

Package ‘ggforestplotR’

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Title Publication-Ready Forest Plots with 'ggplot2'

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Description Transform model coefficients into flexible forest plots using 'ggplot2'. Provides helpers to standardize coefficient data from a range of modelling workflows and render publication-ready forest plots with a consistent interface.

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add_forest_table	<i>Add a summary table to a forest plot</i>
------------------	---

Description

Compose a summary table onto a forest plot.

Usage

```
add_forest_table(  
  plot = NULL,  
  position = c("left", "right"),  
  columns = NULL,  
  term_header = "Term",  
  n_header = "N",  
  events_header = "Events",  
  estimate_label = NULL,  
  p_header = "P-value",  
  column_labels = NULL,  
  digits = NULL,  
  estimate_digits = NULL,  
  interval_digits = NULL,  
  p_digits = NULL,  
  estimate_fmt = NULL,  
  ci_fmt = NULL,  
  text_size = NULL,  
  header_text_size = NULL,  
  header_fontface = "bold",  
  header_family = NULL,  
  striped_rows = NULL,  
  stripe_fill = NULL,  
  stripe_colour = NULL,  
  stripe_alpha = NULL,  
  grid_lines = FALSE,  
  grid_line_colour = "black",  
  grid_line_size = 0.3,  
  grid_line_linetype = 1  
)
```

Arguments

plot	A plot created by <code>ggforestplot()</code> . Leave as NULL to use <code>+ add_forest_table(...)</code> syntax.
position	Whether to place the table on the left or right of the forest plot.
columns	Optional explicit columns to display in the side table, in the order they should appear. Accepts built-in names such as "term", "n", "events", "estimate", "ci", and "p", arbitrary original dataframe columns, or numeric positions in the supplied data. "conf.low" and "conf.high" are accepted as aliases for "ci".
term_header	Header text for the term column.
n_header	Header text for the N column.
events_header	Header text for the Events column.
estimate_label	Header label for the estimate column. Defaults to the model-derived label when available.
p_header	Header text for the p-value column.
column_labels	Optional named vector used to relabel table column headers. Names should match values supplied to columns after column resolution, such as "term", "estimate", "ci", "p", or an arbitrary original dataframe column.
digits	Deprecated. Number of digits used when formatting estimates and p-values. Defaults to 2. Use <code>estimate_digits</code> , <code>interval_digits</code> , and <code>p_digits</code> for separate control.
estimate_digits	Number of digits used for point estimates.
interval_digits	Number of digits used for confidence interval bounds.
p_digits	Number of digits used for p-values.
estimate_fmt	Format string for the estimate column. Use <code>{estimate}</code> , <code>{conf.low}</code> , and <code>{conf.high}</code> as placeholders. The shorthand <code>{conf.low, conf.high}</code> is also supported. Defaults to <code>"{estimate} ({conf.low}, {conf.high})"</code> , or <code>"{estimate}"</code> when columns includes "ci".
ci_fmt	Format string for the confidence interval column when columns includes "ci". Use <code>{conf.low}</code> and <code>{conf.high}</code> as placeholders. The shorthand <code>{conf.low, conf.high}</code> is also supported. Defaults to <code>"({conf.low}, {conf.high})"</code> .
text_size	Text size for table contents. Defaults to 3.2.
header_text_size	Header text size for table column labels. Defaults to 11.
header_fontface	Font face used for table column labels. Defaults to "bold".
header_family	Optional font family used for table column labels.
striped_rows	Whether to draw alternating row stripes behind the table. Defaults to the stripe setting used in <code>ggforestplot()</code> .
stripe_fill	Fill colour used for striped rows. Defaults to the stripe fill used in <code>ggforestplot()</code> .
stripe_colour	Outline colour for striped rows. Defaults to the stripe outline used in <code>ggforestplot()</code> .

stripe_alpha Transparency for striped rows. Defaults to the stripe alpha used in `ggforestplot()`.
grid_lines Whether to draw black horizontal grid lines in the table.
grid_line_colour Colour used for the table grid lines.
grid_line_size Line width used for the table grid lines.
grid_line_linetype Line type used for the table grid lines.

Value

A patchwork-composed plot containing the forest plot and side table, or a ggplot add-on object when `plot = NULL`.

