# Package 'bayesplot'

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```
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      easily used by developers working on a variety of R packages for Bayesian
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```

Type Package

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# **Description**

The **bayesplot** package provides a variety of **ggplot2**-based plotting functions for use after fitting Bayesian models (typically, though not exclusively, via Markov chain Monte Carlo). The package is designed not only to provide convenient functionality for users, but also a common set of functions that can be easily used by developers working on a variety of packages for Bayesian modeling, particularly (but not necessarily) packages powered by **rstan**. Examples of packages that will soon (or already are) using **bayesplot** are **rstan** itself, as well as the **rstan**-dependent **rstanarm** and **brms** packages for applied regression modeling.

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## **Plotting functionality**

The plotting functions in **bayesplot** are organized into several modules:

• MCMC: Visualizations of Markov chain Monte Carlo (MCMC) simulations generated by *any* MCMC algorithm as well as diagnostics. There are also additional functions specifically for use with models fit using the No-U-Turn Sampler (NUTS).

• PPC: Graphical posterior predictive checks (PPCs).

In future releases modules will be added specifically for forecasting/out-of-sample prediction and other inference-related tasks.

#### Resources

• **Bug reports and feature requests**: If you would like to request a new feature or if you have noticed a bug that needs to be fixed please let us know at the **bayesplot** issue tracker on GitHub.

```
https://github.com/stan-dev/bayesplot/issues/.
```

 General questions and help: To ask a question about bayesplot on the Stan Forums forum please visit:

```
http://discourse.mc-stan.org.
```

#### References

Gabry, J., Simpson, D., Vehtari, A., Betancourt, M., and Gelman, A. (2018). Visualization in Bayesian workflow. *Journal of the Royal Statistical Society Series A*, accepted for publication. arXiv preprint: http://arxiv.org/abs/1709.01449.

## See Also

```
theme_default for the default ggplot theme used by bayesplot. bayesplot-colors to set or view the color scheme used for plotting. ggsave in ggplot2 for saving plots.
```

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available\_ppc

Get or view the names of available plotting functions

## **Description**

Get or view the names of available plotting functions

#### Usage

```
available_ppc(pattern, fixed = FALSE, invert = FALSE)
available_mcmc(pattern, fixed = FALSE, invert = FALSE)
```

## **Arguments**

```
pattern, fixed, invert
Passed to grep.
```

## Value

A possibly empty character vector of function names with several additional attributes (for use by a custom print method). If pattern is missing then the returned object contains the names of all available plotting functions in the MCMC or PPC module, depending on which function is called. If pattern is specified then a subset of function names is returned.

```
available_mcmc()
available_mcmc("nuts")
available_mcmc("rhat|neff")
available_ppc("grouped")
available_ppc("grouped", invert = TRUE)
```

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bayesplot-colors

Set, get, or view color schemes

#### **Description**

Set, get, or view color schemes. Choose from a preset scheme or create a custom scheme.

#### Usage

```
color_scheme_set(scheme = "blue")
color_scheme_get(scheme, i)
color_scheme_view(scheme)
```

#### **Arguments**

scheme

For color\_scheme\_set, either a string naming one of the available color schemes or a character vector of *exactly six* colors specifying a custom scheme (see the **Custom Color Schemes** section, below, for more on specifying a custom scheme).

For color\_scheme\_get, scheme can be missing (to get the current color scheme) or a string naming one of the preset schemes.

For color\_scheme\_view, scheme can be missing (to use the current color scheme) or a character vector containing a subset of the available scheme names.

Currently, the available preset color schemes are:

- "blue", "brightblue"
- "gray", "darkgray"
- "green"
- "pink"
- "purple"
- "red"
- "teal"
- "vellow"
- "viridis", "viridisA", "viridisB", "viridisC"
- "mix-x-y", replacing x and y with any two of the scheme names listed above (e.g. "mix-teal-pink", "mix-blue-red", etc.). The order of x and y matters, i.e., the color schemes "mix-blue-red" and "mix-red-blue" are not identical. There is no guarantee that every possible mixed scheme will look good with every possible plot.

If you have a suggestion for a new color scheme please let us know via the **bayesplot** issue tracker.

For color\_scheme\_get, a subset of the integers from 1 (lightest) to 6 (darkest) indicating which of the colors in the scheme to return. If i is not specified then all six colors in the scheme are included.

i

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## Value

color\_scheme\_set has the side effect of setting the color scheme used for plotting. It also returns (invisibly) a list of the hexidecimal color values used in scheme.

color\_scheme\_get returns a list of the hexadecimal color values (without changing the current scheme). If the scheme argument is not specified the returned values correspond to the current color scheme. If the optional argument i is specified then the returned list only contains length(i) elements.

color\_scheme\_view returns a ggplot object if only a single scheme is specified and a gtable object if multiple schemes names are specified.

#### **Custom Color Schemes**

A **bayesplot** color scheme consists of six colors. To specify a custom color scheme simply pass a character vector containing either the names of six colors or six hexidecimal color values (or a mix of names and hex values). The colors should be in order from lightest to darkest. See the end of the **Examples** section for a demonstration.

#### See Also

theme\_default for the default ggplot theme used by **bayesplot**.

```
color_scheme_set("blue")
color_scheme_view()
color_scheme_get()
color\_scheme\_get(i = c(3, 5)) # 3rd and 5th colors only
color_scheme_get("brightblue")
color_scheme_view("brightblue")
# compare multiple schemes
color_scheme_view(c("pink", "gray", "teal"))
color_scheme_view(c("viridis", "viridisA", "viridisB", "viridisC"))
color_scheme_set("pink")
x <- example_mcmc_draws()</pre>
mcmc_intervals(x)
color_scheme_set("teal")
color_scheme_view()
mcmc_intervals(x)
color_scheme_set("red")
mcmc_areas(x, regex_pars = "beta")
color_scheme_set("purple")
color_scheme_view()
y <- example_y_data()</pre>
```

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bayesplot-extractors Extract quantities needed for plotting from model objects

# **Description**

Generics and methods for extracting quantities needed for plotting from various types of model objects. Currently methods are only provided for stanfit (**rstan**) and stanreg (**rstanarm**) objects, but adding new methods should be relatively straightforward.

## Usage

```
log_posterior(object, ...)
nuts_params(object, ...)
rhat(object, ...)
neff_ratio(object, ...)
## S3 method for class 'stanfit'
log_posterior(object, inc_warmup = FALSE, ...)
## S3 method for class 'stanreg'
log_posterior(object, inc_warmup = FALSE, ...)
## S3 method for class 'stanfit'
nuts_params(object, pars = NULL, inc_warmup = FALSE, ...)
## S3 method for class 'stanreg'
```

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```
nuts_params(object, pars = NULL, inc_warmup = FALSE, ...)
## S3 method for class 'list'
nuts_params(object, pars = NULL, ...)
## S3 method for class 'stanfit'
rhat(object, pars = NULL, ...)
## S3 method for class 'stanreg'
rhat(object, pars = NULL, regex_pars = NULL, ...)
## S3 method for class 'stanfit'
neff_ratio(object, pars = NULL, ...)
## S3 method for class 'stanreg'
neff_ratio(object, pars = NULL, regex_pars = NULL, ...)
```

#### **Arguments**

object The object to use.

... Arguments passed to individual methods.

inc\_warmup A logical scalar (defaulting to FALSE) indicating whether to include warmup

draws, if applicable.

pars An optional character vector of parameter names. For nuts\_params these will

be NUTS sampler parameter names rather than model parameters. If pars is

omitted all parameters are included.

regex\_pars An optional regular expression to use for parameter selection. Can be specified

instead of pars or in addition to pars.

#### Value

log\_posterior log\_posterior methods return a molten data frame (see melt). The data frame should have columns "Iteration" (integer), "Chain" (integer), and "Value" (numeric). See **Examples**, below.

nuts\_params nuts\_params methods return a molten data frame (see melt). The data frame should have columns "Parameter" (factor), "Iteration" (integer), "Chain" (integer), and "Value" (numeric). See **Examples**, below.

rhat, neff\_ratio Methods return (named) vectors.

#### See Also

```
MCMC-nuts, MCMC-diagnostics
```

```
## Not run:
library(rstanarm)
fit <- stan_glm(mpg ~ wt, data = mtcars)</pre>
```

```
np <- nuts_params(fit)
head(np)
tail(np)

lp <- log_posterior(fit)
head(lp)
tail(lp)
## End(Not run)</pre>
```

bayesplot-helpers

Convenience functions for adding or changing plot details

# Description

Convenience functions for adding to (and changing details of) ggplot objects (many of the objects returned by **bayesplot** functions). See the **Examples** section, below.

# Usage

```
vline_at(v, fun, ..., na.rm = TRUE)
hline_at(v, fun, ..., na.rm = TRUE)
vline_0(..., na.rm = TRUE)
hline_0(..., na.rm = TRUE)
abline_01(..., na.rm = TRUE)
lbub(p, med = TRUE)
legend_move(position = "right")
legend_none()
legend_text(...)
xaxis_title(on = TRUE, ...)
xaxis_ticks(on = TRUE, ...)
yaxis_title(on = TRUE, ...)
```

```
yaxis_text(on = TRUE, ...)

yaxis_ticks(on = TRUE, ...)

facet_text(on = TRUE, ...)

facet_bg(on = TRUE, ...)

panel_bg(on = TRUE, ...)

plot_bg(on = TRUE, ...)

