes.
reverse  Whether the colour order should be reversed.
...  Additional parameters for scale_fill or scale_colour.

Details

The default for scale_fill_grafify(), scale_colour_grafify() or scale_color_grafify() is a list of 55 colours as part of palette = "all_grafify".

Obviously, it is not recommended to use so many colours, but implementing this was easiest to prevent errors when using a lot of categorical variables.

Colours available can be seen quickly with plot_grafify_palette. There are eight palettes with 5-10 colours each, which are recommended. These can be called by naming the colour scheme using palette = argument. Additional options include "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", and "contrast". These are taken from Paul Taul, Mike Mol and Okabe Ito. scale_fill_grafify2 and scale_colour_grafify2 are identical except that when the number of categorical variables is fewer than the total number of colour shades in the palette (e.g. if you have 3 groups and the "okabe_ito" palette has 7 colours), these functions will pick the most 'distant' colours from the scheme than going sequentially. If you want colours assigned sequentially use scale_fill_grafify or scale_colour_grafify.

Value

ggplot scale_fill function for discrete colours.
Examples

# add a colour scheme to a ggplot object
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
gem_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
  position = position_jitter(0.15)) +
scale_color_grafify2(palette = "bright") + facet_wrap("Sex")

# reverse colour order
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
gem_point(aes(colour = Treatment, shape = Sex), size = 3, alpha = 0.9,
  position = position_jitter(0.1)) +
scale_color_grafify2(palette = "bright", reverse = TRUE) + facet_wrap("Sex")

scale_colour_grafify_c

Scale_colour continuous scheme

Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that this scheme is only for continuous variables and has one palette (yellow_conti) modified from the YlOrBr scheme from RColorBrewer.

Usage

scale_colour_grafify_c(reverse = FALSE, ...)

Arguments

reverse Whether the colour order should be reversed.

... Additional parameters for scale_fill or scale_colour.

Details

Colours available can be seen quickly with plot_grafify_palette.

Value

ggplot scale_fill function for discrete colours.

Examples

# basic usage on mtcars data with x and y quantitative axes
ggplot(mtcars, aes(x = mpg, y = disp)) +
gem_point(aes(colour = disp), size = 3) +
scale_colour_grafify_c()
scale_fill_grafify

Scale_fill colour scheme

Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that these only work for categorical variables. Use the brewer or viridis packages for numeric gradient scales.

Usage

scale_fill_grafify(palette = "all_grafify", reverse = FALSE, ...)

Arguments

palette Name of the colour scheme. Default set to palette = "all_grafify". Provide names as above in quotes.
reverse Whether the colour order should be reversed.
... Additional parameters for scale_fill or scale_colour.

Details

The default for scale_fill_grafify(), scale_colour_grafify() or scale_color_grafify() is a list of 55 colours as part of palette = "all_grafify".

Obviously, it is not recommended to use so many colours, but implementing this was easiest to prevent errors when using a lot of categorical variables.

Colours available can be seen quickly with plot_grafify_palette. There are eight palettes with 5-10 colours each, which are recommended. These can be called by naming the colour scheme using palette = argument. Additional options include "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", and "contrast". These are taken from Paul Taul, Mike Mol and Okabe Ito.

scale_fill_grafify2 and scale_colour_grafify2 are identical except that when the number of categorical variables is fewer than the total number of colour shades in the palette (e.g. if you have 3 groups and the "okabe_ito" palette has 7 colours), these functions will pick the most 'distant' colours from the scheme than going sequentially. If you want colours assigned sequentially use scale_fill_grafify or scale_colour_grafify.

Value

ggplot scale_fill function for discrete colours.

Examples

#add a grafify fill scheme to ggplot
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration))+
  geom_point(aes(fill = Treatment), shape = 21, size = 3,
  position = position_jitter(0.15), alpha = 0.8)+
  scale_fill_grafify(palette = "muted")+facet_wrap("Sex")
# reverse colour order
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
geom_point(aes(fill = Treatment), shape = 21, size = 3,
position = position_jitter(0.15), alpha = 0.8) +
scale_fill_grafify2(palette = "muted", reverse = TRUE) + facet_wrap("Sex")

scale_fill_grafify2  Scale_fill colour scheme

Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics. Note that these only work for categorical variables. Use the brewer or viridis packages for numeric gradient scales.

Usage

scale_fill_grafify2(palette = "all_grafify", reverse = FALSE, ...)

Arguments

- **palette**  Name of the colour scheme. Default set to palette = "all_grafify". Provide names as above in quotes.
- **reverse**  Whether the colour order should be reversed.
- **...**  Additional parameters for scale_fill or scale_colour.

Details

The default for scale_fill_grafify(), scale_colour_grafify() or scale_color_grafify() is a list of 55 colours as part of palette = "all_grafify".

Obviously, it is not recommended to use so many colours, but implementing this was easiest to prevent errors when using a lot of categorical variables.