Examples

```

coefs <- data.frame(
  term = c("Age", "BMI", "Treatment"),
  estimate = c(0.3, -0.2, 0.4),
  conf.low = c(0.1, -0.4, 0.2),
  conf.high = c(0.5, 0.0, 0.6),
  sample_size = c(120, 115, 98),
  p_value = c(0.012, 0.031, 0.004)
)

p <- ggforestplot(coefs, n = "sample_size", p.value = "p_value")
add_forest_table(
  p,
  position = "left",
  columns = c("term", "n", "estimate", "p"),
  estimate_label = "Beta"
)

ggforestplot(coefs, n = "sample_size", p.value = "p_value") +
  add_forest_table(
    position = "right",
    columns = c("term", "n", "estimate", "p"),
    estimate_label = "Beta"
  )

```

add_split_table

Add split tables around a forest plot

Description

Compose split table blocks around a forest plot so that summary data appear on both sides of the plotting panel.

Usage

```

add_split_table(
  plot = NULL,
  left_columns = NULL,
  right_columns = NULL,
  term_header = "Term",
  n_header = "N",
  events_header = "Events",
  estimate_label = NULL,
  p_header = "P-value",
  column_labels = NULL,
  digits = NULL,
  estimate_digits = NULL,
  interval_digits = NULL,
  p_digits = NULL,
  estimate_fmt = NULL,
  ci_fmt = NULL,
  text_size = NULL,
  header_text_size = NULL,
  header_fontface = "bold",
  header_family = NULL,
  striped_rows = NULL,
  stripe_fill = NULL,
  stripe_colour = NULL,
  stripe_alpha = NULL,
  left_width = NULL,
  plot_width = NULL,
  right_width = NULL
)

```

Arguments

plot	A plot created by <code>ggforestplot()</code> . Leave as NULL to use + <code>add_split_table(...)</code> syntax.
left_columns	Optional explicit columns to place on the left side of the forest plot. Accepts built-in names such as "term", "n", "events", "estimate", "ci", and "p", arbitrary original dataframe columns, or numeric positions in the supplied data. "conf.low" and "conf.high" are accepted as aliases for "ci".
right_columns	Optional explicit columns to place on the right side of the forest plot. Accepts built-in names such as "estimate", "ci", and "p", arbitrary original dataframe columns, or numeric positions in the supplied data. "conf.low" and "conf.high" are accepted as aliases for "ci".
term_header	Header text for the term column.
n_header	Header text for the N column.
events_header	Header text for the Events column.
estimate_label	Header label for the estimate column. Defaults to the model-derived label when available.

p_header	Header text for the p-value column.
column_labels	Optional named vector used to relabel table column headers. Names should match values supplied to left_columns or right_columns after column resolution, such as "term", "estimate", "ci", "p", or an arbitrary original dataframe column.
digits	Deprecated. Number of digits used when formatting estimates and p-values. Defaults to 2. Use estimate_digits, interval_digits, and p_digits for separate control.
estimate_digits	Number of digits used for point estimates.
interval_digits	Number of digits used for confidence interval bounds.
p_digits	Number of digits used for p-values.
estimate_fmt	Format string for the estimate column. Use {estimate}, {conf.low}, and {conf.high} as placeholders. The shorthand {conf.low, conf.high} is also supported. Defaults to "{estimate} ({conf.low}, {conf.high})", or "{estimate}" when table columns include "ci".
ci_fmt	Format string for the confidence interval column when table columns include "ci". Use {conf.low} and {conf.high} as placeholders. The shorthand {conf.low, conf.high} is also supported. Defaults to "({conf.low}, {conf.high})".
text_size	Text size for table contents. Defaults to 3.2.
header_text_size	Header text size for table column labels. Defaults to 11.
header_fontface	Font face used for table column labels. Defaults to "bold".
header_family	Optional font family used for table column labels.
striped_rows	Whether to draw alternating row stripes behind the split table layout. Defaults to the stripe setting used in ggforestplot() .
stripe_fill	Fill colour used for striped rows. Defaults to the stripe fill used in ggforestplot() .
stripe_colour	Outline colour for striped rows. Defaults to the stripe outline used in ggforestplot() .
stripe_alpha	Transparency for striped rows. Defaults to the stripe alpha used in ggforestplot() .
left_width	Optional width allocated to the left table block. By default this is derived from the number of displayed left-side columns relative to plot_width.
plot_width	Optional width allocated to the forest plot panel. Defaults to 2.5.
right_width	Optional width allocated to the right table block. By default this is derived from the number of displayed right-side columns relative to plot_width.