grid_lines(color = "gray50", size = 0.2)

overlay_function(...)
```

#### **Arguments**

V	Either a numeric vector specifying the value(s) at which to draw the vertical or
	horizontal line(s), or an object of any type to use as the first argument to fun.

fun A function, or the name of a function, that returns a numeric vector.

... For the various vline\_, hline\_, and abline\_functions, ... is passed to geom\_vline, geom\_hline, and geom\_abline, respectively, to control the appearance of the

line(s).

For functions ending in \_bg, . . . is passed to element\_rect.

For functions ending in \_text or \_title, . . . is passed to element\_text.

For xaxis\_ticks and yaxis\_ticks, . . . is passed to element\_line.

For overlay\_function, ... is passed to stat\_function.

na.rm A logical scalar passed to the appropriate geom (e.g. geom\_vline). The default

is TRUE.

p The probability mass (in [0,1]) to include in the interval.

med Should the median also be included in addition to the lower and upper bounds

of the interval?

position The position of the legend. Either a numeric vector (of length 2) giving the rel-

ative coordinates (between 0 and 1) for the legend, or a string among "right", "left", "top", "bottom". Using position = "none" is also allowed and is

equivalent to using legend\_none().

on For functions modifying ggplot theme elements, set on=FALSE to set the ele-

ment to element\_blank. For example, facet text can be removed by adding facet\_text(on=FALSE), or simply facet\_text(FALSE) to a ggplot object. If on=TRUE (the default), then . . . can be used to customize the appearance of the

theme element.

color, size Passed to element\_line.

#### **Details**

#### Add vertical, horizontal, and diagonal lines to plots:

• vline\_at and hline\_at return an object created by either geom\_vline or geom\_hline that can be added to a ggplot object to draw a vertical or horizontal line (at one or several values). If fun is missing then the lines are drawn at the values in v. If fun is specified then the lines are drawn at the values returned by fun(v).

- vline\_0 and hline\_0 are wrappers for vline\_at and hline\_at with v = 0 and fun missing.
- abline\_01 is a wrapper for geom\_abline with the intercept set to 0 and the slope set to 1.
- 1bub returns a *function* that takes a single argument x and returns the lower and upper bounds (1b, ub) of the 100\*p% central interval of x, as well as the median (if med is TRUE).

# **Control appearance of facet strips:**

• facet\_text and facet\_bg return ggplot2 theme objects that can be added to an existing plot (ggplot object) to format the text and the background for the facet strips.

# Move legend, remove legend, or style the legend text:

• legend\_move and legend\_none return a ggplot2 theme object that can be added to an existing plot (ggplot object) in order to change the position of the legend (legend\_move) or remove the legend (legend\_none). legend\_text works much like facet\_text, except it controls the legend text.

#### Control appearance of x-axis and y-axis features:

- xaxis\_title and yaxis\_title return a ggplot2 theme object that can be added to an existing plot (ggplot object) in order to toggle or format the titles displayed on the x or y axis. (To change the titles themselves use labs.)
- xaxis\_text and yaxis\_text return a ggplot2 theme object that can be added to an existing plot (ggplot object) in order to toggle or format the text displayed on the x or y axis (e.g. tick labels).
- xaxis\_ticks and yaxis\_ticks return a ggplot2 theme object that can be added to an existing plot (ggplot object) to change the appearance of the axis tick marks.

## **Customize plot background:**

- plot\_bg returns a ggplot2 theme object that can be added to an existing plot (ggplot object) to format the background of the *entire* plot.
- panel\_bg returns a ggplot2 theme object that can be added to an existing plot (ggplot object) to format the background of the just the plotting area.
- grid\_lines returns a ggplot2 theme object that can be added to an existing plot (ggplot object) to add grid lines to the plot background.

## Superimpose a function on an existing plot:

• overlay\_function is a simple wrapper for stat\_function but with the inherit.aes argument fixed to FALSE. Fixing inherit.aes=FALSE will avoid potential errors due to the aesthetic mapping used by certain **bayesplot** plotting functions).

## Value

A **ggplot2** layer or theme object that can be added to existing ggplot objects, like those created by many of the **bayesplot** plotting functions. See the **Details** section.

## See Also

theme\_default for the default ggplot theme used by bayesplot.

```
color_scheme_set("gray")
x <- example_mcmc_draws(chains = 1)</pre>
dim(x)
colnames(x)
### vertical & horizontal lines ###
(p <- mcmc_intervals(x, regex_pars = "beta"))</pre>
# vertical line at zero (with some optional styling)
p + vline_0()
p + vline_0(size = 0.25, color = "darkgray", linetype = 2)
# vertical line(s) at specified values
v <- c(-0.5, 0, 0.5)
p + vline_at(v, linetype = 3, size = 0.25)
my_{lines} < vline_{at}(v, alpha = 0.25, size = 0.75 * c(1, 2, 1),
                    color = c("maroon", "skyblue", "violet"))
p + my_lines
# add vertical line(s) at computed values
# (three ways of getting lines at column means)
color_scheme_set("brightblue")
p <- mcmc_intervals(x, regex_pars = "beta")</pre>
p + vline_at(x[, 3:4], colMeans)
p + vline_at(x[, 3:4], "colMeans", color = "darkgray",
            lty = 2, size = 0.25)
p + vline_at(x[, 3:4], function(a) apply(a, 2, mean),
            color = "orange",
            size = 2, alpha = 0.1)
# using the lbub function to get interval lower and upper bounds (lb, ub)
color_scheme_set("pink")
parsed <- ggplot2::label_parsed</pre>
p2 <- mcmc_hist(x, pars = "beta[1]", binwidth = 1/20,</pre>
               facet_args = list(labeller = parsed))
(p2 \leftarrow p2 + facet_text(size = 16))
b1 <- x[, "beta[1]"]
p2 + vline_at(b1, fun = lbub(0.8), color = "gray20",
             size = 2 * c(1,.5,1), alpha = 0.75)
p2 + vline_at(b1, lbub(0.8, med = FALSE), color = "gray20",
```

size = 2, alpha = 0.75)

```
### format axis titles ###
color_scheme_set("green")
y <- example_y_data()</pre>
yrep <- example_yrep_draws()</pre>
(p3 <- ppc_stat(y, yrep, stat = "median", binwidth = 1/4))</pre>
# turn off the legend, turn on x-axis title
p3 +
legend_none() +
xaxis_title(size = 13, family = "sans") +
ggplot2::xlab(expression(italic(T(y)) == median(italic(y))))
### format axis & facet text ###
####################################
color_scheme_set("gray")
p4 <- mcmc_trace(example_mcmc_draws(), pars = c("alpha", "sigma"))
myfacets <-
facet_bg(fill = "gray30", color = NA) +
facet_text(face = "bold", color = "skyblue", size = 14)
p4 + myfacets
### control tick marks ###
#############################
p4 +
myfacets +
yaxis_text(FALSE) +
yaxis_ticks(FALSE) +
xaxis_ticks(size = 1, color = "skyblue")
###################################
### change plot background ###
###################################
color_scheme_set("blue")
# add grid lines
ppc_stat(y, yrep) + grid_lines()
# panel_bg vs plot_bg
ppc_scatter_avg(y, yrep) + panel_bg(fill = "gray90")
ppc_scatter_avg(y, yrep) + plot_bg(fill = "gray90")
color_scheme_set("yellow")
```

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```
p5 <- ppc_scatter_avg(y, yrep, alpha = 1)
p5 + panel_bg(fill = "gray20") + grid_lines(color = "white")
color_scheme_set("purple")
ppc_dens_overlay(y, yrep[1:30, ]) +
legend_text(size = 14) +
legend_move(c(0.75, 0.5)) +
plot_bg(fill = "gray90") +
panel_bg(color = "black", fill = "gray99", size = 3)
### superimpose a function on existing plot ###
# compare posterior of beta[1] to Gaussian with same posterior mean
# and sd as beta[1]
x <- example_mcmc_draws(chains = 4)</pre>
dim(x)
purple_gaussian <-</pre>
 overlay_function(
   fun = dnorm,
   args = list(mean(x[,, "beta[1]"]), sd(x[,, "beta[1]"])),
   color = "purple",
   size = 2
 )
color_scheme_set("gray")
mcmc_hist(x, pars = "beta[1]") + purple_gaussian
mcmc_dens(x, pars = "beta[1]") + purple_gaussian
```

bayesplot\_grid

Arrange plots in a grid

## **Description**

The bayesplot\_grid function makes it simple to juxtapose plots using common x and/or y axes.

## Usage

```
bayesplot_grid(..., plots = list(), xlim = NULL, ylim = NULL,
  grid_args = list(), titles = character(), subtitles = character(),
  legends = TRUE, save_gg_objects = TRUE)
```

## **Arguments**

... One or more ggplot objects.

plots A list of ggplot objects. Can be used as an alternative to specifying plot objects

via . . . .

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xlim, ylim Optionally, numeric vectors of length 2 specifying lower and upper limits for the axes that will be shared across all plots.

grid\_args An optional named list of arguments to pass to arrangeGrob (nrow, ncol, widths, etc.).

titles, subtitles

Optional character vectors of plot titles and subtitles. If specified, titles and subtitles must must have length equal to the number of plots specified.

legends If any of the plots have legends should they be displayed? Defaults to TRUE. save\_gg\_objects

If TRUE, the default, then the ggplot objects specified in ... or via the plots argument are saved in a list in the "bayesplots" component of the returned object. Setting this to FALSE will make the returned object smaller but these individual plot objects will not be available.

