

# Package ‘tidyplots’

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**Title** Tidy Plots for Scientific Papers

**Version** 0.2.1

**Description** The goal of ‘tidyplots’ is to streamline the creation of publication-ready plots for scientific papers. It allows to gradually add, remove and adjust plot components using a consistent and intuitive syntax.

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**Encoding** UTF-8

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**Imports** cli, dplyr, forcats, ggbeeswarm, ggplot2 (>= 3.5.0), ggpubr, ggrastr, ggrepel, glue, Hmisc, htmltools, lifecycle, patchwork (>= 1.2.0), purrr, rlang, scales, stringr, tidyr, tidyselect

**Depends** R (>= 2.10)

**LazyData** true

**URL** <https://github.com/jbengler/tidyplots>,  
<https://jbengler.github.io/tidyplots/>

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**NeedsCompilation** no

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---

add	<i>Add ggplot2 code to a tidyplot</i>
-----	---------------------------------------

---

## Description

Add ggplot2 code to a tidyplot

## Usage

```
add()
```

## Value

A tidyplot object.

## Examples

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add(ggplot2::geom_point())
```

---

add\_annotation\_text    *Add annotation*

---

### Description

Add annotation

### Usage

```
add_annotation_text(plot, text, x, y, fontsize = 7)

add_annotation_rectangle(
  plot,
  xmin,
  xmax,
  ymin,
  ymax,
  fill = "#000000",
  color = NA,
  alpha = 0.1
)

add_annotation_line(plot, x, xend, y, yend, color = "#000000")
```

### Arguments

plot	A tidyplot generated with the function tidyplot().
text	Here goes the description.
x, xmin, xmax, xend, y, ymin, ymax, yend	Here goes the description.
fontsize	Font size in points. Defaults to 7.
fill	A hex color for the fill color. For example, "#FFFFFF" for white.
color	A hex color for the stroke color. For example, "#FFFFFF" for white.
alpha	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.

### Value

A tidyplot object.

### Examples

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_boxplot() %>%
  add_annotation_text("Look here!", x = 2, y = 25)
```

```
eu_countries %>%
  tidyplot(x = area, y = population) %>%
  add_data_points() %>%
  add_annotation_rectangle(xmin = 2.5e5, xmax = Inf, ymin = 42, ymax = Inf)

eu_countries %>%
  tidyplot(x = area, y = population) %>%
  add_data_points() %>%
  add_annotation_rectangle(xmin = 2.5e5, xmax = 6e5, ymin = 42, ymax = 90,
    color = "#E69F00", fill = NA)

eu_countries %>%
  tidyplot(x = area, y = population) %>%
  add_data_points() %>%
  add_annotation_line(x = 0, xend = Inf, y = 0, yend = Inf)
```

---

add\_areastack\_absolute

*Add area stack*

---

## Description

Add area stack

## Usage

```
add_areastack_absolute(
  plot,
  linewidth = 0.25,
  alpha = 0.4,
  reverse = FALSE,
  replace_na = FALSE,
  ...
)
```

```
add_areastack_relative(
  plot,
  linewidth = 0.25,
  alpha = 0.4,
  reverse = FALSE,
  replace_na = FALSE,
  ...
)
```

**Arguments**

plot	A tidyplot generated with the function tidyplot().
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
alpha	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.
reverse	Whether the order should be reversed or not. Defaults to FALSE, meaning not reversed.
replace_na	Whether to replace count = NA with count = 0.
...	Arguments passed on to the geom function.

**Value**

A tidyplot object.

**Examples**

```
# for a `count` provide `x` and `color`
# `count` of the data points in each `energy_type` category
energy %>%
  tidyplot(x = year, color = energy_type) %>%
  add_areastack_absolute()

energy %>%
  tidyplot(x = year, color = energy_type) %>%
  add_areastack_relative()

# for a `sum` provide `x`, `y` and `color`
# `sum` of `energy` in each `energy_type` category
energy %>%
  tidyplot(x = year, y = energy, color = energy_type) %>%
  add_areastack_absolute()

energy %>%
  tidyplot(x = year, y = energy, color = energy_type) %>%
  add_areastack_relative()

# Flip x and y-axis
energy %>%
  tidyplot(x = energy, y = year, color = energy_type) %>%
  add_areastack_absolute(orientation = "y")

energy %>%
  tidyplot(x = energy, y = year, color = energy_type) %>%
  add_areastack_relative(orientation = "y")
```

---

add\_barstack\_absolute *Add bar stack*

---

## Description

Add bar stack

## Usage

```
add_barstack_absolute(plot, width = 0.8, reverse = FALSE, ...)
```

```
add_barstack_relative(plot, width = 0.8, reverse = FALSE, ...)
```

## Arguments

plot	A tidyplot generated with the function tidyplot().
width	Width of the plot area. Defaults to 50.
reverse	Whether the order should be reversed or not. Defaults to FALSE, meaning not reversed.
...	Arguments passed on to the geom function.

## Value

A tidyplot object.

## Examples

```
# for a `count` only provide `color`
# `count` of the data points in each `energy_type` category
energy %>%
  tidyplot(color = energy_type) %>%
  add_barstack_absolute()

energy %>%
  tidyplot(color = energy_type) %>%
  add_barstack_relative()

# for a `sum` provide `color` and `y`
# `sum` of `energy` in each `energy_type` category
energy %>%
  tidyplot(y = energy, color = energy_type) %>%
  add_barstack_absolute()

energy %>%
  tidyplot(y = energy, color = energy_type) %>%
  add_barstack_relative()

# Include variable on second axis
```

```
energy %>%
  tidyplot(x = year, y = energy, color = energy_type) %>%
  add_barstack_absolute()

energy %>%
  tidyplot(x = year, y = energy, color = energy_type) %>%
  add_barstack_relative()

# Flip x and y-axis
energy %>%
  tidyplot(x = energy, y = year, color = energy_type) %>%
  add_barstack_absolute(orientation = "y")

energy %>%
  tidyplot(x = energy, y = year, color = energy_type) %>%
  add_barstack_relative(orientation = "y")
```

---

add\_boxplot

*Add boxplot*

---

## Description

Add boxplot

## Usage

```
add_boxplot(
  plot,
  dodge_width = NULL,
  alpha = 0.3,
  saturation = 1,
  show_whiskers = TRUE,
  show_outliers = TRUE,
  box_width = 0.6,
  whiskers_width = 0.8,
  outlier.size = 0.5,
  coef = 1.5,
  outlier.shape = 19,
  outlier.alpha = 1,
  linewidth = 0.25,
  preserve = "total",
  ...
)
```

## Arguments

**plot** A tidyplot generated with the function tidyplot().



dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
alpha	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
show_whiskers	Whether to show boxplot whiskers. Defaults to TRUE.
show_outliers	Whether to show outliers. Defaults to TRUE.
box_width	Width of the boxplot. Defaults to 0.6.
whiskers_width	Width of the whiskers. Defaults to 0.8.
outlier.size	Size of the outliers. Defaults to 0.5.
coef	Length of the whiskers as multiple of IQR. Defaults to 1.5.
outlier.shape	Shape of the outliers. Defaults to 19.
outlier.alpha	Opacity of the outliers. Defaults to 1.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.

**Value**

A tidyplot object.

**Examples**

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_boxplot()

# Changing arguments:
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_boxplot(show_whiskers = FALSE)

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_boxplot(show_outliers = FALSE)

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_boxplot(box_width = 0.2)

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_boxplot(whiskers_width = 0.2)
```

---

`add_count_bar`*Add count*

---

**Description**

Add count

**Usage**

```
add_count_bar(  
  plot,  
  dodge_width = NULL,  
  width = 0.6,  
  saturation = 1,  
  preserve = "total",  
  ...  
)
```

```
add_count_dash(  
  plot,  
  dodge_width = NULL,  
  width = 0.6,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

```
add_count_dot(plot, dodge_width = NULL, size = 2, preserve = "total", ...)
```

```
add_count_value(  
  plot,  
  dodge_width = NULL,  
  accuracy = 0.1,  
  scale_cut = NULL,  
  fontsize = 7,  
  extra_padding = 0.15,  
  vjust = NULL,  
  hjust = NULL,  
  preserve = "total",  
  ...  
)
```

```
add_count_line(  
  plot,  
  group,  
  dodge_width = NULL,  
  linewidth = 0.25,
```

```

    preserve = "total",
    ...
  )

add_count_area(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  preserve = "total",
  ...
)

```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
width	Width of the plot area. Defaults to 50.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
size	A number representing the size of the plot symbol. Typical values range between 1 and 3.
accuracy	A number to round to. Use (e.g.) 0.01 to show 2 decimal places of precision. If NULL, the default, uses a heuristic that should ensure breaks have the minimum number of digits needed to show the difference between adjacent values. Applied to rescaled data.
scale_cut	Scale cut function to be applied. See <code>scales::cut_short_scale()</code> and friends.
fontsize	Font size in points. Defaults to 7.
extra_padding	Extra padding to create space for the value label.
vjust	Vertical position adjustment of the value label.
hjust	Horizontal position adjustment of the value label.
group	Variable in the dataset to be used for grouping.

### Value

A tidyplot object.

**Examples**

```

dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_bar()

dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_dash()

dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_dot()

dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_value()

dinosaurs %>%
  tidyplot(x = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_line()

dinosaurs %>%
  tidyplot(x = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_area()

# Combination
dinosaurs %>%
  tidyplot(x = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_bar(alpha = 0.4) %>%
  add_count_dash() %>%
  add_count_dot() %>%
  add_count_value() %>%
  add_count_line()

# Changing arguments: alpha
# Makes objects transparent
dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  theme_minimal_y() %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_bar(alpha = 0.4)

# Changing arguments: saturation
# Reduces fill color saturation without making the object transparent
dinosaurs %>%

```

```

tidyplot(x = time_lived, color = time_lived) %>%
  theme_minimal_y() %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_bar(saturation = 0.3)

# Changing arguments: accuracy
dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_value(accuracy = 1)

# Changing arguments: fontsize
dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_value(fontsize = 10)

# Changing arguments: color
dinosaurs %>%
  tidyplot(x = time_lived, color = time_lived) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_count_value(color = "black")

```

---

add\_curve\_fit

*Add curve fit*


---

## Description

Add curve fit

## Usage

```

add_curve_fit(
  plot,
  dodge_width = NULL,
  method = "loess",
  linewidth = 0.25,
  alpha = 0.4,
  preserve = "total",
  ...
)

```

## Arguments

**plot** A tidyplot generated with the function `tidyplot()`.

**dodge\_width** For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.

method	<p>Smoothing method (function) to use, accepts either NULL or a character vector, e.g. "lm", "glm", "gam", "loess" or a function, e.g. MASS::rlm or mgcv::gam, stats::lm, or stats::loess. "auto" is also accepted for backwards compatibility. It is equivalent to NULL.</p> <p>For method = NULL the smoothing method is chosen based on the size of the largest group (across all panels). stats::loess() is used for less than 1,000 observations; otherwise mgcv::gam() is used with formula = y ~ s(x, bs = "cs") with method = "REML". Somewhat anecdotally, loess gives a better appearance, but is <math>O(N^2)</math> in memory, so does not work for larger datasets.</p> <p>If you have fewer than 1,000 observations but want to use the same gam() model that method = NULL would use, then set method = "gam", formula = y ~ s(x, bs = "cs").</p>
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
alpha	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to ggplot2::geom_smooth().