Value

A patchwork-composed plot containing a left table, the forest plot, and a right table, or a ggplot add-on object when plot = NULL.

Examples

```

coefs <- data.frame(
  term = c("Age", "BMI", "Treatment"),
  estimate = c(0.3, -0.2, 0.4),
  conf.low = c(0.1, -0.4, 0.2),
  conf.high = c(0.5, 0.0, 0.6),
  sample_size = c(120, 115, 98),
  p_value = c(0.012, 0.031, 0.004)
)

p <- ggforestplot(coefs, n = "sample_size", p.value = "p_value")
add_split_table(
  p,
  left_columns = c("term", "n"),
  right_columns = c("estimate", "p"),
  estimate_label = "HR"
)

ggforestplot(coefs, n = "sample_size", p.value = "p_value") +
  add_split_table(
    left_columns = c(1, 5),
    right_columns = c(2, 6),
    estimate_label = "HR"
  )

```

as_forest_data

*Standardize coefficient data for forest plots***Description**

Standardizes a coefficient table into the internal forest-plot data structure used throughout ggforestplotR.

Usage

```

as_forest_data(
  data,
  term,
  estimate,
  conf.low,
  conf.high,
  label = term,
  term_labels = NULL,
  group = NULL,
  grouping = NULL,
  separate_groups = NULL,
  n = NULL,
  events = NULL,
  p.value = NULL,

```

```

    exponentiate = FALSE,
    sort_terms = c("none", "descending", "ascending")
  )

```

Arguments

<code>data</code>	A data frame containing coefficient estimates and intervals.
<code>term</code>	Column name holding the model term identifier.
<code>estimate</code>	Column name holding the point estimate.
<code>conf.low</code>	Column name holding the lower confidence bound.
<code>conf.high</code>	Column name holding the upper confidence bound.
<code>label</code>	Optional column name used for the displayed row label.
<code>term_labels</code>	Optional named vector used to relabel displayed terms. Names should match values in the term column and values are the labels to display.
<code>group</code>	Optional column name used for color-grouping multiple estimates per row.
<code>grouping</code>	Optional column name used to split rows into grouped plot sections.
<code>separate_groups</code>	Optional column name used to identify labeled variable blocks that can be outlined with separator lines.
<code>n</code>	Optional column name holding sample sizes or other N labels for table helpers.
<code>events</code>	Optional column name holding event counts or event labels for table helpers.
<code>p.value</code>	Optional column name holding p-values.
<code>exponentiate</code>	Logical; if TRUE, require positive values for estimates and intervals.
<code>sort_terms</code>	How to sort rows: "none", "descending", or "ascending".

Value

A standardized data frame ready for `ggforestplot()` and the table composition helpers. Original dataframe columns are retained for table helpers so they can be displayed with `add_forest_table(columns = ...)`.

Examples

```

raw <- data.frame(
  variable = c("Age", "BMI", "Treatment"),
  beta = c(0.10, -0.08, 0.34),
  lower = c(0.02, -0.16, 0.12),
  upper = c(0.18, 0.00, 0.56)
)

as_forest_data(
  data = raw,
  term = "variable",
  estimate = "beta",
  conf.low = "lower",
  conf.high = "upper"
)

```

bind_forest_models	<i>Bind multiple model summaries for a grouped forest plot</i>
--------------------	--

Description

Tidies multiple fitted models and stacks their fixed-effect coefficient tables into a single forest-plot data frame. The resulting data can be passed directly to `ggforestplot()`, where model labels are used as the grouping variable for dodged, color-coded estimates.