## Value

An object of class "bayesplot\_grid" (essentially a gtable object from arrangeGrob), which has a plot method.

```
y <- example_y_data()</pre>
yrep <- example_yrep_draws()</pre>
stats <- c("sd", "median", "max", "min")</pre>
color_scheme_set("pink")
bayesplot_grid(
 plots = lapply(stats, function(s) ppc_stat(y, yrep, stat = s)),
 titles = stats,
 legends = FALSE,
 grid_args = list(ncol = 1)
## Not run:
library(rstanarm)
mtcars$log_mpg <- log(mtcars$mpg)</pre>
fit1 <- stan_glm(mpg ~ wt, data = mtcars)</pre>
fit2 <- stan_glm(log_mpg ~ wt, data = mtcars)</pre>
y <- mtcars$mpg
yrep1 <- posterior_predict(fit1, draws = 50)</pre>
yrep2 <- posterior_predict(fit2, fun = exp, draws = 50)</pre>
color_scheme_set("blue")
ppc1 <- ppc_dens_overlay(y, yrep1)</pre>
ppc1
ppc1 + yaxis_text()
color_scheme_set("red")
ppc2 <- ppc_dens_overlay(y, yrep2)</pre>
```

bayesplot\_theme\_get

bayesplot\_theme\_get

Get, set, and modify the active bayesplot theme

# **Description**

These functions are the **bayesplot** equivalent to **ggplot2**'s theme\_set and friends. They set, get, and update the active theme but only apply them to bayesplots. The current/active theme is automatically applied to every bayesplot you draw.

Use bayesplot\_theme\_get to get the current **bayesplot** theme, and bayesplot\_theme\_set to change it. bayesplot\_theme\_update and bayesplot\_theme\_replace are shorthands for changing individual elements.

## Usage

```
bayesplot_theme_get()
bayesplot_theme_set(new = theme_default())
bayesplot_theme_update(...)
bayesplot_theme_replace(...)
```

## **Arguments**

The new theme (list of theme elements) to use. This is analogous to the new argument to theme\_set.

... A named list of theme settings.

# **Details**

bayesplot\_theme\_set and friends only apply to bayesplots. However, ggplot2::theme\_set can also be used to change the **bayesplot** theme. Currently, setting a theme with ggplot2::theme\_set (other than the **ggplot2** default theme\_grey) will override the **bayesplot** theme.

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#### Value

bayesplot\_theme\_get returns the current theme. The other three functions (set, update, replace) invisibly return the *previous* theme so it can be saved and easily restored later. This is the same behavior as the **ggplot2** versions of these functions.

#### See Also

theme\_default for the default bayesplot theme.

bayesplot-helpers for a variety of convenience functions, many of which provide shortcuts for tweaking theme elements after creating a plot.

bayesplot-colors to set or view the color scheme used for plotting.

```
library(ggplot2)
# plot using the current value of bayesplot_theme_get()
# (the default is bayesplot::theme_default())
x <- example_mcmc_draws()</pre>
mcmc_hist(x)
# change the bayesplot theme to theme_minimal and save the old theme
old <- bayesplot_theme_set(theme_minimal())</pre>
mcmc_hist(x)
# change back to the previous theme
bayesplot_theme_set(old)
mcmc_hist(x)
# change the default font size and family for bayesplots
bayesplot_theme_update(text = element_text(size = 16, family = "sans"))
mcmc_hist(x)
# change back to the default
bayesplot_theme_set() # same as bayesplot_theme_set(theme_default())
mcmc_hist(x)
# updating theme elements
color_scheme_set("brightblue")
bayesplot_theme_set(theme_dark())
mcmc_hist(x)
bayesplot_theme_update(panel.background = element_rect(fill = "black"))
mcmc_hist(x)
# to get the same plot without updating the theme we could also have
# used the bayeplot convenience function panel_bg()
bayesplot_theme_set(theme_dark())
mcmc_hist(x) + panel_bg(fill = "black")
```

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Combination plots

# **Description**

Combination plots

# Usage

```
mcmc_combo(x, combo = c("dens", "trace"), widths = NULL, gg_theme = NULL,
...)
```

## **Arguments**

widths

gg\_theme

X	A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The
	MCMC-overview page provides details on how to specify each these allowed
	inputs.

A character vector with at least two elements. Each element of combo corresponds to a column in the resulting graphic and should be the name of one of the available MCMC functions (omitting the mcmc\_ prefix).

A numeric vector the same length as combo specifying relative column widths. For example, if the plot has two columns, then widths = c(2,1) will allocate more space for the first column by a factor of 2 (as would widths = c(.3,.15), etc.). The default, NULL, allocates the same horizontal space for each column.

Unlike most of the other **bayesplot** functions, mcmc\_combo returns a gtable object rather than a ggplot object, and so theme objects can't be added directly to the returned plot object. The gg\_theme argument helps get around this problem by accepting a **ggplot2** theme object that is added to each of the plots *before* combining them into the gtable object that is returned. This can be a theme object created by a call to ggplot2::theme or one of the **bayesplot** convenience functions, e.g. legend\_none (see the **Examples** section, below).

Arguments passed to the plotting functions named in combo.

#### Value

A gtable object (the result of calling arrangeGrob) with length(combo) columns and a row for each parameter.

#### See Also

Other MCMC: MCMC-diagnostics, MCMC-distributions, MCMC-intervals, MCMC-nuts, MCMC-overview, MCMC-parcoord, MCMC-recover, MCMC-scatterplots, MCMC-traces

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## **Examples**

```
# some parameter draws to use for demonstration
x <- example_mcmc_draws()</pre>
dim(x)
dimnames(x)
mcmc_combo(x, pars = c("alpha", "sigma"))
mcmc\_combo(x, pars = c("alpha", "sigma"), widths = c(1, 2))
# change second plot, show log(sigma) instead of sigma,
# and remove the legends
color_scheme_set("mix-blue-red")
mcmc_combo(
 х,
 combo = c("dens_overlay", "trace"),
 pars = c("alpha", "sigma"),
 transformations = list(sigma = "log"),
 gg_theme = legend_none()
# same thing but this time also change the entire ggplot theme
mcmc_combo(
х,
 combo = c("dens_overlay", "trace"),
 pars = c("alpha", "sigma"),
 transformations = list(sigma = "log"),
 gg_theme = ggplot2::theme_gray() + legend_none()
)
```

MCMC-diagnostics

General MCMC diagnostics

## **Description**

Plots of Rhat statistics, ratios of effective sample size to total sample size, and autocorrelation of MCMC draws. See the **Plot Descriptions** section, below, for details. For models fit using the No-U-Turn-Sampler, see also MCMC-nuts for additional MCMC diagnostic plots.

# Usage

```
mcmc_rhat(rhat, ..., size = NULL)
mcmc_rhat_hist(rhat, ..., binwidth = NULL, breaks = NULL)
mcmc_rhat_data(rhat, ...)
```

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```
mcmc_neff(ratio, ..., size = NULL)
mcmc_neff_hist(ratio, ..., binwidth = NULL, breaks = NULL)
mcmc_neff_data(ratio, ...)
mcmc_acf(x, pars = character(), regex_pars = character(),
  facet_args = list(), ..., lags = 20, size = NULL)
mcmc_acf_bar(x, pars = character(), regex_pars = character(),
  facet_args = list(), ..., lags = 20)
```

# Arguments

rhat	A vector of Rhat estimates.
	Currently ignored.
size	An optional value to override <pre>geom_point's</pre> default size (for mcmc_rhat, mcmc_neff) or <pre>geom_line's</pre> default size (for mcmc_acf).
binwidth	Passed to geom_histogram to override the default binwidth.
breaks	Passed to geom_histogram as an alternative to binwidth.
ratio	A vector of <i>ratios</i> of effective sample size estimates to total sample size. See neff_ratio.
Х	A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.
pars	An optional character vector of parameter names. If neither pars nor regex_pars is specified then the default is to use <i>all</i> parameters.
regex_pars	An optional regular expression to use for parameter selection. Can be specified instead of pars or in addition to pars.
facet_args	A named list of arguments (other than facets) passed to facet_wrap or facet_grid to control faceting.
lags	The number of lags to show in the autocorrelation plot.

# Value

A ggplot object that can be further customized using the **ggplot2** package. The \_data functions return the data that would have been drawn by the plotting function.

## **Plot Descriptions**

mcmc\_rhat, mcmc\_rhat\_hist Rhat values as either points or a histogram. Values are colored using different shades (lighter is better). The chosen thresholds are somewhat arbitrary, but can be useful guidelines in practice.

- *light*: below 1.05 (good)
- *mid*: between 1.05 and 1.1 (ok)
- dark: above 1.1 (too high)

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mcmc\_neff, mcmc\_neff\_hist Ratios of effective sample size to total sample size as either points or a histogram. Values are colored using different shades (lighter is better). The chosen thresholds are somewhat arbitrary, but can be useful guidelines in practice.

- *light*: between 0.5 and 1 (high)
- *mid*: between 0.1 and 0.5 (good)
- *dark*: below 0.1 (low)

mcmc\_acf Grid of autocorrelation plots by chain and parameter. The lags argument gives the maximum number of lags at which to calculate the autocorrelation function. mcmc\_acf is a line plot whereas mcmc\_acf\_bar is a barplot.

#### References

Stan Development Team. (2016). *Stan Modeling Language Users Guide and Reference Manual*. http://mc-stan.org/documentation/

Gelman, A. and Rubin, D. B. (1992). Inference from iterative simulation using multiple sequences. *Statistical Science*. 7(4), 457–472.

#### See Also

- The Visual MCMC Diagnostics vignette.
- MCMC-nuts for additional MCMC diagnostic plots for models fit using the No-U-Turn-Sampler.

Other MCMC: MCMC-combos, MCMC-distributions, MCMC-intervals, MCMC-nuts, MCMC-overview, MCMC-parcoord, MCMC-recover, MCMC-scatterplots, MCMC-traces