## Value

A tidyplot object.

## Examples

```
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_curve_fit()

# Changing arguments
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_curve_fit(linewidth = 1)

time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_curve_fit(alpha = 0.8)

# Remove confidence interval
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_curve_fit(se = FALSE)
```

---

add_data_labels	<i>Add data labels</i>
-----------------	------------------------

---

## Description

Add data labels

## Usage

```
add_data_labels(  
  plot,  
  label,  
  data = all_rows(),  
  fontsize = 7,  
  background = FALSE,  
  background_color = "#FFFFFF",  
  background_alpha = 0.6,  
  label_position = c("below", "above", "left", "right", "center"),  
  ...  
)
```

```
add_data_labels_repel(  
  plot,  
  label,  
  data = all_rows(),  
  fontsize = 7,  
  segment.size = 0.2,  
  box.padding = 0.2,  
  max.overlaps = Inf,  
  background = FALSE,  
  background_color = "#FFFFFF",  
  background_alpha = 0.6,  
  ...  
)
```

## Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
label	Variable in the dataset to be used for the text label.
data	The data to be displayed in this layer. There are three options: <ul style="list-style-type: none"><li>• If <code>all_rows()</code> (the default) the complete dataset is displayed.</li><li>• A function to subset the plot data. See <code>filter_rows()</code> and friends.</li><li>• A <code>data.frame</code> to override the plot data.</li></ul>
fontsize	Font size in points. Defaults to 7.
background	Whether to include semitransparent background box behind the labels to improve legibility. Defaults to <code>FALSE</code> .

background_color	Hex color of the background box. Defaults to "#FFFFFF" for white.
background_alpha	Opacity of the background box. Defaults to 0.6.
label_position	Position of the label in relation to the data point. Can be one of c("below", "above", "left", "right", "center").
...	Arguments passed on to the geom function.
segment.size	Thickness of the line connecting the label with the data point. Defaults to 0.2.
box.padding	Amount of padding around bounding box, as unit or number. Defaults to 0.25. (Default unit is lines, but other units can be specified by passing unit(x, "units")).
max.overlaps	Exclude text labels when they overlap too many other things. For each text label, we count how many other text labels or other data points it overlaps, and exclude the text label if it has too many overlaps. Defaults to 10.

### Details

- `add_data_labels_repel()` uses `ggrepel::geom_text_repel()`. Check there and in [ggrepel examples](#) for additional arguments.
- `add_data_labels()` and `add_data_labels_repel()` support data subsetting. See [Advanced plotting](#).

### Value

A tidyplot object.

### Examples

```
# Create plot and increase padding to make more space for labels
p <-
  animals %>%
  dplyr::slice_head(n = 5) %>%
  tidyplot(x = weight, y = speed) %>%
  theme_ggplot2() %>%
  add_data_points() %>%
  adjust_padding(all = 0.3)

# Default label position is `below` the data point
p %>% add_data_labels(label = animal)

# Alternative label positions
p %>% add_data_labels(label = animal, label_position = "above")

p %>% add_data_labels(label = animal, label_position = "right")

p %>% add_data_labels(label = animal, label_position = "left")

# Include white background box
p %>% add_data_labels(label = animal, background = TRUE)
```



```
p %>% add_data_labels(label = animal, background = TRUE,
  background_color = "pink")

# Black labels
p %>% add_data_labels(label = animal, color = "black")

# Use repelling data labels
p %>% add_data_labels_repel(label = animal, color = "black")

p %>% add_data_labels_repel(label = animal, color = "black",
  background = TRUE)

p %>% add_data_labels_repel(label = animal, color = "black",
  background = TRUE, min.segment.length = 0)
```

---

add_data_points	<i>Add data points</i>
-----------------	------------------------

---

## Description

Add data points

## Usage

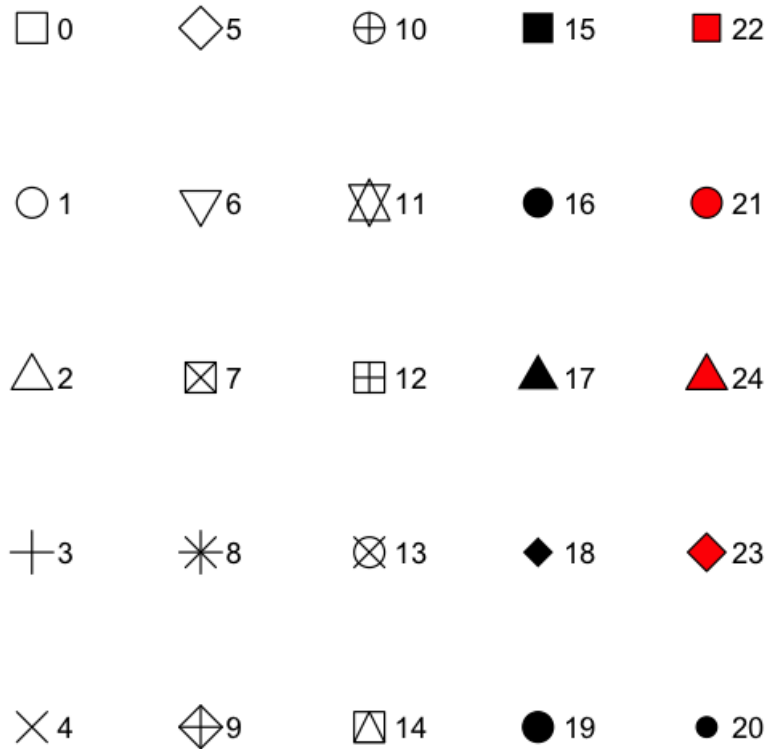
```
add_data_points(
  plot,
  data = all_rows(),
  shape = 19,
  size = 1,
  white_border = FALSE,
  dodge_width = NULL,
  preserve = "total",
  rasterize = FALSE,
  rasterize_dpi = 300,
  ...
)

add_data_points_jitter(
  plot,
  data = all_rows(),
  shape = 19,
  size = 1,
  white_border = FALSE,
  dodge_width = NULL,
  jitter_width = 0.2,
  jitter_height = 0,
  preserve = "total",
```

```
rasterize = FALSE,  
rasterize_dpi = 300,  
...  
)  
  
add_data_points_beeswarm(  
  plot,  
  data = all_rows(),  
  shape = 19,  
  size = 1,  
  white_border = FALSE,  
  cex = 3,  
  corral = "wrap",  
  corral.width = 0.5,  
  dodge_width = NULL,  
  preserve = "total",  
  rasterize = FALSE,  
  rasterize_dpi = 300,  
  ...  
)
```

## Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
data	The data to be displayed in this layer. There are three options: <ul style="list-style-type: none"><li>• If <code>all_rows()</code> (the default) the complete dataset is displayed.</li><li>• A function to subset the plot data. See <code>filter_rows()</code> and friends.</li><li>• A <code>data.frame</code> to override the plot data.</li></ul>
shape	An integer between 0 and 24, representing the shape of the plot symbol.



size	A number representing the size of the plot symbol. Typical values range between 1 and 3.
white_border	Whether to include a white border around data points. Defaults to FALSE.
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
rasterize	If FALSE (the default) the layer will be constructed of vector shapes. If TRUE the layer will be rasterized to a pixel image. This can be useful when plotting many individual objects (1,000 or more) compromises the performance of the generated PDF file.
rasterize_dpi	The resolution in dots per inch (dpi) used for rastering the layer if rasterize is TRUE. The default is 300 dpi.
...	Arguments passed on to the geom function.
jitter_width	Amount of random noise to be added to the horizontal position of the of the data points. This can be useful to deal with overplotting. Typical values range between 0 and 1.
jitter_height	Amount of random noise to be added to the vertical position of the of the data points. This can be useful to deal with overplotting. Typical values range between 0 and 1.
cex	Scaling for adjusting point spacing (see <a href="#">beeswarm::swarmx()</a> ). Values between 1 (default) and 3 tend to work best.

`corral` string. Method used to adjust points that would be placed too wide horizontally, default is "none". See details below.

`corral.width` numeric. Width of the corral, default is 0.9.

### Details

- `add_data_points_beeswarm()` is based on `ggbeeswarm::geom_beeswarm()`. Check there for additional arguments.
- `add_data_points()` and friends support rasterization. See examples and [Advanced plotting](#).
- `add_data_points()` and friends support data subsetting. See examples and [Advanced plotting](#).

### Value

A `tidyplot` object.

### Examples

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_jitter()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm()

# Changing arguments
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_jitter(jitter_width = 1)

animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points(white_border = TRUE)

animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points(alpha = 0.4)

# Rasterization
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points(rasterize = TRUE, rasterize_dpi = 50)

# Data subsetting
animals %>%
  tidyplot(x = weight, y = size) %>%
```

```
add_data_points() %>%  
add_data_points(data = filter_rows(size > 300), color = "red")
```

---

**add\_heatmap***Add heatmap*

---

## Description

Add heatmap

## Usage

```
add_heatmap(  
  plot,  
  scale = c("none", "row", "column"),  
  rotate_labels = 90,  
  rasterize = FALSE,  
  rasterize_dpi = 300,  
  ...  
)
```

## Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
scale	Whether to compute row z scores for "row" or "column". Defaults to "none".
rotate_labels	Degree to rotate the x-axis labels. Defaults to 90.
rasterize	If FALSE (the default) the layer will be constructed of vector shapes. If TRUE the layer will be rasterized to a pixel image. This can be useful when plotting many individual objects (1,000 or more) compromises the performance of the generated PDF file.
rasterize_dpi	The resolution in dots per inch (dpi) used for rastering the layer if rasterize is TRUE. The default is 300 dpi.
...	Arguments passed on to the geom function.

## Details

- `add_heatmap()` supports rasterization. See examples and [Advanced plotting](#).

## Value

A tidyplot object.

**Examples**

```

climate %>%
  tidyplot(x = month, y = year, color = max_temperature) %>%
  add_heatmap()

# Calculate row-wise z score
climate %>%
  tidyplot(x = month, y = year, color = max_temperature) %>%
  add_heatmap(scale = "row")

# Calculate column-wise z score
climate %>%
  tidyplot(x = month, y = year, color = max_temperature) %>%
  add_heatmap(scale = "column")

# Rasterize heatmap
climate %>%
  tidyplot(x = month, y = year, color = max_temperature) %>%
  add_heatmap(rasterize = TRUE, rasterize_dpi = 20)

```

---

`add_histogram`*Add histogram*

---

**Description**

Add histogram

**Usage**`add_histogram(plot, binwidth = NULL, bins = NULL, ...)`**Arguments**

<code>plot</code>	A tidyplot generated with the function <code>tidyplot()</code> .
<code>binwidth</code>	The width of the bins. Can be specified as a numeric value or as a function that calculates width from unscaled x. Here, "unscaled x" refers to the original x values in the data, before application of any scale transformation. When specifying a function along with a grouping structure, the function will be called once per group. The default is to use the number of bins in <code>bins</code> , covering the range of the data. You should always override this value, exploring multiple widths to find the best to illustrate the stories in your data. The bin width of a date variable is the number of days in each time; the bin width of a time variable is the number of seconds.
<code>bins</code>	Number of bins. Overridden by <code>binwidth</code> . Defaults to 30.
<code>...</code>	Arguments passed on to the geom function.

**Value**

A tidyplot object.

**Examples**