Usage

```
bind_forest_models(models, model_labels = NULL, exponentiate = NULL, ...)
```

Arguments

<code>models</code>	A non-empty list of fitted model objects supported by <code>tidy_forest_model()</code> .
<code>model_labels</code>	Optional labels used to identify each model in the forest plot. Defaults to list names when present, otherwise "Model 1", "Model 2", and so on.
<code>exponentiate</code>	NULL, a single logical value, or one logical value per model. NULL uses the canonical scale inferred by <code>tidy_forest_model()</code> for each model.
<code>...</code>	Additional arguments passed to <code>tidy_forest_model()</code> , such as <code>conf.level</code> , <code>intercept</code> , <code>term_labels</code> , or <code>sort_terms</code> .

Value

A standardized forest-plot data frame with one row per model term and a group column containing the model labels.

Examples

```
if (requireNamespace("broom", quietly = TRUE)) {
  fit1 <- lm(mpg ~ wt + hp, data = mtcars)
  fit2 <- lm(mpg ~ wt + qsec, data = mtcars)

  bound <- bind_forest_models(
    list(Base = fit1, Adjusted = fit2)
  )

  ggforestplot(bound)
}
```

`ggforestplot`*Draw a ggplot2 forest plot*

Description

Builds a forest plot from standardized coefficient data or directly from a fitted model.

Usage

```
ggforestplot(  
  data,  
  term = "term",  
  estimate = "estimate",  
  conf.low = "conf.low",  
  conf.high = "conf.high",  
  label = term,  
  term_labels = NULL,  
  group = NULL,  
  facet = NULL,  
  facet_strip_position = c("left", "right"),  
  grouping = NULL,  
  grouping_strip_position = NULL,  
  separate_groups = NULL,  
  n = NULL,  
  events = NULL,  
  p.value = NULL,  
  exponentiate = NULL,  
  sort_terms = c("none", "descending", "ascending"),  
  point_size = 2.3,  
  point_shape = 19,  
  linewidth = 0.5,  
  line_size = NULL,  
  staple_width = 0.2,  
  ci_limits = NULL,  
  ci_arrows = TRUE,  
  ci_arrow_length = 0.08,  
  ci_arrow_type = c("closed", "open"),  
  dodge_width = 0.6,  
  separate_lines = FALSE,  
  separator_line_linetype = 2,  
  separator_line_colour = "black",  
  separator_line_size = 0.4,  
  striped_rows = FALSE,  
  stripe_fill = "grey95",  
  stripe_colour = NA,  
  stripe_alpha = 1,  
  ref_line = NULL,
```

```

    ref_label = NULL,
    ref_linetype = 2,
    ref_color = "grey60"
  )

```

Arguments

<code>data</code>	Either a tidy coefficient data frame or a model object supported by <code>broom::tidy()</code> .
<code>term</code>	Column name holding the model term identifiers.
<code>estimate</code>	Column name holding the point estimates.
<code>conf.low</code>	Column name holding the lower confidence bounds.
<code>conf.high</code>	Column name holding the upper confidence bounds.
<code>label</code>	Optional column name used for the displayed row labels.
<code>term_labels</code>	Optional named vector used to relabel displayed terms. Names should match values in the term column and values are the labels to display.
<code>group</code>	Optional column name used for color-grouping estimates.
<code>facet</code>	Optional column name used to split rows into faceted plot sections. If this column is a factor, its levels control facet order.
<code>facet_strip_position</code>	Positioning for facet strip labels.
<code>grouping</code>	Deprecated. Use <code>facet</code> instead.
<code>grouping_strip_position</code>	Deprecated. Use <code>facet_strip_position</code> instead.
<code>separate_groups</code>	Optional column name used to identify labeled variable blocks that can be outlined with grid lines.
<code>n</code>	Optional column name holding sample sizes or other N labels for table helpers.
<code>events</code>	Optional column name holding event counts or event labels for table helpers.
<code>p.value</code>	Optional column name holding p-values.
<code>exponentiate</code>	Logical; if TRUE, transform the estimates and draw the axis on the log scale with the reference line at 1. For model objects, NULL uses the canonical scale when it can be inferred, such as hazard ratios for Cox models.
<code>sort_terms</code>	How to sort rows: "none", "descending", or "ascending".
<code>point_size</code>	Point size for coefficient markers.
<code>point_shape</code>	Shape used for coefficient markers.
<code>linewidth</code>	Line width for confidence intervals.
<code>line_size</code>	Deprecated. Use <code>linewidth</code> instead.
<code>staple_width</code>	Width of the terminal staples on confidence interval lines.
<code>ci_limits</code>	Optional numeric vector of length 2 used to truncate displayed confidence intervals. Intervals extending beyond these limits are clipped to the boundary and marked with arrows when <code>ci_arrows</code> is TRUE.