```
# autocorrelation
x <- example_mcmc_draws()</pre>
dim(x)
dimnames(x)
color_scheme_set("green")
mcmc_acf(x, pars = c("alpha", "beta[1]"))
color_scheme_set("pink")
(p <- mcmc_acf_bar(x, pars = c("alpha", "beta[1]")))</pre>
# add horiztonal dashed line at 0.5
p + hline_at(0.5, linetype = 2, size = 0.15, color = "gray")
# fake rhat values to use for demonstration
rhat <- c(runif(100, 1, 1.15))
mcmc_rhat_hist(rhat)
mcmc_rhat(rhat)
# lollipops
color_scheme_set("purple")
```

```
mcmc_rhat(rhat[1:10], size = 5)
color_scheme_set("blue")
mcmc_rhat(runif(1000, 1, 1.07))
mcmc_rhat(runif(1000, 1, 1.3)) + legend_move("top") # add legend above plot
# fake neff ratio values to use for demonstration
ratio <- c(runif(100, 0, 1))
mcmc_neff_hist(ratio)
mcmc_neff(ratio)
## Not run:
# Example using rstanarm model (requires rstanarm package)
library(rstanarm)
# intentionally use small 'iter' so there are some
# problems with rhat and neff for demonstration
fit <- stan_glm(mpg ~ ., data = mtcars, iter = 50)</pre>
rhats <- rhat(fit)</pre>
ratios <- neff_ratio(fit)</pre>
mcmc_rhat(rhats)
mcmc_neff(ratios, size = 3)
# there's a small enough number of parameters in the
# model that we can display their names on the y-axis
mcmc_neff(ratios) + yaxis_text(hjust = 1)
# can also look at autocorrelation
draws <- as.array(fit)</pre>
mcmc_acf(draws, pars = c("wt", "cyl"), lags = 10)
# increase number of iterations and plots look much better
fit2 <- update(fit, iter = 500)</pre>
mcmc_rhat(rhat(fit2))
mcmc_neff(neff_ratio(fit2))
mcmc_acf(as.array(fit2), pars = c("wt", "cyl"), lags = 10)
## End(Not run)
```

 ${\tt MCMC-} distributions$ 

Histograms and kernel density plots of MCMC draws

## **Description**

Various types of histograms and kernel density plots of MCMC draws. See the **Plot Descriptions** section, below, for details.

## **Usage**

```
mcmc_hist(x, pars = character(), regex_pars = character(),
  transformations = list(), facet_args = list(), ..., binwidth = NULL,
 breaks = NULL, freq = TRUE)
mcmc_dens(x, pars = character(), regex_pars = character(),
  transformations = list(), facet_args = list(), ..., trim = FALSE)
mcmc_hist_by_chain(x, pars = character(), regex_pars = character(),
  transformations = list(), facet_args = list(), ..., binwidth = NULL,
  freq = TRUE)
mcmc_dens_overlay(x, pars = character(), regex_pars = character(),
  transformations = list(), facet_args = list(), color_chains = TRUE, ...,
  trim = FALSE)
mcmc_dens_chains(x, pars = character(), regex_pars = character(),
  transformations = list(), color_chains = TRUE, ..., bw = NULL,
  adjust = NULL, kernel = NULL, n_dens = NULL)
mcmc_dens_chains_data(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., bw = NULL, adjust = NULL,
  kernel = NULL, n_dens = NULL)
mcmc_violin(x, pars = character(), regex_pars = character(),
  transformations = list(), facet_args = list(), ..., probs = c(0.1, 0.5, 0.5)
 0.9))
```

# Arguments

Х

A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.

pars

An optional character vector of parameter names. If neither pars nor regex\_pars is specified then the default is to use *all* parameters.

regex\_pars

An optional regular expression to use for parameter selection. Can be specified instead of pars or in addition to pars.

## transformations

Optionally, transformations to apply to parameters before plotting. If transformations is a function or a single string naming a function then that function will be used to transform all parameters. To apply transformations to particular parameters, the transformations argument can be a named list with length equal to the number of parameters to be transformed. Currently only univariate transformations of scalar parameters can be specified (multivariate transformations will be implemented in a future release). If transformations is a list, the name of each list element should be a parameter name and the content of each list element should be a function (or any item to match as a function via match.fun,

e.g. a string naming a function). If a function is specified by its name as a string (e.g. "log"), then it can be used to construct a new parameter label for the appropriate parameter (e.g. "log(sigma)"). If a function itself is specified (e.g. log or function(x) log(x)) then "t" is used in the new parameter label to indicate that the parameter is transformed (e.g. "t(sigma)").

Note: due to partial argument matching transformations can be abbreviated

for convenience in interactive use (e.g., transform, trans, etc.).

facet\_args A named list of arguments (other than facets) passed to facet\_wrap or facet\_grid

to control faceting.

... Currently ignored.

binwidth Passed to geom\_histogram to override the default binwidth.

Passed to geom\_histogram as an alternative to binwidth.

freq For histograms, freq=TRUE (the default) puts count on the y-axis. Setting freq=FALSE

puts density on the y-axis. (For many plots the y-axis text is off by default. To view the count or density labels on the y-axis see the yaxis\_text convenience

function.)

trim A logical scalar passed to geom\_density.
color\_chains option for whether to separately color chains.

bw, adjust, kernel, n\_dens

Optional arguments passed to density to override default kernel density esti-

mation parameters. n\_dens defaults to 1024.

probs A numeric vector passed to geom\_violin's draw\_quantiles argument to spec-

ify at which quantiles to draw horizontal lines. Set to NULL to remove the lines.

## Value

A ggplot object that can be further customized using the **ggplot2** package.

#### **Plot Descriptions**

mcmc\_hist Histograms of posterior draws with all chains merged.

mcmc\_dens Kernel density plots of posterior draws with all chains merged.

mcmc\_hist\_by\_chain Histograms of posterior draws with chains separated via faceting.

mcmc\_dens\_overlay Kernel density plots of posterior draws with chains separated but overlaid on a single plot.

mcmc\_violin The density estimate of each chain is plotted as a violin with horizontal lines at notable quantiles.

mcmc\_dens\_chains Ridgeline kernel density plots of posterior draws with chains separated but overlaid on a single plot. In mcmc\_dens\_overlay parameters appear in separate facets; in mcmc\_dens\_chains they appear in the same panel and can overlap vertically.

## See Also

Other MCMC: MCMC-combos, MCMC-diagnostics, MCMC-intervals, MCMC-nuts, MCMC-overview, MCMC-parcoord, MCMC-recover, MCMC-scatterplots, MCMC-traces

```
set.seed(9262017)
# some parameter draws to use for demonstration
x <- example_mcmc_draws()</pre>
dim(x)
dimnames(x)
### Histograms ###
######################
# histograms of all parameters
color_scheme_set("brightblue")
mcmc_hist(x)
# histograms of some parameters
color_scheme_set("pink")
mcmc_hist(x, pars = c("alpha", "beta[2]"))
mcmc_hist(x, pars = "sigma", regex_pars = "beta")
# example of using 'transformations' argument to plot log(sigma),
# and parsing facet labels (e.g. to get greek letters for parameters)
mcmc_hist(x, transformations = list(sigma = "log"),
          facet_args = list(labeller = ggplot2::label_parsed)) +
          facet_text(size = 15)
# instead of list(sigma = "log"), you could specify the transformation as
# list(sigma = log) or list(sigma = function(x) log(x)), but then the
# label for the transformed sigma is 't(sigma)' instead of 'log(sigma)'
mcmc_hist(x, transformations = list(sigma = log))
# separate histograms by chain
color_scheme_set("pink")
mcmc_hist_by_chain(x, regex_pars = "beta")
###################
### Densities ###
#################
mcmc_dens(x, pars = c("sigma", "beta[2]"),
          facet_args = list(nrow = 2))
# separate and overlay chains
color_scheme_set("mix-teal-pink")
mcmc_dens_overlay(x, pars = c("sigma", "beta[2]"),
                  facet_args = list(nrow = 2)) +
                  facet_text(size = 14)
x2 <- example_mcmc_draws(params = 6)</pre>
mcmc_dens_chains(x2, pars = c("beta[1]", "beta[2]", "beta[3]"))
```

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```
# separate chains as violin plots
color_scheme_set("green")
mcmc_violin(x) + panel_bg(color = "gray20", size = 2, fill = "gray30")
```

MCMC-intervals

Plot interval estimates from MCMC draws

## Description

Plot central (quantile-based) posterior interval estimates from MCMC draws. See the **Plot Descriptions** section, below, for details.

## Usage

```
mcmc_intervals(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., prob = 0.5, prob_outer = 0.9,
  point_est = c("median", "mean", "none"), rhat = numeric())
mcmc_areas(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., prob = 0.5, prob_outer = 1,
  point_est = c("median", "mean", "none"), rhat = numeric(), bw = NULL,
  adjust = NULL, kernel = NULL, n_dens = NULL)
mcmc_areas_ridges(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., prob_outer = 1, prob = 1, bw = NULL,
  adjust = NULL, kernel = NULL, n_dens = NULL)
mcmc_intervals_data(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., prob = 0.5, prob_outer = 0.9,
  point_est = c("median", "mean", "none"), rhat = numeric())
mcmc_areas_data(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., prob = 0.5, prob_outer = 1,
 point_est = c("median", "mean", "none"), rhat = numeric(), bw = NULL.
  adjust = NULL, kernel = NULL, n_dens = NULL)
mcmc_areas_ridges_data(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., prob_outer = 1, prob = 1, bw = NULL,
  adjust = NULL, kernel = NULL, n_dens = NULL)
```

# Arguments

Χ

A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.

pars

An optional character vector of parameter names. If neither pars nor regex\_pars is specified then the default is to use *all* parameters.