Colours available can be seen quickly with plot_grafify_palette. There are eight palettes with 5-10 colours each, which are recommended. These can be called by naming the colour scheme using palette = argument. Additional options include "okabe_ito", "vibrant", "bright", "pale", "muted", "dark", "light", and "contrast". These are taken from Paul Taul, Mike Mol and Okabe Ito. scale_fill_grafify2 and scale_colour_grafify2 are identical except that when the number of categorical variables is fewer than the total number of colour shades in the palette (e.g. if you have 3 groups and the "okabe_ito" palette has 7 colours), these functions will pick the most 'distant' colours from the scheme than going sequentially. If you want colours assigned sequentially use scale_fill_grafify or scale_colour_grafify.

Value

ggplot scale_fill function for discrete colours.
scale_fill_grafify_c

Examples

# add a grafify fill scheme to ggplot
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
  geom_point(aes(fill = Treatment), shape = 21, size = 3,
  position = position_jitter(0.15), alpha = 0.8) +
  scale_fill_grafify2(palette = "muted") +
  facet_wrap("Sex")
# reverse colour order
ggplot(emmeans::neuralgia, aes(x = Treatment, y = Duration)) +
  geom_point(aes(fill = Treatment), shape = 21, size = 3,
  position = position_jitter(0.15), alpha = 0.8) +
  scale_fill_grafify2(palette = "muted", reverse = TRUE) +
  facet_wrap("Sex")

scale_fill_grafify_c  Scale_fill continuous scheme

Description

grafify internally includes colour-blind compatible schemes for fill and colour/color aesthetics.
Note that this scheme is only for continuous variables and has one palette (yellow_conti) modified
from the YlOrBr scheme from RColorBrewer.

Usage

scale_fill_grafify_c(reverse = FALSE, ...)

Arguments

reverse  Whether the colour order should be reversed.
...

Additional parameters for scale_fill or scale_colour.

Details

Colours available can be seen quickly with plot_grafify_palette.

Value

ggplot scale_fill function for continuous colours.

Examples

# basic usage on mtcars data with x and y quantitative axes
ggplot(mtcars, aes(x = mpg, y = disp)) +
  geom_point(aes(fill = disp), shape = 21, size = 3) +
  scale_fill_grafify_c()
simple_anova

ANOVA table from a linear model fit to data.

Description

Update in v0.2.1: This function uses \texttt{lm} to fit a linear model to data, passes it on to \texttt{Anova}, and outputs the ANOVA table with type II sum of squares with F statistics and \textit{P} values. (Previous versions produced type I sum of squares using \texttt{anova} call.)

Usage

\texttt{simple_anova(data, Y\_value, Fixed\_Factor, ...)}

Arguments

- \texttt{data}: a data table object, e.g. \texttt{data.frame} or \texttt{tibble}.
- \texttt{Y\_value}: name of column containing quantitative (dependent) variable, provided within "quotes".
- \texttt{Fixed\_Factor}: name(s) of categorical fixed factors (independent variables) provided as a vector if more than one or within "quotes".
- \texttt{...}: any additional argument to pass on to \texttt{lm} if required.

Details

It requires a data table, one quantitative dependent variable and one or more independent variables. If your experiment design has random factors, use the related function \texttt{mixed_anova}.

This function is related to \texttt{link{simple_model}}.

Value

ANOVA table of class "anova" and "data.frame".

Examples

# Basic usage
simple_anova(data = data\_doubling\_time,
Y\_value = "Doubling\_time",
Fixed\_Factor = "Student")
simple_model

Model from a linear model fit to data.

Description

This function uses \texttt{lm} to fit a linear model to data and outputs the model object. It requires a data table, one quantitative dependent variable and one or more independent variables. The model output can be used to extract coefficients and other information, including post-hoc comparisons. If your experiment design has random factors, use the related function \texttt{mixed_model}.

Usage

\begin{verbatim}
simple_model(data, Y_value, Fixed_Factor, ...)
\end{verbatim}

Arguments

- \texttt{data} a data table object, e.g. data.frame or tibble.
- \texttt{Y_value} name of column containing quantitative (dependent) variable, provided within "quotes".
- \texttt{Fixed_Factor} name(s) of categorical fixed factors (independent variables) provided as a vector if more than one or within "quotes".
- \texttt{...} any additional arguments to pass on to \texttt{lm} if required.

Details

This function is related to \texttt{link{simple_anova}}. Output of this function can be used with \texttt{posthoc_Pairwise}, \texttt{posthoc_Levelwise} and \texttt{posthoc_vsRef}, or with \texttt{emmeans}.

Value

This function returns an object of class "lm".

Examples

\begin{verbatim}
#fixed factors provided as a vector
Doubmodel <- simple_model(data = data_doubling_time, Y_value = "Doubling_time", Fixed_Factor = "Student")
#get summary
summary(Doubmodel)
\end{verbatim}
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