<code>ci_arrows</code>	Logical; if TRUE, draw outward-facing arrows for confidence intervals truncated by <code>ci_limits</code> .
<code>ci_arrow_length</code>	Length of CI truncation arrows in inches.
<code>ci_arrow_type</code>	Arrowhead type for truncated confidence intervals. Passed to <code>grid::arrow()</code> .
<code>dodge_width</code>	Horizontal dodging used for grouped estimates.
<code>separate_lines</code>	Logical; if TRUE, draw grid lines around each labeled block identified by <code>separate_groups</code> .
<code>separator_line_linetype</code>	Line type used for separator lines.
<code>separator_line_colour</code>	Colour used for separator lines.
<code>separator_line_size</code>	Line width used for separator lines.
<code>striped_rows</code>	Logical; if TRUE, shade alternating rows.
<code>stripe_fill</code>	Fill color used for shaded rows.
<code>stripe_colour</code>	Border color for shaded rows.
<code>stripe_alpha</code>	Transparency for shaded rows.
<code>ref_line</code>	Numeric x-value where the reference line is drawn, or NULL to hide it. When omitted, defaults to 0 for additive effects and 1 for exponentiated effects.
<code>ref_label</code>	Optional label drawn alongside the reference line.
<code>ref_linetype</code>	Line type for the reference line.
<code>ref_color</code>	Color for the reference line.

Value

A ggplot object. Use standard ggplot2 functions such as `ggplot2::labs()` for plot labels, and add composition helpers after styling the main plot.

Examples

```

coefs <- data.frame(
  term = c("Age", "BMI", "Treatment"),
  estimate = c(0.10, -0.08, 0.34),
  conf.low = c(0.02, -0.16, 0.12),
  conf.high = c(0.18, 0.00, 0.56)
)

ggforestplot(coefs)

ggforestplot(coefs, striped_rows = TRUE, point_shape = 17)

```

tidy_forest_model	<i>Tidy a model object for forest plotting</i>
-------------------	--

Description

Uses `broom::tidy()` to convert a fitted model into forest-plot data. Mixed models are supported through `broom.mixed` tidy methods when that package is installed.

Usage

```
tidy_forest_model(
  model,
  conf.int = TRUE,
  conf.level = 0.95,
  exponentiate = NULL,
  intercept = FALSE,
  term_labels = NULL,
  sort_terms = c("none", "descending", "ascending")
)
```

Arguments

<code>model</code>	A fitted model object supported by <code>broom::tidy()</code> or, for mixed models, a <code>broom.mixed</code> tidy method.
<code>conf.int</code>	Logical; if TRUE, request confidence intervals from <code>broom::tidy()</code> .
<code>conf.level</code>	Confidence level for intervals.
<code>exponentiate</code>	Logical; passed through to <code>broom::tidy()</code> .
<code>intercept</code>	Logical; if FALSE, drop the intercept term.
<code>term_labels</code>	Optional named vector used to relabel displayed terms. Names should match model term names and values are the labels to display.
<code>sort_terms</code>	How to sort rows: "none", "descending", or "ascending".

Value

A standardized coefficient data frame ready for `ggforestplot()`.

Examples

```
if (requireNamespace("broom", quietly = TRUE)) {
  fit <- lm(mpg ~ wt + hp + qsec, data = mtcars)
  tidy_forest_model(fit)

  set.seed(123)
  logit_data <- data.frame(
    age = rnorm(250, mean = 62, sd = 8),
    bmi = rnorm(250, mean = 28, sd = 4),
```

```
    treatment = factor(rbinom(250, 1, 0.45), labels = c("Control", "Treatment"))
  )
  linpred <- -9 + 0.09 * logit_data$age + 0.11 * logit_data$bmi +
    0.9 * (logit_data$treatment == "Treatment")
  logit_data$event <- rbinom(250, 1, plogis(linpred))
  logit_fit <- glm(event ~ age + bmi + treatment, data = logit_data, family = binomial())

  tidy_forest_model(logit_fit, exponentiate = TRUE)
}
```

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