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regex\_pars

An optional regular expression to use for parameter selection. Can be specified instead of pars or in addition to pars.

transformations

Optionally, transformations to apply to parameters before plotting. If transformations is a function or a single string naming a function then that function will be used to transform all parameters. To apply transformations to particular parameters, the transformations argument can be a named list with length equal to the number of parameters to be transformed. Currently only univariate transformations of scalar parameters can be specified (multivariate transformations will be implemented in a future release). If transformations is a list, the name of each list element should be a parameter name and the content of each list element should be a function (or any item to match as a function via match. fun, e.g. a string naming a function). If a function is specified by its name as a string (e.g. "log"), then it can be used to construct a new parameter label for the appropriate parameter (e.g. "log(sigma)"). If a function itself is specified (e.g. log or function(x) log(x)) then "t" is used in the new parameter label to indicate that the parameter is transformed (e.g. "t(sigma)").

Note: due to partial argument matching transformations can be abbreviated for convenience in interactive use (e.g., transform, trans, etc.).

. .

Currently unused.

prob

The probability mass to include in the inner interval (for mcmc\_intervals) or in the shaded region (for mcmc\_areas). The default is 0.5 (50% interval) and 1 for mcmc\_areas\_ridges.

prob\_outer

The probability mass to include in the outer interval. The default is 0.9 for mcmc\_intervals (90% interval) and 1 for mcmc\_areas and for mcmc\_areas\_ridges.

point\_est

The point estimate to show. Either "median" (the default), "mean", or "none".

rhat

An optional numeric vector of  $\hat{R}$  estimates, with one element per parameter included in x. If rhat is provided, the intervals/areas and point estimates in the resulting plot are colored based on  $\hat{R}$  value. See rhat for methods for extracting  $\hat{R}$  estimates.

bw, adjust, kernel, n\_dens

Optional arguments passed to density to override default kernel density estimation parameters. n\_dens defaults to 1024.

# Value

A ggplot object that can be further customized using the **ggplot2** package. The \_data functions return the data that would have been drawn by the plotting function.

## **Plot Descriptions**

mcmc\_intervals Plots of uncertainty intervals computed from posterior draws with all chains merged.

mcmc\_areas Density plots computed from posterior draws with all chains merged, with uncertainty intervals shown as shaded areas under the curves.

mcmc\_areas\_ridges Density plot, as in mcmc\_areas, but drawn with overlapping ridgelines. This plot provides a compact display of (hierarchically) related distributions.

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## See Also

 $Other\ MCMC: \texttt{MCMC-combos}, \texttt{MCMC-diagnostics}, \texttt{MCMC-distributions}, \texttt{MCMC-nuts}, \texttt{MCMC-overview}, \texttt{MCMC-parcoord}, \texttt{MCMC-recover}, \texttt{MCMC-scatterplots}, \texttt{MCMC-traces}$ 

```
set.seed(9262017)
# some parameter draws to use for demonstration
x <- example_mcmc_draws(params = 6)</pre>
dim(x)
dimnames(x)
color_scheme_set("brightblue")
mcmc_intervals(x)
mcmc_intervals(x, pars = c("beta[1]", "beta[2]"))
mcmc_areas(x, regex_pars = "beta\[[1-3]", prob = 0.8) +
 ggplot2::labs(
   title = "Posterior distributions",
   subtitle = "with medians and 80% intervals"
color_scheme_set("red")
mcmc_areas(
  х,
  pars = c("alpha", "beta[4]"),
   prob = 2/3,
  prob_outer = 0.9,
  point_est = "mean"
)
# color by rhat value
color_scheme_set("blue")
fake_rhat_values <- c(1, 1.07, 1.3, 1.01, 1.15, 1.005)
mcmc_intervals(x, rhat = fake_rhat_values)
mcmc_intervals_data(x)
mcmc_intervals_data(x, rhat = fake_rhat_values)
mcmc_areas_data(x, pars = "alpha")
color_scheme_set("gray")
p \leftarrow mcmc\_areas(x, pars = c("alpha", "beta[4]"), rhat = c(1, 1.1))
p + legend_move("bottom")
p + legend_move("none") # or p + legend_none()
# apply transformations
mcmc_intervals(
  Х,
  pars = c("beta[2]", "sigma"),
  transformations = list("sigma" = "log", "beta[2]" = function(x) x + 3)
)
```

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```
# apply same transformation to all selected parameters
mcmc_intervals(x, regex_pars = "beta", transformations = "exp")
## Not run:
# example using fitted model from rstanarm package
library(rstanarm)
fit <- stan_glm(</pre>
 mpg \sim 0 + wt + factor(cyl),
 data = mtcars,
 iter = 500
x <- as.matrix(fit)</pre>
color_scheme_set("teal")
mcmc_intervals(x, point_est = "mean", prob = 0.8, prob_outer = 0.95)
mcmc_areas(x, regex_pars = "cyl", bw = "SJ",
           rhat = rhat(fit, regex_pars = "cyl"))
## End(Not run)
## Not run:
# Example of hierarchically related parameters
# plotted with ridgelines
m <- shinystan::eight_schools@posterior_sample</pre>
mcmc_areas_ridges(m, pars = "mu", regex_pars = "theta") +
 ggplot2::ggtitle("Treatment effect on eight schools (Rubin, 1981)")
## End(Not run)
```

MCMC-nuts

Diagnostic plots for the No-U-Turn-Sampler (NUTS)

# Description

Diagnostic plots for the No-U-Turn-Sampler (NUTS), the default MCMC algorithm used by Stan. See the **Plot Descriptions** section, below.

## Usage

```
mcmc_nuts_acceptance(x, lp, chain = NULL, ..., binwidth = NULL)
mcmc_nuts_divergence(x, lp, chain = NULL, ...)
mcmc_nuts_stepsize(x, lp, chain = NULL, ...)
mcmc_nuts_treedepth(x, lp, chain = NULL, ...)
mcmc_nuts_energy(x, ..., binwidth = NULL, alpha = 0.5,
    merge_chains = FALSE)
```

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## **Arguments**

A molten data frame of NUTS sampler parameters, either created by nuts\_params Х or in the same form as the object returned by nuts\_params. A molten data frame of draws of the log-posterior or, more commonly, of a 1p quantity equal to the log-posterior up to a constant. 1p should either be created via log\_posterior or be an object with the same form as the object returned by log\_posterior. chain A positive integer for selecting a particular chain. The default (NULL) is to merge the chains before plotting. If chain = k then the plot for chain k is overlaid (in a darker shade but with transparency) on top of the plot for all chains. The chain argument is not used by mcmc\_nuts\_energy. Currently ignored. binwidth An optional value passed to geom\_histogram to override the default binwidth. For mcmc\_nuts\_energy only, the transparency (alpha) level in [0,1] used for the alpha overlaid histogram. For mcmc\_nuts\_energy only, should all chains be merged or displayed sepamerge\_chains rately? The default is FALSE, i.e., to show the chains separately.

#### Value

A gtable object (the result of calling arrangeGrob) created from several ggplot objects, except for mcmc\_nuts\_energy, which returns a ggplot object.

# **Quick Definitions**

For more details see Stan Development Team (2016) and Betancourt (2017).

- accept\_stat\_\_: the average acceptance probabilities of all possible samples in the proposed tree.
- divergent\_\_: the number of leapfrog transitions with diverging error. Because NUTS terminates at the first divergence this will be either 0 or 1 for each iteration.
- stepsize\_\_: the step size used by NUTS in its Hamiltonian simulation.
- treedepth\_\_: the depth of tree used by NUTS, which is the log (base 2) of the number of leapfrog steps taken during the Hamiltonian simulation.
- energy\_\_: the value of the Hamiltonian (up to an additive constant) at each iteration.

## **Plot Descriptions**

mcmc\_nuts\_acceptance Three plots:

- Histogram of accept\_stat\_\_ with vertical lines indicating the mean (solid line) and median (dashed line).
- Histogram of 1p\_ with vertical lines indicating the mean (solid line) and median (dashed line).
- Scatterplot of accept\_stat\_\_ vs lp\_\_.

mcmc\_nuts\_divergence Two plots:

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- Violin plots of lp\_\_|divergent\_\_=1 and lp\_\_|divergent\_\_=0.
- Violin plots of accept\_stat\_\_|divergent\_\_=1 and accept\_stat\_\_|divergent\_\_=0.

mcmc\_nuts\_stepsize Two plots:

- Violin plots of lp\_\_ by chain ordered by stepsize\_\_ value.
- Violin plots of accept\_stat\_\_ by chain ordered by stepsize\_\_ value.

mcmc\_nuts\_treedepth Three plots:

- Violin plots of lp\_\_ by value of treedepth\_\_.
- Violin plots of accept\_stat\_\_ by value of treedepth\_\_.
- Histogram of treedepth\_\_.

mcmc\_nuts\_energy Overlaid histograms showing energy\_\_ vs the change in energy\_\_. See Betancourt (2016) for details.

#### References

Betancourt, M. (2017). A conceptual introduction to Hamiltonian Monte Carlo. https://arxiv.org/abs/1701.02434

Betancourt, M. and Girolami, M. (2013). Hamiltonian Monte Carlo for hierarchical models. https://arxiv.org/abs/1312.0906

Hoffman, M. D. and Gelman, A. (2014). The No-U-Turn Sampler: adaptively setting path lengths in Hamiltonian Monte Carlo. *Journal of Machine Learning Research*. 15:1593–1623.

Stan Development Team. (2016). *Stan Modeling Language Users Guide and Reference Manual*. http://mc-stan.org/documentation/

#### See Also

- The Visual MCMC Diagnostics vignette.
- Several other plotting functions in the bayesplot package are not NUTS-specific but take optional extra arguments if the model was fit using NUTS:
  - mcmc\_trace: show divergences as tick marks below the trace plot.
  - mcmc\_parcoord: change the color/size/transparency of lines correspondending to divergences.
  - mcmc\_scatter: change the color/size/shape of points corresponding to divergences.
  - mcmc\_pairs: change the color/size/shape of points corresponding divergences and/or max treedepth saturation.

 $\label{lem:combos} Other\ MCMC-combos,\ MCMC-diagnostics,\ MCMC-distributions,\ MCMC-intervals,\ MCMC-overview,\ MCMC-parcoord,\ MCMC-recover,\ MCMC-scatterplots,\ MCMC-traces$ 