```
energy %>%
  tidyplot(x = energy) %>%
  add_histogram()

energy %>%
  tidyplot(x = energy, color = energy_type) %>%
  add_histogram()
```

---

add_line	<i>Add line or area</i>
----------	-------------------------

---

**Description**

add\_line() and add\_area() connect individual data points, which is rarely needed. In most cases, you are probably looking for add\_sum\_line(), add\_mean\_line(), add\_sum\_area() or add\_mean\_area().

**Usage**

```
add_line(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  preserve = "total",
  ...
)

add_area(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  alpha = 0.4,
  preserve = "total",
  ...
)
```

**Arguments**

plot	A tidyplot generated with the function tidyplot().
group	Variable in the dataset to be used for grouping.
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.
alpha	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.

**Value**

A tidyplot object.

**Examples**

```
# Paired data points
study %>%
  tidyplot(x = treatment, y = score, color = group) %>%
  reorder_x_axis_labels("A", "C", "B", "D") %>%
  add_data_points() %>%
  add_line(group = participant, color = "grey")

study %>%
  tidyplot(x = treatment, y = score) %>%
  reorder_x_axis_labels("A", "C", "B", "D") %>%
  add_data_points() %>%
  add_area(group = participant)
```

---

add\_mean\_bar

*Add mean*

---

**Description**

Add mean

**Usage**

```
add_mean_bar(
  plot,
  dodge_width = NULL,
  width = 0.6,
  saturation = 1,
```



```
    preserve = "total",
    ...
  )

add_mean_dash(
  plot,
  dodge_width = NULL,
  width = 0.6,
  linewidth = 0.25,
  preserve = "total",
  ...
)

add_mean_dot(plot, dodge_width = NULL, size = 2, preserve = "total", ...)

add_mean_value(
  plot,
  dodge_width = NULL,
  accuracy = 0.1,
  scale_cut = NULL,
  fontsize = 7,
  extra_padding = 0.15,
  vjust = NULL,
  hjust = NULL,
  preserve = "total",
  ...
)

add_mean_line(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  preserve = "total",
  ...
)

add_mean_area(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  preserve = "total",
  ...
)
```

### Arguments

`plot` A tidyplot generated with the function `tidyplot()`.

dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
width	Width of the plot area. Defaults to 50.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
size	A number representing the size of the plot symbol. Typical values range between 1 and 3.
accuracy	A number to round to. Use (e.g.) 0.01 to show 2 decimal places of precision. If NULL, the default, uses a heuristic that should ensure breaks have the minimum number of digits needed to show the difference between adjacent values. Applied to rescaled data.
scale_cut	Scale cut function to be applied. See scales::cut_short_scale() and friends.
fontsize	Font size in points. Defaults to 7.
extra_padding	Extra padding to create space for the value label.
vjust	Vertical position adjustment of the value label.
hjust	Horizontal position adjustment of the value label.
group	Variable in the dataset to be used for grouping.

### Value

A tidyplot object.

### Examples

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar()
```

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_dash()
```

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_dot()
```

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_value()
```

```
study %>%
  tidyplot(x = treatment, y = score) %>%
```

```

    add_mean_line()

study %>%
  tidyplot(x = treatment, y = score) %>%
  add_mean_area()

# Combination
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_mean_dash() %>%
  add_mean_dot() %>%
  add_mean_value() %>%
  add_mean_line()

# Changing arguments: alpha
# Makes objects transparent
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  theme_minimal_y() %>%
  add_mean_bar(alpha = 0.4)

# Changing arguments: saturation
# Reduces fill color saturation without making the object transparent
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  theme_minimal_y() %>%
  add_mean_bar(saturation = 0.3)

# Changing arguments: accuracy
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_value(accuracy = 0.01)

# Changing arguments: fontsize
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_value(fontsize = 10)

# Changing arguments: color
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_value(color = "black")

```

---

add\_median\_bar

*Add median*


---

## Description

Add median

**Usage**

```
add_median_bar(  
  plot,  
  dodge_width = NULL,  
  width = 0.6,  
  saturation = 1,  
  preserve = "total",  
  ...  
)
```

```
add_median_dash(  
  plot,  
  dodge_width = NULL,  
  width = 0.6,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

```
add_median_dot(plot, dodge_width = NULL, size = 2, preserve = "total", ...)
```

```
add_median_value(  
  plot,  
  dodge_width = NULL,  
  accuracy = 0.1,  
  scale_cut = NULL,  
  fontsize = 7,  
  extra_padding = 0.15,  
  vjust = NULL,  
  hjust = NULL,  
  preserve = "total",  
  ...  
)
```

```
add_median_line(  
  plot,  
  group,  
  dodge_width = NULL,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

```
add_median_area(  
  plot,  
  group,  
  dodge_width = NULL,  
  linewidth = 0.25,
```

```

    preserve = "total",
    ...
)

```

### Arguments

plot	A tidyplot generated with the function tidyplot().
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
width	Width of the plot area. Defaults to 50.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
size	A number representing the size of the plot symbol. Typical values range between 1 and 3.
accuracy	A number to round to. Use (e.g.) 0.01 to show 2 decimal places of precision. If NULL, the default, uses a heuristic that should ensure breaks have the minimum number of digits needed to show the difference between adjacent values. Applied to rescaled data.
scale_cut	Scale cut function to be applied. See scales::cut_short_scale() and friends.
fontsize	Font size in points. Defaults to 7.
extra_padding	Extra padding to create space for the value label.
vjust	Vertical position adjustment of the value label.
hjust	Horizontal position adjustment of the value label.
group	Variable in the dataset to be used for grouping.

### Value

A tidyplot object.

### Examples

```

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_bar()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_dash()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_dot()

```

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_value()

study %>%
  tidyplot(x = treatment, y = score) %>%
  add_median_line()

study %>%
  tidyplot(x = treatment, y = score) %>%
  add_median_area()

# Combination
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_median_bar(alpha = 0.4) %>%
  add_median_dash() %>%
  add_median_dot() %>%
  add_median_value() %>%
  add_median_line()

# Changing arguments: alpha
# Makes objects transparent
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  theme_minimal_y() %>%
  add_median_bar(alpha = 0.4)

# Changing arguments: saturation
# Reduces fill color saturation without making the object transparent
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  theme_minimal_y() %>%
  add_median_bar(saturation = 0.3)

# Changing arguments: accuracy
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_value(accuracy = 0.01)

# Changing arguments: fontsize
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_value(fontsize = 10)

# Changing arguments: color
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_median_value(color = "black")
```

---

add_pie	<i>Add pie or donut chart</i>
---------	-------------------------------

---

### Description

Add pie or donut chart

### Usage

```
add_pie(plot, width = 1, reverse = FALSE, ...)
```

```
add_donut(plot, width = 1, reverse = FALSE, ...)
```

### Arguments

plot	A tidyplot generated with the function tidyplot().
width	Width of the plot area. Defaults to 50.
reverse	Whether the order should be reversed or not. Defaults to FALSE, meaning not reversed.
...	Arguments passed on to the geom function.

### Value

A tidyplot object.

### Examples

```
# for a `count` only provide `color`
# `count` of the data points in each `energy_type` category
energy %>%
  tidyplot(color = energy_type) %>%
  add_pie()

energy %>%
  tidyplot(color = energy_type) %>%
  add_donut()

# for a `sum` provide `color` and `y`
# `sum` of `energy` in each `energy_type` category
energy %>%
  tidyplot(y = energy, color = energy_type) %>%
  add_pie()

energy %>%
  tidyplot(y = energy, color = energy_type) %>%
  add_donut()
```

---

add\_reference\_lines    *Add reference lines*

---

## Description

Add reference lines

## Usage

```
add_reference_lines(
  plot,
  x = NULL,
  y = NULL,
  linetype = "dashed",
  linewidth = 0.25,
  ...
)
```

## Arguments

plot	A tidyplot generated with the function tidyplot().
x	Numeric values where the reference lines should meet the x-axis. For example, x = 4 or x = c(2, 3, 4).
y	Numeric values where the reference lines should meet the y-axis. For example, y = 4 or y = c(2, 3, 4).
linetype	Either an integer (0-6) or a name (0 = blank, 1 = solid, 2 = dashed, 3 = dotted, 4 = dotdash, 5 = longdash, 6 = twodash).
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
...	Arguments passed on to the geom function.

## Value

A tidyplot object.

## Examples

```
animals %>%
  tidyplot(x = weight, y = speed) %>%
  add_reference_lines(x = 4000, y = c(100, 200)) %>%
  add_data_points()

animals %>%
  tidyplot(x = weight, y = speed) %>%
  add_reference_lines(x = 4000, y = c(100, 200), linetype = "dotdash") %>%
  add_data_points()
```



---

add_sem_errorbar	<i>Add error bar</i>
------------------	----------------------

---

**Description**

- add\_sem\_errorbar() adds the standard error of mean.
- add\_range\_errorbar() adds the range from smallest to largest value.
- add\_sd\_errorbar() adds the standard deviation.
- add\_ci95\_errorbar() adds the 95% confidence interval.

**Usage**

```
add_sem_errorbar(  
  plot,  
  dodge_width = NULL,  
  width = 0.4,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

```
add_range_errorbar(  
  plot,  
  dodge_width = NULL,  
  width = 0.4,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

```
add_sd_errorbar(  
  plot,  
  dodge_width = NULL,  
  width = 0.4,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

```
add_ci95_errorbar(  
  plot,  
  dodge_width = NULL,  
  width = 0.4,  
  linewidth = 0.25,  
  preserve = "total",  
  ...  
)
```

**Arguments**

plot	A tidyplot generated with the function tidyplot().
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
width	Width of the plot area. Defaults to 50.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.

**Value**

A tidyplot object.

**Examples**

```
# Standard error of the mean
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar()

# Range from minimum to maximum value
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_range_errorbar()

# Standard deviation
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sd_errorbar()

# 95% confidence interval
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_ci95_errorbar()

# Changing arguments: error bar width
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar(width = 0.8)

# Changing arguments: error bar line width
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
```

```
add_sem_errorbar(linewidth = 1)
```

---

```
add_sem_ribbon          Add ribbon
```

---

### Description

- `add_sem_ribbon()` adds the standard error of mean.
- `add_range_ribbon()` adds the range from smallest to largest value.
- `add_sd_ribbon()` adds the standard deviation.
- `add_ci95_ribbon()` adds the 95% confidence interval.

### Usage

```
add_sem_ribbon(plot, dodge_width = NULL, alpha = 0.4, color = NA, ...)
```

```
add_range_ribbon(plot, dodge_width = NULL, alpha = 0.4, color = NA, ...)
```

```
add_sd_ribbon(plot, dodge_width = NULL, alpha = 0.4, color = NA, ...)
```

```
add_ci95_ribbon(plot, dodge_width = NULL, alpha = 0.4, color = NA, ...)
```

### Arguments

<code>plot</code>	A tidyplot generated with the function <code>tidyplot()</code> .
<code>dodge_width</code>	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
<code>alpha</code>	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.
<code>color</code>	A hex color for the stroke color. For example, "#FFFFFF" for white.
<code>...</code>	Arguments passed on to the geom function.

### Value

A tidyplot object.

### Examples

```
# Standard error of the mean
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_mean_line() %>%
  add_sem_ribbon()

# Range from minimum to maximum value
```

```

time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_mean_line() %>%
  add_range_ribbon()