```
## Not run:
library(ggplot2)
library(rstanarm)
fit <- stan_glm(mpg ~ wt + am, data = mtcars, iter = 1000)
np <- nuts_params(fit)
lp <- log_posterior(fit)</pre>
```

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```
color_scheme_set("brightblue")
mcmc_nuts_acceptance(np, lp)
mcmc_nuts_acceptance(np, lp, chain = 2)

mcmc_nuts_divergence(np, lp)
mcmc_nuts_stepsize(np, lp)
mcmc_nuts_treedepth(np, lp)

color_scheme_set("red")
mcmc_nuts_energy(np)
mcmc_nuts_energy(np, merge_chains = TRUE, binwidth = .15)
mcmc_nuts_energy(np) +
  facet_wrap(~ Chain, nrow = 1) +
  coord_fixed(ratio = 150) +
  ggtitle("NUTS Energy Diagnostic")

## End(Not run)
```

MCMC-overview

Plots for Markov chain Monte Carlo simulations

## **Description**

The **bayesplot** MCMC module provides various plotting functions for creating graphical displays of Markov chain Monte Carlo (MCMC) simulations. The **MCMC plotting functions** section, below, provides links to the documentation for various categories of MCMC plots. Currently the MCMC plotting functions accept posterior draws provided in one of the following formats:

- 3-D array: An array with dimensions [Iteration, Chain, Parameter] in that order.
- **list**: A list of matrices, where each matrix corresponds to a Markov chain. All of the matrices should have the same number of iterations (rows) and parameters (columns), and parameters should have the same names and be in the same order.
- matrix: A matrix with one column per parameter. If using matrix there should only be a single Markov chain or all chains should already be merged (stacked).
- data frame: There are two types of data frames allowed. Either a data frame with one column per parameter (if only a single chain or all chains have already been merged), or a data frame with one column per parameter plus an additional column "Chain" that contains the chain number (an integer) corresponding to each row in the data frame.

**Note**: typically the user should *not* include warmup iterations in the object passed to **bayesplot** plotting functions, although for certain plots (e.g. trace plots) it can occasionally be useful to include the warmup iterations for diagnostic purposes.

MCMC-parcoord 33

## MCMC plotting functions

**Posterior distributions** Histograms and kernel density plots of parameter draws, optionally showing each Markov chain separately.

**Uncertainty intervals** Uncertainty intervals computed from parameter draws.

**Trace plots** Times series of parameter draws, optionally including with HMC/NUTS diagnostic information.

**Scatterplots** Scatterplots, heatmaps, and pairs plots of parameter draws, optionally including with HMC/NUTS diagnostic information.

**Parallel coordinates plots** Parallel coordinates plot of MCMC draws (one dimension per parameter), optionally including with HMC/NUTS diagnostic information.

**Combinations** Combination plots (e.g. trace plot + histogram).

General MCMC diagnostics MCMC diagnostic plots including Rhat, effective sample size, autocorrelation.

**NUTS diagnostics** Special diagnostic plots for the No-U-Turn Sampler.

Comparisons to "true" values Plots comparing MCMC estimates to "true" parameter values (e.g., values used to simulate data).

#### References

Gabry, J., Simpson, D., Vehtari, A., Betancourt, M., and Gelman, A. (2018). Visualization in Bayesian workflow. *Journal of the Royal Statistical Society Series A*, accepted for publication. arXiv preprint: http://arxiv.org/abs/1709.01449.

#### See Also

 $\label{lem:combos} Other\ MCMC-combos,\ MCMC-diagnostics,\ MCMC-distributions,\ MCMC-intervals,\ MCMC-nuts,\ MCMC-parcoord,\ MCMC-recover,\ MCMC-scatterplots,\ MCMC-traces$ 

MCMC-parcoord

Parallel coordinates plot of MCMC draws

# Description

Parallel coordinates plot of MCMC draws (one dimension per parameter). See the **Plot Descriptions** section, below, for details.

## Usage

```
mcmc_parcoord(x, pars = character(), regex_pars = character(),
    transformations = list(), ..., size = 0.2, alpha = 0.3, np = NULL,
    np_style = parcoord_style_np())

mcmc_parcoord_data(x, pars = character(), regex_pars = character(),
    transformations = list(), np = NULL)

parcoord_style_np(div_color = "red", div_size = 0.2, div_alpha = 0.2)
```

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#### **Arguments**

Х

A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.

pars

An optional character vector of parameter names. If neither pars nor regex\_pars is specified then the default is to use *all* parameters.

regex\_pars

An optional regular expression to use for parameter selection. Can be specified instead of pars or in addition to pars.

#### transformations

Optionally, transformations to apply to parameters before plotting. If transformations is a function or a single string naming a function then that function will be used to transform all parameters. To apply transformations to particular parameters, the transformations argument can be a named list with length equal to the number of parameters to be transformed. Currently only univariate transformations of scalar parameters can be specified (multivariate transformations will be implemented in a future release). If transformations is a list, the name of each list element should be a parameter name and the content of each list element should be a function (or any item to match as a function via match. fun, e.g. a string naming a function). If a function is specified by its name as a string (e.g. "log"), then it can be used to construct a new parameter label for the appropriate parameter (e.g. "log(sigma)"). If a function itself is specified (e.g. log or function(x) log(x)) then "t" is used in the new parameter label to indicate that the parameter is transformed (e.g. "t(sigma)").

Note: due to partial argument matching transformations can be abbreviated for convenience in interactive use (e.g., transform, trans, etc.).

... Currently ignored.

size, alpha Arguments passed on to geom\_line.

np

For models fit using NUTS (more generally, any symplectic integrator), an optional data frame providing NUTS diagnostic information. The data frame should be the object returned by nuts\_params or one with the same structure.

np\_style

A call to the parcoord\_style\_np helper function to specify arguments controlling the appearance of superimposed lines representing NUTS diagnostics (in this case divergences) if the np argument is specified.

div\_color, div\_size, div\_alpha

Optional arguments to the parcoord\_style\_np helper function that are eventually passed to <code>geom\_line</code> if the np argument is also specified. They control the color, size, and transparency specifications for showing divergences in the plot. The default values are displayed in the <code>Usage</code> section above.

#### Value

A ggplot object that can be further customized using the **ggplot2** package. The \_data functions return the data that would have been drawn by the plotting function.

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## **Plot Descriptions**

mcmc\_parcoord (Parallel coordinates plot) of MCMC draws. There is one dimension per parameter along the horizontal axis and each set of connected line segments represents a single MCMC draw (i.e., a vector of length equal to the number of parameters).

The parallel coordinates plot is most useful if the optional HMC/NUTS diagnostic information is provided via the np argument. In that case divergences are highlighted in the plot. The appearance of the divergences can be customized using the np\_style argument and the parcoord\_style\_np helper function.

When the plotted model parameters are on very different scales the transformations argument can be useful. For example, to standardize all variables before plotting you could use function (x - mean(x))/sd(x) when specifying the transformations argument to mcmc\_parcoord. See the **Examples** section for how to do this.

#### References

Gabry, J., Simpson, D., Vehtari, A., Betancourt, M., and Gelman, A. (2018). Visualization in Bayesian workflow. *Journal of the Royal Statistical Society Series A*, accepted for publication. arXiv preprint: http://arxiv.org/abs/1709.01449.

Hartikainen, A. (2017, Aug 23). Concentration of divergences [Msg 21]. Message posted to The Stan Forums: http://discourse.mc-stan.org/t/concentration-of-divergences/1590/21.

#### See Also

Other MCMC: MCMC-combos, MCMC-diagnostics, MCMC-distributions, MCMC-intervals, MCMC-nuts, MCMC-overview, MCMC-recover, MCMC-scatterplots, MCMC-traces

```
color_scheme_set("pink")
x <- example_mcmc_draws(params = 5)</pre>
mcmc_parcoord(x)
mcmc_parcoord(x, regex_pars = "beta")
## Not run:
# Example using a Stan demo model
library(rstan)
fit <- stan_demo("eight_schools")</pre>
draws <- as.array(fit, pars = c("mu", "tau", "theta", "lp__"))</pre>
np <- nuts_params(fit)</pre>
str(np)
levels(np$Parameter)
color_scheme_set("brightblue")
mcmc_parcoord(draws, alpha = 0.05)
mcmc_parcoord(draws, np = np)
# customize appearance of divergences
color_scheme_set("darkgray")
div_style <- parcoord_style_np(div_color = "green", div_size = 0.05, div_alpha = 0.4)
mcmc_parcoord(draws, size = 0.25, alpha = 0.1,
```

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```
np = np, np_style = div_style)
# to use a transformation (e.g., to standarde all the variables)
# specify the 'transformations' argument (though partial argument name
# matching means we can just use 'trans' or 'transform')
mcmc_parcoord(
 draws,
 transform = function(x) \{(x - mean(x)) / sd(x)\},\
 size = 0.25,
 alpha = 0.1,
 np = np,
 np_style = div_style
# mcmc_parcoord_data returns just the data in a conventient form for plotting
d <- mcmc_parcoord_data(x, np = np)</pre>
head(d)
tail(d)
## End(Not run)
```

MCMC-recover

Compare MCMC estimates to "true" parameter values

# **Description**

Plots comparing MCMC estimates to "true" parameter values. Before fitting a model to real data it is useful to simulate data according to the model using known (fixed) parameter values and to check that these "true" parameter values are (approximately) recovered by fitting the model to the simulated data. See the **Plot Descriptions** section, below, for details on the available plots.

## Usage

```
mcmc_recover_intervals(x, true, batch = rep(1, length(true)),
  facet_args = list(), ..., prob = 0.5, prob_outer = 0.9,
  point_est = c("median", "mean", "none"), size = 4, alpha = 1)

mcmc_recover_scatter(x, true, batch = rep(1, length(true)),
  facet_args = list(), ..., point_est = c("median", "mean"), size = 3,
  alpha = 1)

mcmc_recover_hist(x, true, facet_args = list(), ..., binwidth = NULL,
  breaks = NULL)
```

#### **Arguments**

Х

A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.

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true	A numeric vector of "true" values of the parameters in x. There should be one value in true for each parameter included in x and the order of the parameters in true should be the same as the order of the parameters in x.
batch	Optionally, a vector-like object (numeric, character, integer, factor) used to split the parameters into batches. If batch is specified, it must have the same length as true and be in the same order as true. Parameters in the same batch will be grouped together in the same facet in the plot (see the <b>Examples</b> section, below). The default is to group all parameters together into a single batch. Changing the default is most useful when parameters are on very different scales, in which case batch can be used to group them into batches within which it makes sense to use the same <i>y</i> -axis.
facet_args	A named list of arguments (other than facets) passed to facet_wrap or facet_grid to control faceting.
	Currently unused.
prob	The probability mass to include in the inner interval. The default is $0.5 (50\%$ interval).
prob_outer	The probability mass to include in the outer interval. The default is $0.9 (90\%$ interval).
point_est	The point estimate to show. Either "median" (the default), "mean", or "none".
size, alpha	Passed to geom_point to control the appearance of plotted points.
binwidth	Passed to geom_histogram to override the default binwidth.
breaks	Passed to geom_histogram as an alternative to binwidth.

## Value

A ggplot object that can be further customized using the ggplot2 package.

# **Plot Descriptions**

mcmc\_recover\_intervals Central intervals and point estimates computed from MCMC draws, with "true" values plotted using a different shape.

mcmc\_recover\_scatter Scatterplot of posterior means (or medians) against "true" values.

mcmc\_recover\_hist Histograms of the draws for each parameter with the "true" value overlaid as a vertical line.

# See Also

 $Other\ MCMC-combos, \ MCMC-diagnostics, \ MCMC-distributions, \ MCMC-intervals, \ MCMC-nuts, \ MCMC-overview, \ MCMC-parcoord, \ MCMC-scatterplots, \ MCMC-traces$ 

```
## Not run:
library(rstanarm)
alpha <- 1; beta <- rnorm(10, 0, 3); sigma <- 2
X <- matrix(rnorm(1000), 100, 10)</pre>
```

```
y <- rnorm(100, mean = c(alpha + X %*% beta), sd = sigma)
fit <- stan_glm(y ~ ., data = data.frame(y, X))</pre>
draws <- as.matrix(fit)</pre>
print(colnames(draws))
true <- c(alpha, beta, sigma)</pre>
mcmc_recover_intervals(draws, true)
# put the coefficients on X into the same batch
mcmc_recover_intervals(draws, true, batch = c(1, rep(2, 10), 1))
# equivalent
mcmc_recover_intervals(draws, true, batch = grepl("X", colnames(draws)))
# same but facets stacked vertically
mcmc_recover_intervals(draws, true,
                       batch = grepl("X", colnames(draws)),
                       facet_args = list(ncol = 1),
                       size = 3)
# each parameter in its own facet
mcmc_recover_intervals(draws, true, batch = 1:ncol(draws))
# same but in a different order
mcmc_recover_intervals(draws, true, batch = c(1, 3, 4, 2, 5:12))
# present as bias by centering with true values
mcmc_recover_intervals(sweep(draws, 2, true), rep(0, ncol(draws))) + hline_0()
# scatterplot of posterior means vs true values
mcmc_recover_scatter(draws, true, point_est = "mean")
# histograms of parameter draws with true value added as vertical line
color_scheme_set("brightblue")
mcmc_recover_hist(draws[, 1:4], true[1:4])
## End(Not run)
```

MCMC-scatterplots

Scatterplots of MCMC draws

# **Description**

Scatterplots, hexagonal heatmaps, and pairs plots from MCMC draws. See the **Plot Descriptions** section, below, for details.

# Usage

```
mcmc_scatter(x, pars = character(), regex_pars = character(),
  transformations = list(), ..., size = 2.5, alpha = 0.8, np = NULL,
  np_style = scatter_style_np())
```

```
mcmc_hex(x, pars = character(), regex_pars = character(),
    transformations = list(), ..., binwidth = NULL)

mcmc_pairs(x, pars = character(), regex_pars = character(),
    transformations = list(), ..., diag_fun = c("hist", "dens"),
    off_diag_fun = c("scatter", "hex"), diag_args = list(),
    off_diag_args = list(), condition = pairs_condition(), lp = NULL,
    np = NULL, np_style = pairs_style_np(), max_treedepth = NULL,
    grid_args = list(), save_gg_objects = TRUE)

scatter_style_np(div_color = "red", div_shape = 16, div_size = 2.5,
    div_alpha = 1)

pairs_style_np(div_color = "red", div_shape = 4, div_size = 1,
    div_alpha = 1, td_color = "yellow2", td_shape = 3, td_size = 1,
    td_alpha = 1)