# Standard deviation
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_mean_line() %>%
  add_sd_ribbon()

# 95% confidence interval
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_mean_line() %>%
  add_ci95_ribbon()

# Changing arguments: alpha
time_course %>%
  tidyplot(x = day, y = score, color = treatment) %>%
  add_mean_line() %>%
  add_sem_ribbon(alpha = 0.7)

```

---

add\_sum\_bar

*Add sum*


---

## Description

Add sum

## Usage

```

add_sum_bar(
  plot,
  dodge_width = NULL,
  width = 0.6,
  saturation = 1,
  preserve = "total",
  ...
)

```

```

add_sum_dash(
  plot,
  dodge_width = NULL,
  width = 0.6,
  linewidth = 0.25,
  preserve = "total",
  ...
)

```

```

)

add_sum_dot(plot, dodge_width = NULL, size = 2, preserve = "total", ...)

add_sum_value(
  plot,
  dodge_width = NULL,
  accuracy = 0.1,
  scale_cut = NULL,
  fontsize = 7,
  extra_padding = 0.15,
  vjust = NULL,
  hjust = NULL,
  preserve = "total",
  ...
)

add_sum_line(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  preserve = "total",
  ...
)

add_sum_area(
  plot,
  group,
  dodge_width = NULL,
  linewidth = 0.25,
  preserve = "total",
  ...
)

```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
width	Width of the plot area. Defaults to 50.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
preserve	Should dodging preserve the "total" width of all elements at a position, or the width of a "single" element?
...	Arguments passed on to the geom function.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.

size	A number representing the size of the plot symbol. Typical values range between 1 and 3.
accuracy	A number to round to. Use (e.g.) 0.01 to show 2 decimal places of precision. If NULL, the default, uses a heuristic that should ensure breaks have the minimum number of digits needed to show the difference between adjacent values. Applied to rescaled data.
scale_cut	Scale cut function to be applied. See <code>scales::cut_short_scale()</code> and friends.
fontsize	Font size in points. Defaults to 7.
extra_padding	Extra padding to create space for the value label.
vjust	Vertical position adjustment of the value label.
hjust	Horizontal position adjustment of the value label.
group	Variable in the dataset to be used for grouping.

### Value

A tidyplot object.

### Examples

```
spendings %>%
  tidyplot(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_bar()
```

```
spendings %>%
  tidyplot(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_dash()
```

```
spendings %>%
  tidyplot(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_dot()
```

```
spendings %>%
  tidyplot(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_value()
```

```
spendings %>%
  tidyplot(x = category, y = amount) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_line()
```

```
spendings %>%
  tidyplot(x = category, y = amount) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_area()
```

```
# Combination
spendings %>%
  tidypLOT(x = category, y = amount) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_median_bar(alpha = 0.4) %>%
  add_median_dash() %>%
  add_median_dot() %>%
  add_median_value() %>%
  add_median_line()

# Changing arguments: alpha
# Makes objects transparent
spendings %>%
  tidypLOT(x = category, y = amount, color = category) %>%
  theme_minimal_y() %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_bar(alpha = 0.4)

# Changing arguments: saturation
# Reduces fill color saturation without making the object transparent
spendings %>%
  tidypLOT(x = category, y = amount, color = category) %>%
  theme_minimal_y() %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_bar(saturation = 0.3)

# Changing arguments: accuracy
spendings %>%
  tidypLOT(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_value(accuracy = 1)

# Changing arguments: fontsize
spendings %>%
  tidypLOT(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_value(fontsize = 10)

# Changing arguments: color
spendings %>%
  tidypLOT(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_value(color = "black")

# Changing arguments: extra_padding
spendings %>%
  tidypLOT(x = category, y = amount, color = category) %>%
  adjust_x_axis(rotate_labels = TRUE) %>%
  add_sum_value(extra_padding = 0.5)
```

---

add_test_pvalue	<i>Add statistical test</i>
-----------------	-----------------------------

---

**Description**

Add statistical test

**Usage**

```
add_test_pvalue(  
  plot,  
  padding_top = 0.15,  
  method = "t.test",  
  p.adjust.method = "none",  
  ref.group = NULL,  
  label = "{format_p_value(p.adj, 0.0001)}",  
  label.size = 7/ggplot2::.pt,  
  step.increase = 0.15,  
  vjust = -0.25,  
  bracket.nudge.y = 0.1,  
  hide.ns = FALSE,  
  p.adjust.by = "panel",  
  symnum.args = list(cutpoints = c(0, 0.001, 0.01, 0.05, Inf), symbols = c("***", "**",  
    "*", "ns")),  
  hide_info = FALSE,  
  ...  
)  
  
add_test_asterisks(  
  plot,  
  padding_top = 0.1,  
  method = "t.test",  
  p.adjust.method = "none",  
  ref.group = NULL,  
  label = "p.adj.signif",  
  label.size = 10/ggplot2::.pt,  
  step.increase = 0.2,  
  vjust = 0.3,  
  bracket.nudge.y = 0.15,  
  hide.ns = TRUE,  
  p.adjust.by = "panel",  
  symnum.args = list(cutpoints = c(0, 0.001, 0.01, 0.05, Inf), symbols = c("***", "**",  
    "*", "ns")),  
  hide_info = FALSE,  
  ...  
)
```



**Arguments**

plot	A tidyplot generated with the function tidyplot().
padding_top	Extra padding above the data points to accommodate the statistical comparisons.
method	a character string indicating which method to be used for pairwise comparisons. Default is "wilcox_test". Allowed methods include pairwise comparisons methods implemented in the rstatix R package. These methods are: "wilcox_test", "t_test", "sign_test", "dunn_test", "emmeans_test", "tukey_hsd", "games_howell_test".
p.adjust.method	method for adjusting p values (see <a href="#">p.adjust</a> ). Has impact only in a situation, where multiple pairwise tests are performed; or when there are multiple grouping variables. Ignored when the specified method is "tukey_hsd" or "games_howell_test" because they come with internal p adjustment method. Allowed values include "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none". If you don't want to adjust the p value (not recommended), use p.adjust.method = "none".
ref.group	a character string or a numeric value specifying the reference group. If specified, for a given grouping variable, each of the group levels will be compared to the reference group (i.e. control group). ref.group can be also "all". In this case, each of the grouping variable levels is compared to all (i.e. basemean). Allowed values can be: <ul style="list-style-type: none"> <li>• <b>numeric value:</b> specifying the rank of the reference group. For example, use ref.group = 1 when the first group is the reference; use ref.group = 2 when the second group is the reference, and so on. This works for all situations, including i) when comparisons are performed between x-axis groups and ii) when comparisons are performed between legend groups.</li> <li>• <b>character value:</b> For example, you can use ref.group = "ctrl" instead of using the numeric rank value of the "ctrl" group.</li> <li>• <b>"all":</b> In this case, each of the grouping variable levels is compared to all (i.e. basemean).</li> </ul>
label	character string specifying label. Can be: <ul style="list-style-type: none"> <li>• the column containing the label (e.g.: label = "p" or label = "p.adj"), where p is the p-value. Other possible values are "p.signif", "p.adj.signif", "p.format", "p.adj.format".</li> <li>• an expression that can be formatted by the <a href="#">glue()</a> package. For example, when specifying label = "Wilcoxon, p = <math>\{p\}</math>", the expression {p} will be replaced by its value.</li> <li>• a combination of plotmath expressions and glue expressions. You may want some of the statistical parameter in italic; for example: label = "Wilcoxon, <i>italic(p)</i> = {p}"</li> </ul>
label.size	change the size of the label text
step.increase	numeric vector with the increase in fraction of total height for every additional comparison to minimize overlap.

vjust	move the text up or down relative to the bracket.
bracket.nudge.y	Vertical adjustment to nudge brackets by (in fraction of the total height). Useful to move up or move down the bracket. If positive value, brackets will be moved up; if negative value, brackets are moved down.
hide.ns	can be logical value (TRUE or FALSE) or a character vector ("p.adj" or "p").
p.adjust.by	possible value is one of c("group", "panel"). Default is "group": for a grouped data, if pairwise test is performed, then the p-values are adjusted for each group level independently. P-values are adjusted by panel when p.adjust.by = "panel".
symnum.args	a list of arguments to pass to the function <code>symnum</code> for symbolic number coding of p-values. For example, <code>symnum.args &lt;- list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, Inf), symbols = c("****", "***", "**", "*", "ns"))</code> . In other words, we use the following convention for symbols indicating statistical significance: <ul style="list-style-type: none"> <li>• ns: <math>p &gt; 0.05</math></li> <li>• *: <math>p \leq 0.05</math></li> <li>• **: <math>p \leq 0.01</math></li> <li>• ***: <math>p \leq 0.001</math></li> <li>• ****: <math>p \leq 0.0001</math></li> </ul>
hide_info	Whether to hide details about the statistical testing as caption. Defaults to FALSE.
...	Arguments passed on to <code>ggpubr::geom_pwc()</code> .

### Details

- `add_test_pvalue()` and `add_test_asterisks()` use `ggpubr::geom_pwc()`. Check there for additional arguments.

### Value

A `tidyplot` object.

### Examples

```
study %>%
  tidyplot(x = dose, y = score, color = group) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points() %>%
  add_test_pvalue()

# Change stat methods
study %>%
  tidyplot(x = dose, y = score, color = group) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points() %>%
```

```

    add_test_pvalue(method = "wilcoxon", p.adjust.method = "BH")

# Define reference group to test against
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points() %>%
  add_test_pvalue(ref.group = "A")

# hide non-significant p values
gene_expression %>%
  # filter to one gene
  dplyr::filter(external_gene_name == "Apol6") %>%
  # start plotting
  tidyplot(x = condition, y = expression, color = sample_type) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points() %>%
  add_test_pvalue(hide.ns = TRUE)

# Adjust top padding for statistical comparisons
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points() %>%
  add_test_pvalue(padding_top = 0.08)

# Hide stats information
study %>%
  tidyplot(x = dose, y = score, color = group) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points() %>%
  add_test_pvalue(hide_info = TRUE)

```

---

add_title	<i>Add plot title or caption</i>
-----------	----------------------------------

---

## Description

Add plot title or caption

## Usage

```
add_title(plot, title = ggplot2::waiver())
```

```
add_caption(plot, caption = ggplot2::waiver())
```

**Arguments**

plot	A tidyplot generated with the function tidyplot().
title	Title of the plot.
caption	Caption of the plot.

**Details**

- add\_title() and add\_caption() support **plotmath expressions** to include special characters. See examples and [Advanced plotting](#).

**Value**

A tidyplot object.

**Examples**

```
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points_beeswarm() %>%
  add_title("This is my title")

study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points_beeswarm() %>%
  add_caption("This is the fine print in the caption")

# Plotmath expression
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points_beeswarm() %>%
  add_title("$H[2]*0~and~E==m*c^{2}$")
```

---

add\_violin

*Add violin plot*

---

**Description**

Add violin plot

**Usage**

```
add_violin(
  plot,
  dodge_width = NULL,
  alpha = 0.3,
  saturation = 1,
  draw_quantiles = NULL,
```

```

    trim = FALSE,
    linewidth = 0.25,
    scale = "width",
    ...
  )

```

### Arguments

plot	A tidyplot generated with the function tidyplot().
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.
alpha	A number between 0 and 1 for the opacity of an object. A value of 0 is completely transparent, 1 is completely opaque.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
draw_quantiles	If not(NULL) (default), draw horizontal lines at the given quantiles of the density estimate.
trim	If TRUE (default), trim the tails of the violins to the range of the data. If FALSE, don't trim the tails.
linewidth	Thickness of the line in points (pt). Typical values range between 0.25 and 1.
scale	if "area" (default), all violins have the same area (before trimming the tails). If "count", areas are scaled proportionally to the number of observations. If "width", all violins have the same maximum width.
...	Arguments passed on to the geom function.

### Value

A tidyplot object.

### Examples

```

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_violin()

# Changing arguments:
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_violin(saturation = 0.6)

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_violin(draw_quantiles = c(0.25, 0.5, 0.75))

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_violin(trim = TRUE)

```

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_violin(linewidth = 1)
```

---

adjust_colors	<i>Adjust colors</i>
---------------	----------------------

---

## Description

Adjust colors

## Usage

```
adjust_colors(
  plot,
  new_colors = NULL,
  saturation = 1,
  labels = tidyplot_parse_labels(),
  downsample = c("evenly", "first", "last", "middle"),
  ...
)
```

## Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
new_colors	A character vector of new hex colors to use. Can be a named character vector of hex colors to assign certain data labels to specific colors.
saturation	A number between 0 and 1 for the color saturation of an object. A value of 0 is completely desaturated (white), 1 is the original color.
labels	One of: <ul style="list-style-type: none"> <li>• NULL for no labels</li> <li>• <code>waiver()</code> for the default labels computed by the transformation object</li> <li>• A character vector giving labels (must be same length as breaks)</li> <li>• An expression vector (must be the same length as breaks). See <code>?plotmath</code> for details.</li> <li>• A function that takes the breaks as input and returns labels as output. Also accepts <code>rlang</code> <a href="#">lambda</a> function notation.</li> </ul>
downsample	If too many colors are provided, whether to downsample evenly, or use the first, the last or the middle colors of the color vector. Defaults to evenly.
...	Arguments passed on to the <code>ggplot2</code> scale function.

## Value

A tidyplot object.

**See Also**

[colors\\_discrete\\_friendly\(\)](#), [colors\\_continuous\\_viridis\(\)](#), [colors\\_diverging\\_blue2brown\(\)](#), and [new\\_color\\_scheme\(\)](#)

**Examples**

```
# Plot without adjustments
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Provide hex colors
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_colors(new_colors = c("#644296", "#F08533", "#3B78B0", "#D1352C"))

# Provide discrete color scheme
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_colors(new_colors = colors_discrete_seaside)

# Provide named vector
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_colors(new_colors = c(
    "A" = "pink",
    "B" = "purple",
    "C" = "grey",
    "D" = "blue"))

# Provide continuous color scheme
climate %>%
  tidyplot(x = month, y = year, color = max_temperature) %>%
  add_heatmap() %>%
  adjust_colors(new_colors = colors_continuous_turbo)
```

**Description**

Adjust font

**Usage**

```
adjust_font(plot, fontsize = 7, family = NULL, face = NULL, color = "black")
```

**Arguments**

plot	A tidyplot generated with the function tidyplot().
fontsize	Font size in points. Defaults to 7.
family	Font family
face	Font face ("plain", "italic", "bold", "bold.italic")
color	A hex color for the stroke color. For example, "#FFFFFF" for white.

**Value**

A tidyplot object.

**Examples**

```
# Plot without adjustments
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Increase font size
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_font(fontsize = 16)

# Change font family
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_font(family = "mono")

# Change font face
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
```



```
adjust_font(face = "bold")
```

---

adjust\_legend\_title *Adjust legend*

---

## Description

Adjust legend

## Usage

```
adjust_legend_title(  
  plot,  
  title = ggplot2::waiver(),  
  fontsize = NULL,  
  family = NULL,  
  face = NULL,  
  color = "black",  
  ...  
)  
  
adjust_legend_position(plot, position = "right")
```

## Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
title	Legend title.
fontsize	Font size in points. Defaults to 7.
family	Font family
face	Font face ("plain", "italic", "bold", "bold.italic")
color	A hex color for the stroke color. For example, "#FFFFFF" for white.
...	Arguments passed on to <code>ggplot2::element_text()</code> .
position	The position of the legend. Can be one of <code>c("right", "left", "bottom", "top", "none")</code> . Defaults to "right".

## Details

- The title argument of `adjust_legend_title()` supports **plotmath expressions** to include special characters. See examples and [Advanced plotting](#).

## Value

A tidyplot object.

**Examples**

```

# Plot without adjustments
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# New title
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_legend_title("My new legend title")

# New title with plotmath expression
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_legend_title("$E=mc^2$")

# Alternative legend positions
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_legend_position("left")

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_legend_position("top")

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_legend_position("bottom")

# `position = "none"` hides the legend
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%

```

```
adjust_legend_position("none")
```

---

adjust_padding	<i>Adjust plot area padding</i>
----------------	---------------------------------

---

### Description

Adjust plot area padding

### Usage

```
adjust_padding(
  plot,
  top = NA,
  right = NA,
  bottom = NA,
  left = NA,
  all = NA,
  force_continuous = FALSE,
  ...
)
```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
top	Extra space between the data points and the top. Defaults to NA, which does not change the padding.
right	Extra space between the data points and the right. Defaults to NA, which does not change the padding.
bottom	Extra space between the data points and the bottom. Defaults to NA, which does not change the padding.
left	Extra space between the data points and the left. Defaults to NA, which does not change the padding.
all	Extra space around the data points. Overwrites top, right, bottom, left if set. Defaults to NA, which does not change the padding.
force_continuous	Whether to force the axis to be continuous. Defaults to FALSE.
...	Arguments passed on to the geom function.

### Value

A tidyplot object.

## Examples

```
# Plot without adjustments
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_padding()

# Increase plot area padding
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_padding(all = 0.2)

animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_padding(top = 0.8)

animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_padding(bottom = 0.8)

animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_padding(right = 0.8)

animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_padding(left = 0.8)
```

---

adjust\_size

*Adjust plot area size*

---

## Description

Adjust plot area size

## Usage

```
adjust_size(plot, width = 50, height = 50, unit = "mm")
```

## Arguments

plot	A tidyplot generated with the function tidyplot().
width	Width of the plot area. Defaults to 50.

height	Height of the plot area. Defaults to 50.
unit	Unit of the plot area width and height. Defaults to mm.

**Value**

A tidyplot object.

**Examples**

```
# Plot without adjustments
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm(shape = 1) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Resize to 20 x 20 mm
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm(shape = 1) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_size(width = 20, height = 20)

# Resize to 4 x 4 cm
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm(shape = 1) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_size(width = 4, height = 4, unit = "cm")

# Remove absolute dimensions and take all available space. This is the ggplot2 default.
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm(shape = 1) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_size(width = NA, height = NA)
```

---

adjust\_theme\_details *Adjust theme details*

---

**Description**

This function is a wrapper around `ggplot2::theme()`. To use the required theme helper functions `ggplot2::element_blank()`, `ggplot2::element_rect()`, `ggplot2::element_line()`, and `ggplot2::element_text()` you need to either load the `ggplot2` package via `library(ggplot2)` or use the `ggplot2::` prefix as shown above.

**Usage**

```
adjust_theme_details(plot, ...)
```

**Arguments**

```
plot          A tidyplot generated with the function tidyplot().  
...          Arguments passed on to the geom function.
```

**Value**

A tidyplot object.

**Examples**

```
study %>%  
  tidyplot(x = treatment, y = score, color = treatment) %>%  
  add_data_points_beeswarm() %>%  
  add_mean_bar(alpha = 0.4) %>%  
  adjust_theme_details(plot.background = ggplot2::element_rect(fill = "#FFEBFF"))
```

---

adjust_title	<i>Adjust titles and caption</i>
--------------	----------------------------------

---

**Description**

Adjust titles and caption

**Usage**

```
adjust_title(  
  plot,  
  title = ggplot2::waiver(),  
  fontsize = NULL,  
  family = NULL,  
  face = NULL,  
  color = "black",  
  ...  
)
```

```
adjust_x_axis_title(  
  plot,  
  title = ggplot2::waiver(),  
  fontsize = NULL,  
  family = NULL,  
  face = NULL,  
  color = "black",
```

```

    ...
  )

adjust_y_axis_title(
  plot,
  title = ggplot2::waiver(),
  fontsize = NULL,
  family = NULL,
  face = NULL,
  color = "black",
  ...
)

adjust_caption(
  plot,
  caption = ggplot2::waiver(),
  fontsize = NULL,
  family = NULL,
  face = NULL,
  color = "black",
  ...
)

```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
title	Plot or axes title.
fontsize	Font size in points. Defaults to 7.
family	Font family
face	Font face ("plain", "italic", "bold", "bold.italic")
color	A hex color for the stroke color. For example, "#FFFFFF" for white.
...	Arguments passed on to <code>ggplot2::element_text()</code> .
caption	Plot caption.

### Details

Adjust the plot title, axis titles and caption

- All functions support **plotmath expressions** to include special characters. See examples and [Advanced plotting](#).

### Value

A tidyplot object.

**Examples**

```

# Plot without adjustments
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Adjust description
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_title("This is my fantastic plot title") %>%
  adjust_x_axis_title("Treatment group") %>%
  adjust_y_axis_title("Disease score") %>%
  adjust_legend_title("Legend title") %>%
  adjust_caption("Here goes the caption")

# Plotmath expressions
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  adjust_title("$H[2]*0$") %>%
  adjust_x_axis_title("$H[2]*0$") %>%
  adjust_y_axis_title("$H[2]*0$") %>%
  adjust_legend_title("$H[2]*0$") %>%
  adjust_caption("$H[2]*0$")

```

---

adjust\_x\_axis

*Adjust axes*


---

**Description**

Adjust axes

**Usage**

```

adjust_x_axis(
  plot,
  title = ggplot2::waiver(),
  breaks = ggplot2::waiver(),
  labels = ggplot2::waiver(),
  limits = NULL,
  padding = c(NA, NA),

```



```

    rotate_labels = FALSE,
    transform = "identity",
    cut_short_scale = FALSE,
    force_continuous = FALSE,
    ...
  )

adjust_y_axis(
  plot,
  title = ggplot2::waiver(),
  breaks = ggplot2::waiver(),
  labels = ggplot2::waiver(),
  limits = NULL,
  padding = c(NA, NA),
  rotate_labels = FALSE,
  transform = "identity",
  cut_short_scale = FALSE,
  force_continuous = FALSE,
  ...
)

```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
title	Axis title.
breaks	One of: <ul style="list-style-type: none"> <li>• NULL for no breaks</li> <li>• <code>waiver()</code> for the default breaks computed by the <a href="#">transformation object</a></li> <li>• A numeric vector of positions</li> <li>• A function that takes the limits as input and returns breaks as output (e.g., a function returned by <code>scales::extended_breaks()</code>). Note that for position scales, limits are provided after scale expansion. Also accepts <a href="#">rlang lambda</a> function notation.</li> </ul>
labels	One of: <ul style="list-style-type: none"> <li>• NULL for no labels</li> <li>• <code>waiver()</code> for the default labels computed by the transformation object</li> <li>• A character vector giving labels (must be same length as breaks)</li> <li>• An expression vector (must be the same length as breaks). See <code>?plotmath</code> for details.</li> <li>• A function that takes the breaks as input and returns labels as output. Also accepts <a href="#">rlang lambda</a> function notation.</li> </ul>
limits	Axis limits. For example, with <code>limits = c(20, 90)</code> the axis starts at 20 and ends at 90.
padding	Extra space between the data points and the axes. Defaults to <code>c(NA, NA)</code> , which does not change the padding.

rotate_labels	Whether to rotate axis labels. If TRUE is set to 45 degrees. You can also provide custom degree values, for example, rotate_labels = 90. Defaults to FALSE.
transform	For continuous scales, the name of a transformation object or the object itself. Built-in transformations include "asn", "atanh", "boxcox", "date", "exp", "hms", "identity", "log", "log10", "log1p", "log2", "logit", "modulus", "probability", "probit", "pseudo_log", "reciprocal", "reverse", "sqrt" and "time". A transformation object bundles together a transform, its inverse, and methods for generating breaks and labels. Transformation objects are defined in the scales package, and are called transform_<name>. If transformations require arguments, you can call them from the scales package, e.g. <code>scales::transform_boxcox(p = 2)</code> . You can create your own transformation with <code>scales::new_transform()</code> .
cut_short_scale	Whether to shorten axis labels using K for thousand, M for million, and so on. Defaults to FALSE.
force_continuous	Whether to force the axis to be continuous. Defaults to FALSE.
...	Arguments passed on to ggplot2 scale function.

### Details

- The title argument of `adjust_x_axis()` and `adjust_y_axis()` supports **plotmath expressions** to include special characters. See examples and **Advanced plotting**.

### Value

A tidyplot object.

### Examples

```
# Plot without adjustments
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points()

# New titles
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_x_axis(title = "My new x-axis title") %>%
  adjust_y_axis(title = "My new y-axis title")

# New titles with plotmath expressions
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_x_axis(title = "$H[2]*O$") %>%
  adjust_y_axis(title = "$E==m*c^{2}$")

# Axes limits
animals %>%
```

```
tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_x_axis(limits = c(-1000, 4000)) %>%
  adjust_y_axis(limits = c(-200, 600))

# Rotate labels
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_x_axis(rotate_labels = 90) %>%
  adjust_y_axis(rotate_labels = 90)

# Increase plot area padding
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_x_axis(padding = c(0.2, 0.2)) %>%
  adjust_y_axis(padding = c(0.2, 0.2))

# Scale transformation
animals %>%
  tidyplot(x = weight, y = size, color = family) %>%
  add_data_points() %>%
  adjust_x_axis(transform = "log10") %>%
  adjust_y_axis(transform = "log2")
```

---

all\_rows

*Subset data rows*

---

## Description

Subset data rows

## Usage

all\_rows()

filter\_rows(..., .by = NULL)

max\_rows(order\_by, n, by = NULL, with\_ties = TRUE, na\_rm = FALSE)

min\_rows(order\_by, n, by = NULL, with\_ties = TRUE, na\_rm = FALSE)

first\_rows(n, by = NULL)

last\_rows(n, by = NULL)

sample\_rows(n, by = NULL)

**Arguments**

...	<a href="#">&lt;data-masking&gt;</a> Expressions that return a logical value, and are defined in terms of the variables in <code>.data</code> . If multiple expressions are included, they are combined with the <code>&amp;</code> operator. Only rows for which all conditions evaluate to <code>TRUE</code> are kept.
<code>.by</code> , <code>by</code>	<b>[Experimental]</b> <a href="#">&lt;tidy-select&gt;</a> Optionally, a selection of columns to group by for just this operation, functioning as an alternative to <code>group_by()</code> . For details and examples, see <code>?dplyr_by</code> .
<code>order_by</code>	<a href="#">&lt;data-masking&gt;</a> Variable or function of variables to order by. To order by multiple variables, wrap them in a data frame or tibble.
<code>n</code>	The number of rows to select. If not are supplied, <code>n = 1</code> will be used. If <code>n</code> is greater than the number of rows in the group, the result will be silently truncated to the group size.  A negative value of <code>n</code> will be subtracted from the group size. For example, <code>n = -2</code> with a group of 5 rows will select $5 - 2 = 3$ rows.
<code>with_ties</code>	Should ties be kept together? The default, <code>TRUE</code> , may return more rows than you request. Use <code>FALSE</code> to ignore ties, and return the first <code>n</code> rows.
<code>na_rm</code>	Should missing values in <code>order_by</code> be removed from the result? If <code>FALSE</code> , <code>NA</code> values are sorted to the end (like in <code>dplyr::arrange()</code> ), so they will only be included if there are insufficient non-missing values to reach <code>n</code> .

**Value**

A function to achieve the desired data subsetting.

**Examples**

```
# Highlight all animals
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points() %>%
  add_data_points(data = all_rows(),
    color = "red", shape = 1, size = 3)

# Highlight 3 animals with the highest weight
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points() %>%
  add_data_points(data = max_rows(weight, n = 3),
    color = "red", shape = 1, size = 3)

# Highlight 3 animals with the lowest weight
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points() %>%
  add_data_points(data = min_rows(weight, n = 3),
    color = "red", shape = 1, size = 3)
```

```
# Highlight the first 3 animals in the dataset
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points() %>%
  add_data_points(data = first_rows(n = 3),
    color = "red", shape = 1, size = 3)

# Highlight the last 3 animals in the dataset
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points() %>%
  add_data_points(data = last_rows(n = 3),
    color = "red", shape = 1, size = 3)

# Highlight 3 random animals
animals %>%
  tidyplot(x = weight, y = size) %>%
  add_data_points() %>%
  add_data_points(data = sample_rows(n = 3),
    color = "red", shape = 1, size = 3)
```

---

animals

*Animals data*

---

## Description

Animals data

## Usage

```
animals
```

## Format

A data frame.

## Source

ChatGPT-3.5, *Caution: The accuracy of the data has not been verified.*

## Examples

```
dplyr::glimpse(animals)
```

---

as_tidyplot	<i>Convert ggplot to tidyplot</i>
-------------	-----------------------------------

---

**Description**

Convert ggplot to tidyplot

**Usage**

```
as_tidyplot(gg, width = 50, height = 50, dodge_width = NULL)
```

**Arguments**

gg	A ggplot.
width	Width of the plot area. Defaults to 50.
height	Height of the plot area. Defaults to 50.
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.

**Value**

A tidyplot object.

**Examples**

```
gg <-  
  study %>%  
  ggplot2::ggplot(ggplot2::aes(x = treatment, y = score, color = treatment)) +  
  ggplot2::geom_point()  
  
gg  
  
gg %>% as_tidyplot()
```

---

climate	<i>Climate data</i>
---------	---------------------

---

**Description**

Climate data

**Usage**

```
climate
```

**Format**

A data frame.

**Source**

[National Oceanic and Atmospheric Administration](#), Temperature data, weather station Hamburg Fuhlsbüttel, Germany

**Examples**

```
dplyr::glimpse(climate)
```

---

```
colors_continuous_viridis
```

*Continuous color schemes*

---

**Description**

For more information about the use of color schemes in tidypLOTS, check out this article: [Color schemes](#)

**Usage**

```
colors_continuous_viridis  
colors_continuous_magma  
colors_continuous_inferno  
colors_continuous_plasma  
colors_continuous_cividis  
colors_continuous_rocket  
colors_continuous_mako  
colors_continuous_turbo  
colors_continuous_bluepinkyellow
```

**Format**

An object of class `tidycolor` (inherits from `character`) of length 265.  
An object of class `tidycolor` (inherits from `character`) of length 265.  
An object of class `tidycolor` (inherits from `character`) of length 265.

An object of class tidycolor (inherits from character) of length 265.  
An object of class tidycolor (inherits from character) of length 265.  
An object of class tidycolor (inherits from character) of length 265.  
An object of class tidycolor (inherits from character) of length 265.  
An object of class tidycolor (inherits from character) of length 265.  
An object of class tidycolor (inherits from character) of length 11.

### Details

Color schemes can be conveniently previewed by using the print method of the tidycolor class. This will send a html preview to the RStudio Viewer pane.

```
colors_continuous_viridis  
  
colors_continuous_magma  
  
colors_continuous_inferno  
  
colors_continuous_plasma  
  
colors_continuous_cividis  
  
colors_continuous_rocket  
  
colors_continuous_mako  
  
colors_continuous_turbo  
  
colors_continuous_bluepinkyellow
```

---

```
colors_discrete_friendly
```

*Discrete color schemes*

---

### Description

For more information about the use of color schemes in tidyplots, check out this article: [Color schemes](#)



### Usage

colors\_discrete\_friendly  
colors\_discrete\_seaside  
colors\_discrete\_apple  
colors\_discrete\_friendly\_long  
colors\_discrete\_okabeito  
colors\_discrete\_ibm  
colors\_discrete\_metro  
colors\_discrete\_candy  
colors\_discrete\_alger  
colors\_discrete\_rainbow

### Format

An object of class tidycolor (inherits from character) of length 6.  
An object of class tidycolor (inherits from character) of length 5.  
An object of class tidycolor (inherits from character) of length 7.  
An object of class tidycolor (inherits from character) of length 7.  
An object of class tidycolor (inherits from character) of length 7.  
An object of class tidycolor (inherits from character) of length 5.  
An object of class tidycolor (inherits from character) of length 5.  
An object of class tidycolor (inherits from character) of length 5.  
An object of class tidycolor (inherits from character) of length 5.  
An object of class tidycolor (inherits from character) of length 9.

### Details

The signature theme of tidypplots colors\_discrete\_friendly was adapted from the [Okabe & Ito](#) color palette that was designed to work well for people with color vision deficiency.

Color schemes can be conveniently previewed by using the print method of the tidycolor class. This will send a html preview to the RStudio Viewer pane.

colors\_discrete\_friendly  
colors\_discrete\_seaside

colors\_discrete\_apple  
colors\_discrete\_friendly\_long  
colors\_discrete\_okabeito  
colors\_discrete\_ibm  
colors\_discrete\_metro  
colors\_discrete\_candy  
colors\_discrete\_alger  
colors\_discrete\_rainbow

---

colors\_diverging\_blue2red  
*Diverging color schemes*

---

### **Description**

For more information about the use of color schemes in tidypLOTS, check out this article: [Color schemes](#)

### **Usage**

colors\_diverging\_blue2red  
colors\_diverging\_blue2brown  
colors\_diverging\_BuRd  
colors\_diverging\_BuYlRd  
colors\_diverging\_spectral  
colors\_diverging\_icefire

### **Format**

An object of class tidycolor (inherits from character) of length 17.  
An object of class tidycolor (inherits from character) of length 17.  
An object of class tidycolor (inherits from character) of length 11.  
An object of class tidycolor (inherits from character) of length 11.  
An object of class tidycolor (inherits from character) of length 96.  
An object of class tidycolor (inherits from character) of length 96.

**Details**

Color schemes can be conveniently previewed by using the `print` method of the `tidycolor` class. This will send a html preview to the RStudio Viewer pane.

```
colors_diverging_blue2red
```

```
colors_diverging_blue2brown
```

```
colors_diverging_BuRd
```

```
colors_diverging_BuYlRd
```

```
colors_diverging_spectral
```

```
colors_diverging_icefire
```

---

dinosaurus

*Dinosaurs data*

---

**Description**

Dinosaurs data

**Usage**

```
dinosaurus
```

**Format**

A data frame.

**Source**

ChatGPT-3.5, *Caution: The accuracy of the data has not been verified.*

**Examples**

```
dplyr::glimpse(dinosaurus)
```

---

distributions	<i>Distributions data</i>
---------------	---------------------------

---

**Description**

Distributions data

**Usage**

```
distributions
```

**Format**

A data frame.

**Source**

tidyplots package

**Examples**

```
dplyr::glimpse(distributions)
```

---

energy	<i>Energy data</i>
--------	--------------------

---

**Description**

Energy data

**Usage**

```
energy
```

**Format**

A data frame.

**Source**

[Energy-Charts](#), Energy production data, Germany

**Examples**

```
dplyr::glimpse(energy)
```

---

energy_week	<i>Energy week data</i>
-------------	-------------------------

---

**Description**

Energy week data

**Usage**

```
energy_week
```

**Format**

A data frame.

**Source**

[Energy-Charts](#), Energy production data, Germany

**Examples**

```
dplyr::glimpse(energy_week)
```

---

eu_countries	<i>EU countries data</i>
--------------	--------------------------

---

**Description**

EU countries data

**Usage**

```
eu_countries
```

**Format**

A data frame.

**Source**

ChatGPT-3.5, *Caution: The accuracy of the data has not been verified.*

**Examples**

```
dplyr::glimpse(eu_countries)
```

flip\_plot

*Flip x and y-axis***Description****[Superseded]**

This function is superseded because in many cases, `flip_plot()` can easily be replaced by swapping the x and y axis. Some plot components additionally require to set the orientation argument to "y".

**Usage**

```
flip_plot(plot, ...)
```

**Arguments**

<code>plot</code>	A tidyplot generated with the function <code>tidyplot()</code> .
<code>...</code>	Arguments passed on to <code>ggplot2::coord_flip()</code> .

**Value**

A tidyplot object.

**Examples**

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  flip_plot()

energy %>%
  tidyplot(x = year, y = energy, color = energy_type) %>%
  add_barstack_absolute() %>%
  flip_plot()

# Better solutions without `flip_plot()`
study %>%
  tidyplot(x = score, y = treatment, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

energy %>%
  tidyplot(x = energy, y = year, color = energy_type) %>%
  add_barstack_absolute(orientation = "y")
```

---

format_number	<i>Format numbers or p values</i>
---------------	-----------------------------------

---

**Description**

Format numbers or p values

**Usage**

```
format_number(x, accuracy = 0.1, big.mark = ",", scale_cut = NULL, ...)
```

```
format_p_value(x, accuracy = 1e-04)
```

**Arguments**

x	A number to format.
accuracy	A number to round to. Use (e.g.) 0.01 to show 2 decimal places of precision. If NULL, the default, uses a heuristic that should ensure breaks have the minimum number of digits needed to show the difference between adjacent values. Applied to rescaled data.
big.mark	Character used between every 3 digits to separate thousands.
scale_cut	Named numeric vector that allows you to rescale large (or small) numbers and add a prefix. Built-in helpers include: <ul style="list-style-type: none"> <li>cut_short_scale(): [10<sup>3</sup>, 10<sup>6</sup>) = K, [10<sup>6</sup>, 10<sup>9</sup>) = M, [10<sup>9</sup>, 10<sup>12</sup>) = B, [10<sup>12</sup>, Inf) = T.</li> <li>cut_long_scale(): [10<sup>3</sup>, 10<sup>6</sup>) = K, [10<sup>6</sup>, 10<sup>12</sup>) = M, [10<sup>12</sup>, 10<sup>18</sup>) = B, [10<sup>18</sup>, Inf) = T.</li> <li>cut_si(unit): uses standard SI units.</li> </ul> <p>If you supply a vector c(a = 100, b = 1000), absolute values in the range [0, 100) will not be rescaled, absolute values in the range [100, 1000) will be divided by 100 and given the suffix "a", and absolute values in the range [1000, Inf) will be divided by 1000 and given the suffix "b". If the division creates an irrational value (or one with many digits), the cut value below will be tried to see if it improves the look of the final label.</p>
...	Arguments passed on to <code>scales::number</code>
scale	A scaling factor: x will be multiplied by scale before formatting. This is useful if the underlying data is very small or very large.
style_positive	A string that determines the style of positive numbers: <ul style="list-style-type: none"> <li>"none" (the default): no change, e.g. 1.</li> <li>"plus": preceded by +, e.g. +1.</li> <li>"space": preceded by a Unicode "figure space", i.e., a space equally as wide as a number or +. Compared to "none", adding a figure space can ensure numbers remain properly aligned when they are left- or right-justified.</li> </ul>

`style_negative` A string that determines the style of negative numbers:

- "hyphen" (the default): preceded by a standard hyphen -, e.g. -1.
- "minus", uses a proper Unicode minus symbol. This is a typographical nicety that ensures - aligns with the horizontal bar of the the horizontal bar of +.
- "parens", wrapped in parentheses, e.g. (1).

### Value

Formatted number as character string.

### Examples

```
format_number(232342.3443)
```

```
format_number(232342.3443, accuracy = 0.01)
```

```
format_number(232342.3443, accuracy = 1, big.mark = "")
```

```
format_p_value(0.03445553)
```

```
format_p_value(0.0003445553)
```

```
format_p_value(0.00003445553)
```

---

gene_expression	<i>RNA-Seq expression data</i>
-----------------	--------------------------------

---

### Description

RNA-Seq expression data

### Usage

```
gene_expression
```

### Format

A data frame.

### Source

[Bassoon proteinopathy drives neurodegeneration in multiple sclerosis](#), Nature Neuroscience  
[GSE104899](#), Gene Expression Omnibus

### Examples

```
dplyr::glimpse(gene_expression)
```



---

new_color_scheme	<i>New color scheme</i>
------------------	-------------------------

---

### Description

For more information about the use of color schemes in tidyplots, check out this article: [Color schemes](#)

### Usage

```
new_color_scheme(x, name = "Untitled color scheme", reverse = FALSE)
```

### Arguments

x	Character vector of hex colors. For example <code>x = c("#FF00FF", "#00FFFF")</code> .
name	Name of the custom color scheme.
reverse	Whether the order should be reversed or not. Defaults to FALSE, meaning not reversed.

### Value

A tidyplot object.

### Examples

```
new_color_scheme(c("#ECA669", "#E06681", "#8087E2", "#E2D269"))

new_color_scheme(c("#ECA669", "#E06681", "#8087E2", "#E2D269"),
  name = "my_custom_color_scheme")
```

---

remove_legend	<i>Remove legend or legend title</i>
---------------	--------------------------------------

---

### Description

Remove legend or legend title

### Usage

```
remove_legend(plot)

remove_legend_title(plot)
```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
------	--

**Value**

A tidyplot object.

**Examples**

```
# Before removing
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar()

# After removing
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_legend_title()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_legend()
```

---

remove_padding	<i>Remove plot area padding</i>
----------------	---------------------------------

---

**Description**

Remove plot area padding

**Usage**

```
remove_padding(plot, force_continuous = FALSE)
```

**Arguments**

**plot** A tidyplot generated with the function tidyplot().

**force\_continuous** Whether to force the axis to be continuous. Defaults to FALSE.

**Value**

A tidyplot object.

**Examples**

```
# Before removing
animals %>%
  tidyplot(x = weight, y = speed, color = family) %>%
  add_data_points()

# After removing
animals %>%
  tidyplot(x = weight, y = speed, color = family) %>%
  add_data_points() %>%
  remove_padding()
```

---

remove_title	<i>Remove plot title or caption</i>
--------------	-------------------------------------

---

**Description**

Remove plot title or caption

**Usage**

```
remove_title(plot)

remove_caption(plot)
```

**Arguments**

plot            A tidyplot generated with the function tidyplot().

**Value**

A tidyplot object.

**Examples**

```
# Before removing
animals %>%
  tidyplot(x = weight, y = speed, color = family) %>%
  add_data_points() %>%
  add_title("Name of the plot") %>%
  add_caption("This is the caption")

# After removing
animals %>%
  tidyplot(x = weight, y = speed, color = family) %>%
  add_data_points() %>%
  add_title("Name of the plot") %>%
  add_caption("This is the caption") %>%
```

```
remove_title()

animals %>%
  tidyplot(x = weight, y = speed, color = family) %>%
  add_data_points() %>%
  add_title("Name of the plot") %>%
  add_caption("This is the caption") %>%
  remove_caption()
```

---

remove\_x\_axis

*Remove x-axis or parts of it*

---

## Description

Remove x-axis or parts of it

## Usage

```
remove_x_axis(plot)

remove_x_axis_line(plot)

remove_x_axis_ticks(plot)

remove_x_axis_labels(plot)

remove_x_axis_title(plot)
```

## Arguments

plot            A tidyplot generated with the function tidyplot().

## Value

A tidyplot object.

## Examples

```
# Before removing
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar()

# After removing
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_x_axis_line()
```

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_x_axis_ticks()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_x_axis_labels()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_x_axis_title()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_x_axis()
```

---

remove_y_axis	<i>Remove y-axis or parts of it</i>
---------------	-------------------------------------

---

## Description

Remove y-axis or parts of it

## Usage

```
remove_y_axis(plot)

remove_y_axis_line(plot)

remove_y_axis_ticks(plot)

remove_y_axis_labels(plot)

remove_y_axis_title(plot)
```

## Arguments

plot            A tidyplot generated with the function tidyplot().

## Value

A tidyplot object.

## Examples

```
# Before removing
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar()

# After removing
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_y_axis_line()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_y_axis_ticks()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_y_axis_labels()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_y_axis_title()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar() %>%
  remove_y_axis()
```

---

rename\_x\_axis\_labels *Rename axis or color labels*

---

## Description

Rename axis or color labels

## Usage

```
rename_x_axis_labels(plot, new_names)
```

```
rename_y_axis_labels(plot, new_names)
```

```
rename_color_labels(plot, new_names)
```

**Arguments**

`plot` A tidyplot generated with the function `tidyplot()`.  
`new_names` Named character vector in the format `c("old1" = "new1", "old2" = "new2")`.

**Value**

A tidyplot object.

**Examples**

```
# Before adjustments
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Rename x-axis labels
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  rename_x_axis_labels(new_names = c(
    "A" = "This",
    "B" = "is",
    "C" = "totally",
    "D" = "new"))

# Before adjustments
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Rename y-axis labels
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  rename_y_axis_labels(new_names = c(
    "A" = "This",
    "B" = "is",
    "C" = "totally",
    "D" = "new"))

# Before adjustment
study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_data_points() %>%
```

```
add_mean_bar(alpha = 0.4) %>%
add_sem_errorbar()

# Rename color labels
study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  rename_color_labels(new_names = c(
    "high" = "Sky high",
    "low" = "Deep low"))
```

---

reorder\_x\_axis\_labels *Reorder axis or color labels*

---

## Description

Reorder axis or color labels

## Usage

```
reorder_x_axis_labels(plot, ...)

reorder_y_axis_labels(plot, ...)

reorder_color_labels(plot, ...)
```

## Arguments

plot	A tidyplot generated with the function tidyplot().
...	Arguments passed on to forcats::fct_relevel().

## Value

A tidyplot object.

## Examples

```
# Before adjustments
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Reorder x-axis labels
study %>%
```



```
tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  reorder_x_axis_labels("D", "B", "A")

# Before adjustments
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Reorder y-axis labels
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  reorder_y_axis_labels("D", "B", "A")

# Before adjustment
study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Reorder color labels
study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  reorder_color_labels("low")
```

---

reverse\_x\_axis\_labels *Reverse axis or color labels*

---

## Description

Reverse axis or color labels

## Usage

```
reverse_x_axis_labels(plot)
```

```
reverse_y_axis_labels(plot)
```

```
reverse_color_labels(plot)
```

**Arguments**

plot                    A tidyplot generated with the function tidyplot().

**Value**

A tidyplot object.

**Examples**

```
# Before adjustments
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Reverse x-axis labels
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  reverse_x_axis_labels()

# Before adjustments
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Reverse y-axis labels
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  reverse_y_axis_labels()

# Before adjustment
study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Reverse color labels
study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
```

```
reverse_color_labels()
```

---

save_plot	<i>Save plots to file</i>
-----------	---------------------------

---

### Description

This function takes a plot or list of plots and writes them to a (multipage) file.

### Usage

```
save_plot(
  plot = ggplot2::last_plot(),
  filename,
  width = NA,
  height = NA,
  units = c("mm", "cm", "in"),
  multiple_files = FALSE,
  view_plot = TRUE,
  bg = "transparent",
  ...
)
```

### Arguments

plot	Plot to save, defaults to last plot displayed.
filename	File name to create on disk.
width, height	Dimensions of the graphic device to save the plot. Defaults to NA. In case of NA, the dimensions are inferred from the incoming plot object (see Details).
units	Units of length. Defaults to "mm".
multiple_files	Whether to save multiple pages as individual files.
view_plot	Whether to view the plot on screen after saving.
bg	Background colour. If NULL, uses the plot.background fill value from the plot theme.
...	Other arguments passed on to the graphics device function, as specified by device.

### Details

**Handling of file dimensions.** Output file dimensions are determined according to the following precedence.

1. The width and height arguments.
2. Dimensions inferred from the incoming plot object with absolute dimensions.
3. System default device dimensions.

**Value**

A tidyplot object.

**Examples**

```
# Save plot to file
study %>%
  tidyplot(treatment, score) %>%
  add_data_points() %>%
  save_plot("single_plot.pdf")

# Save intermediate stages to file
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  add_data_points_beeswarm() %>%
  save_plot("before.pdf") %>%
  adjust_colors(colors_discrete_seaside) %>%
  save_plot("after.pdf")

# Save multipage PDF file
gene_expression %>%
  .[1:160,] %>%
  tidyplot(group, expression, color = sample_type) %>%
  add_data_points() %>%
  split_plot(by = external_gene_name, nrow = 2, ncol = 2) %>%
  save_plot("multipage_plot.pdf")

# Save multiple PDF files
gene_expression %>%
  .[1:160,] %>%
  tidyplot(group, expression, color = sample_type) %>%
  add_data_points() %>%
  split_plot(by = external_gene_name, nrow = 2, ncol = 2) %>%
  save_plot("plot.pdf", multiple_files = TRUE)
```

---

sort\_x\_axis\_labels      *Sort axis or color labels*

---

**Description**

Sort axis or color labels

**Usage**

```
sort_x_axis_labels(plot, ..., .fun = NULL, .reverse = FALSE)
```

```
sort_y_axis_labels(plot, ..., .fun = NULL, .reverse = FALSE)
```

```
sort_color_labels(plot, ..., .fun = NULL, .reverse = FALSE)
```

**Arguments**

<code>plot</code>	A tidyplot generated with the function <code>tidyplot()</code> .
<code>...</code>	Optional variables to use for sorting.
<code>.fun</code>	Override the function used for sorting. Is automatically determined from the plot.
<code>.reverse</code>	Whether the order should be reversed or not. Defaults to <code>FALSE</code> , meaning not reversed.

**Value**

A tidyplot object.

**Examples**

```
# Before adjustments
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Sort x-axis labels by score
study %>%
  tidyplot(x = treatment, y = score) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  sort_x_axis_labels()

# Before adjustments
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Sort y-axis labels by score
study %>%
  tidyplot(x = score, y = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
```

```
sort_y_axis_labels()

# Before adjustment
study %>%
  tidyplot(x = group, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar()

# Sort color labels by score
study %>%
  tidyplot(x = group, y = score, color = treatment) %>%
  add_data_points() %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  sort_color_labels()
```

---

spendings

*Spending data*

---

### **Description**

Spending data

### **Usage**

spendings

### **Format**

A data frame.

### **Source**

tidyplots package

### **Examples**

```
dplyr::glimpse(spendings)
```

---

split_plot	<i>Split plot into multiple subplots</i>
------------	--

---

### Description

Split plot into multiple subplots

### Usage

```
split_plot(
  plot,
  by,
  ncol = NULL,
  nrow = NULL,
  byrow = NULL,
  widths = 30,
  heights = 25,
  guides = "collect",
  tag_level = NULL,
  design = NULL,
  unit = "mm"
)
```

### Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
by	Variable that should be used for splitting.
ncol, nrow	The number of columns and rows per page.
byrow	Analogous to <code>byrow</code> in <code>matrix()</code> . If <code>FALSE</code> the plots will be filled in in column-major order
widths, heights	The relative widths and heights of each column and row in the grid. Will get repeated to match the dimensions of the grid. The special value of <code>NA/1null</code> will behave as <code>1null</code> unless a fixed aspect plot is inserted in which case it will allow the dimension to expand or contract to match the aspect ratio of the content
guides	A string specifying how guides should be treated in the layout. <code>'collect'</code> will collect guides below to the given nesting level, removing duplicates. <code>'keep'</code> will stop collection at this level and let guides be placed alongside their plot. <code>auto</code> will allow guides to be collected if a upper level tries, but place them alongside the plot if not. If you modify default guide <code>"position"</code> with <code>theme(legend.position=...)</code> while also collecting guides you must apply that change to the overall patchwork (see example).
tag_level	A string ( <code>'keep'</code> or <code>'new'</code> ) to indicate how auto-tagging should behave. See <code>plot_annotation()</code> .
design	Specification of the location of areas in the layout. Can either be specified as a text string or by concatenating calls to <code>area()</code> together. See the examples for further information on use.

unit                    Unit of length. Defaults to "mm".

### Value

A tidypilot object.

### Examples

```
# Before splitting
energy %>%
  dplyr::filter(year %in% c(2005, 2010, 2015, 2020)) %>%
  tidypilot(y = energy, color = energy_source) %>%
  add_donut()

# Split by year
energy %>%
  dplyr::filter(year %in% c(2005, 2010, 2015, 2020)) %>%
  tidypilot(y = energy, color = energy_source) %>%
  add_donut() %>%
  split_plot(by = year)

# Change dimensions of subplots
energy %>%
  dplyr::filter(year %in% c(2005, 2010, 2015, 2020)) %>%
  tidypilot(y = energy, color = energy_source) %>%
  add_donut() %>%
  split_plot(by = year, widths = 15, heights = 15)

# Spread plots across multiple pages
energy %>%
  dplyr::filter(year %in% c(2005, 2010, 2015, 2020)) %>%
  tidypilot(y = energy, color = energy_source) %>%
  add_donut() %>%
  split_plot(by = year, ncol = 2, nrow = 1)
```

---

study

*Study data*

---

### Description

Study data

### Usage

```
study
```

### Format

A data frame.



**Source**

tidyplots package

**Examples**

```
dplyr::glimpse(study)
```

---

theme_tidyplot	<i>Themes</i>
----------------	---------------

---

**Description**

Themes

**Usage**

```
theme_tidyplot(plot, fontsize = 7)
theme_ggplot2(plot, fontsize = 7)
theme_minimal_xy(plot, fontsize = 7)
theme_minimal_x(plot, fontsize = 7)
theme_minimal_y(plot, fontsize = 7)
```

**Arguments**

plot	A tidyplot generated with the function tidyplot().
fontsize	Font size in points. Defaults to 7.

**Value**

A tidyplot object.

**Examples**

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar() %>%
  add_mean_dash() %>%
  theme_tidyplot()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar() %>%
```

```

add_mean_dash() %>%
theme_ggplot2()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar() %>%
  add_mean_dash() %>%
  theme_minimal_xy()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar() %>%
  add_mean_dash() %>%
  theme_minimal_x()

study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points() %>%
  add_sem_errorbar() %>%
  add_mean_dash() %>%
  theme_minimal_y()

```

---

tidyplot

*Create a new tidyplot*


---

## Description

Create a new tidyplot

## Usage

```
tidyplot(data, ..., width = 50, height = 50, dodge_width = NULL)
```

## Arguments

data	A tidy data frame to use for plotting.
...	Mappings for the x axis, y axis and color, see examples. Additional argument are passed to <code>ggplot2::aes()</code> .
width	Width of the plot area. Defaults to 50.
height	Height of the plot area. Defaults to 50.
dodge_width	For adjusting the distance between grouped objects. Defaults to 0.8 for plots with at least one discrete axis and 0 for plots with two continuous axes.

## Value

A tidyplot object.

**Examples**

```
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_data_points_beeswarm()

study %>%
  tidyplot(x = group, y = score, color = dose) %>%
  add_mean_bar()

# Change plot area size
study %>%
  tidyplot(x = treatment, y = score, color = treatment,
           width = 35, height = 35) %>%
  add_data_points_beeswarm()

# Change dodge_width
study %>%
  tidyplot(x = group, y = score, color = dose, dodge_width = 0.3) %>%
  add_mean_bar()
```

---

time\_course

*Time course data*

---

**Description**

Time course data

**Usage**

```
time_course
```

**Format**

A data frame.

**Source**

tidyplots package

**Examples**

```
dplyr::glimpse(time_course)
```

---

 view\_plot

*View plot on screen*


---

## Description

View plot on screen

## Usage

```
view_plot(plot, data = all_rows(), title = ggplot2::waiver(), ...)
```

## Arguments

plot	A tidyplot generated with the function <code>tidyplot()</code> .
data	The data to be displayed in this layer. There are three options: <ul style="list-style-type: none"> <li>• If <code>all_rows()</code> (the default) the complete dataset is displayed.</li> <li>• A function to subset the plot data. See <code>filter_rows()</code> and friends.</li> <li>• A <code>data.frame</code> to override the plot data.</li> </ul>
title	Plot title.
...	Arguments passed on to <code>print()</code> .

## Details

- `view_plot()` supports data subsetting. See examples and [Advanced plotting](#).

## Value

A tidyplot object.

## Examples

```
# View intermediate stages on screen
study %>%
  tidyplot(x = treatment, y = score, color = treatment) %>%
  add_mean_bar(alpha = 0.4) %>%
  add_sem_errorbar() %>%
  add_data_points_beeswarm() %>%
  view_plot(title = "Before changing color scheme") %>%
  adjust_colors(colors_discrete_seaside) %>%
  view_plot(title = "After changing color scheme")

# View data subsets on screen
gene_expression %>%
  tidyplot(x = condition, y = expression, color = sample_type) %>%
  add_mean_dash() %>%
  add_sem_errorbar() %>%
  add_data_points_beeswarm() %>%
```

```
view_plot(data = filter_rows(external_gene_name == "Apol6"),  
          title = "Apol6") %>%  
view_plot(data = filter_rows(external_gene_name == "Bsn"),  
          title = "Bsn")
```

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