pairs_condition(chains = NULL, draws = NULL, nuts = NULL)
```

#### **Arguments**

Х

A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.

pars

An optional character vector of parameter names. For mcmc\_scatter and mcmc\_hex only two parameters can be selected. To plot more than two parameters use mcmc\_pairs.

regex\_pars

An optional regular expression to use for parameter selection. Can be specified instead of pars or in addition to pars.

#### transformations

Optionally, transformations to apply to parameters before plotting. If transformations is a function or a single string naming a function then that function will be used to transform all parameters. To apply transformations to particular parameters, the transformations argument can be a named list with length equal to the number of parameters to be transformed. Currently only univariate transformations of scalar parameters can be specified (multivariate transformations will be implemented in a future release). If transformations is a list, the name of each list element should be a parameter name and the content of each list element should be a function (or any item to match as a function via match. fun, e.g. a string naming a function). If a function is specified by its name as a string (e.g. "log"), then it can be used to construct a new parameter label for the appropriate parameter (e.g. "log(sigma)"). If a function itself is specified (e.g. log or function(x) log(x)) then "t" is used in the new parameter label to indicate that the parameter is transformed (e.g. "t(sigma)").

Note: due to partial argument matching transformations can be abbreviated for convenience in interactive use (e.g., transform, trans, etc.).

... Currently ignored.

size, alpha For mcmc\_scatter, passed to geom\_point to control the appearance of the

points.

Optionally, a data frame of NUTS sampler parameters, either created by nuts\_params np

or in the same form as the object returned by nuts\_params. The colors, shapes, and sizes of the superimposed points can be customized using the np\_style

argument.

np\_style If np is specified, np\_style can be a call to the scatter\_style\_np helper func-

tion (for mcmc\_scatter) or the pairs\_style\_np helper function (for mcmc\_pars) to specify arguments controlling the appearance of superimposed points representing NUTS diagnostic information. (Note: for pairs\_style\_np the size

arguments are interpreted as scaling factors).

binwidth For mcmc\_hex, an optional numeric vector of *length two* passed to geom\_hex to

override the default binwidth in both the vertical and horizontal directions.

diag\_fun, off\_diag\_fun

For mcmc\_pairs, the plotting function to use for the plots along the diagonal and for the off-diagonal plots, respectively. Currently diag\_fun can be "hist" for histogram or "dens" for density, and off\_diag\_fun can be "scatter" for

scatterplot or "hex" for a hexagonal heatmap.

diag\_args, off\_diag\_args

For mcmc\_pairs, optional named lists of arguments to pass to the functions implied by the diag\_fun and off\_diag\_fun arguments, respectively. For example, if off\_diag\_fun is "scatter" then off\_diag\_args could include optional

arguments to mcmc\_scatter like size and alpha.

condition

For mcmc\_pairs, a call to the pairs\_condition helper function, which is used to specify a criterion for determining which chains (or iterations) are shown in the plots above the diagonal and which are shown in the plots below the diagonal. The histograms (or density plots) along the diagonal are always made using all chains and iterations, but the scatterplots (or hex plots) above and below the diagonal show different combinations of chains/iterations depending on condition. The default is a call to pairs\_condition with none of its arguments specified. In this case half of the chains (or roughly half if there are an odd number) will be used in the plots above the diagonal and the rest in the plots below the diagonal. The chains, draws, and nuts arguments to pairs\_condition, which are documented below, can be used to change this

default.

For mcmc\_pairs, a molten data frame of draws of the log-posterior or, more commonly, of a quantity equal to the log-posterior up to a constant. 1p should either be created via log\_posterior or be an object with the same form as the

object returned by log\_posterior.

max\_treedepth For mcmc\_pairs, an integer representing the maximum treedepth allowed when fitting the model (if fit using NUTS). This is only needed for detecting which

transitions (if any) hit the maximum treedepth.

grid\_args, save\_gg\_objects

For mcmc\_pairs, arguments to pass to bayesplot\_grid. For example, since mcmc\_pairs returns more than a single ggplot object, using ggtitle afterwards will not work. But you you can still add a title to the plot using grid\_args = list(top="My title").

1p

div\_color, div\_shape, div\_size, div\_alpha, td\_color, td\_shape, td\_size, td\_alpha

Optional arguments to the scatter\_style\_np or pairs\_style\_np helper functions that are eventually passed to geom\_point. The default values are displayed
in the Usage section above.

chains, draws, nuts

Optional arguments to the pairs\_condition helper function, which is used to specify the condition argument for mcmc\_pairs.

- The chains argument can be used to select some subset of the chains. If chains is an integer vector then the behavior is the same as the default (half the chains above the diagonal and half below) except using only the specified subset of chains. Alternatively, chains can be a list of two integer vectors with the first specifying the chains to be shown in the plots above the diagonal and the second for below the diagonal.
- The draws argument to pairs\_condition can be used to directly specify which realizations are plotted above and below the diagonal. draws can be a single proportion, which is interpreted as the proportion of realizations (among all chains) to plot in the lower panel starting with the first realization in each chain, with the complement (from the end of each chain) plotted in the upper panel. Alternatively draws can be a logical vector with length equal to the product of the number of iterations and the number of chains, in which case realizations corresponding to FALSE and TRUE will be plotted in the lower and upper panels, respectively.
- For models fit using NUTS, the nuts argument to pairs\_condition can be used. It takes a (possibly abbreviated) string to select among "accept\_stat\_\_", "stepsize\_\_", "treedepth\_\_", "n\_leapfrog\_\_", "divergent\_\_", "energy\_\_", and "lp\_\_". These are the sampler parameters associated with NUTS (and "lp\_\_" is the log-posterior up to an additive constant). In this case, plots below the diagonal will contain realizations that are below the median of the indicated variable (or are zero in the case of "divergent\_\_"), and plots above the diagonal will contain realizations that are greater than or equal to the median of the indicated variable (or are one in the case of "divergent\_\_"). If "lp\_\_" is used then the lp argument to mcmc\_pairs must also be specified. For the other NUTS parameters the np argument to mcmc\_pairs must also be specified.

#### Value

mcmc\_scatter and mcmc\_hex return a ggplot object that can be further customized using the **gg-plot2** package.

mcmc\_pairs returns many ggplot objects organized into a grid via bayesplot\_grid.

## **Plot Descriptions**

mcmc\_scatter Bivariate scatterplot of posterior draws. If using a very large number of posterior draws then mcmc\_hex may be preferable to avoid overplotting. For models fit using NUTS the np, and np\_style arguments can be used to add additional information in the plot (in this case the approximate location of divergences).

mcmc\_hex Hexagonal heatmap of 2-D bin counts. This plot is useful in cases where the posterior sample size is large enough that mcmc\_scatter suffers from overplotting.

mcmc\_pairs A square plot matrix with univariate marginal distributions along the diagonal (as histograms or kernel density plots) and bivariate distributions off the diagonal (as scatterplots or hex heatmaps).

For the off-diagonal plots, the default is to split the chains so that (roughly) half are displayed above the diagonal and half are below (all chains are always merged together for the plots along the diagonal). Other possibilities are available by setting the condition argument.

Additionally, extra diagnostic information for models fit using NUTS can be added to the pairs plot using the lp, np, and np\_style arguments. If np is specified (and condition is *not* "divergent\_\_"), then points (red, by default) will be superimposed onto the off-diagonal plots indicating which (if any) iterations encountered a divergent transition. Also, if both np and max\_treedepth are specified then points (yellow, by default) will be superimposed to indicate a transition that hit the maximum treedepth rather than terminated its evolution normally. The np\_style argument can be used with the pairs\_style\_np convenience function to change the appearance of these overlaid points. See the **Examples** section.

#### References

Gabry, J., Simpson, D., Vehtari, A., Betancourt, M., and Gelman, A. (2018). Visualization in Bayesian workflow. *Journal of the Royal Statistical Society Series A*, accepted for publication. arXiv preprint: http://arxiv.org/abs/1709.01449.

#### See Also

Other MCMC: MCMC-combos, MCMC-diagnostics, MCMC-distributions, MCMC-intervals, MCMC-nuts, MCMC-overview, MCMC-parcoord, MCMC-recover, MCMC-traces

```
library("ggplot2")
# some parameter draws to use for demonstration
x <- example_mcmc_draws(params = 6)</pre>
dimnames(x)
# scatterplot of alpha vs log(sigma)
color_scheme_set("teal")
(p <- mcmc_scatter(x, pars = c("alpha", "sigma"),</pre>
                  transform = list(sigma = "log")))
p +
 labs(
    title = "Insert your own headline-grabbing title",
    subtitle = "with a provocative subtitle",
    caption = "and a controversial caption",
    x = expression(alpha),
   y = expression(log(sigma))
# add ellipse
```

```
p + stat_ellipse(level = 0.9, color = "gray20", size = 1)
# add contour
color_scheme_set("red")
p2 <- mcmc_scatter(x, pars = c("alpha", "sigma"), size = 3.5, alpha = 0.25)
p2 + stat_density_2d(color = "black", size = .5)
# can also add lines/smooths
color_scheme_set("pink")
(p3 <- mcmc_scatter(x, pars = c("alpha", "beta[3]"), alpha = 0.25, size = 3))</pre>
p3 + geom_smooth(method = "lm", se = FALSE, color = "gray20",
                 size = .75, linetype = 2)
# hexagonal heatmap
color_scheme_set("brightblue")
(p <- mcmc_hex(x, pars = c("sigma", "alpha"), transform = list(sigma = "log")))</pre>
p + plot_bg(fill = "gray95")
p + plot_bg(fill = "gray95") + panel_bg(fill = "gray70")
color_scheme_set("purple")
# pairs plots
# default of condition=NULL implies splitting chains between upper and lower panels
mcmc_pairs(x, pars = "alpha", regex_pars = "beta\\[[1,4]\\]",
           off_diag_args = list(size = 1, alpha = 0.5))
# change to density plots instead of histograms and hex plots instead of
# scatterplots
mcmc_pairs(x, pars = "alpha", regex_pars = "beta\\[[1,4]\\]",
           diag_fun = "dens", off_diag_fun = "hex")
# plot chain 1 above diagonal and chains 2, 3, and 4 below
color_scheme_set("brightblue")
mcmc_pairs(x, pars = "alpha", regex_pars = "beta\\[[1,4]\\]",
           diag_fun = "dens", off_diag_fun = "hex",
           condition = pairs_condition(chains = list(1, 2:4)))
### Adding NUTS diagnostics to scatterplots and pairs plots
# examples using rstanarm package
library(rstanarm)
# for demonstration purposes, intentionally fit a model that
# will (almost certainly) have some divergences
fit <- stan_glm(</pre>
  mpg \sim ., data = mtcars,
  iter = 1000,
  # this combo of prior and adapt_delta should lead to some divergences
  prior = hs(),
```

```
adapt_delta = 0.9
posterior <- as.array(fit)</pre>
np <- nuts_params(fit)</pre>
# mcmc_scatter with divergences highlighted
color_scheme_set("brightblue")
mcmc_scatter(posterior, pars = c("wt", "sigma"), np = np)
color_scheme_set("darkgray")
div_style <- scatter_style_np(div_color = "green", div_shape = 4, div_size = 4)</pre>
mcmc_scatter(posterior, pars = c("sigma", "(Intercept)"),
             np = np, np_style = div_style)
# split the draws according to above/below median accept_stat__
# and show approximate location of divergences (red points)
color_scheme_set("brightblue")
mcmc_pairs(
  posterior,
  pars = c("wt", "cyl", "sigma"),
  off_diag_args = list(size = 1, alpha = 1/3),
  condition = pairs_condition(nuts = "accept_stat__"),
  np = np
)
# more customizations:
# - transform sigma to log(sigma)
# - median log-posterior as 'condition'
# - hex instead of scatter for off-diagonal plots
# - show points where max treedepth hit in blue
color_scheme_set("darkgray")
mcmc_pairs(
  posterior,
  pars = c("wt", "cyl", "sigma"),
  transform = list(sigma = "log"),
  off_diag_fun = "hex",
  condition = pairs_condition(nuts = "lp__"),
  lp = log_posterior(fit),
  np = np,
  np_style = pairs_style_np(div_color = "firebrick",
                            td_color = "blue",
                            td_size = 2),
  # for demonstration purposes, set max_treedepth to a value that will
  # result in at least a few max treedepth warnings
  max_treedepth = with(np, -1 + max(Value[Parameter == "treedepth__"]))
)
## End(Not run)
```

#### **Description**

Trace plot (or traceplot) of MCMC draws. See the **Plot Descriptions** section, below, for details.

## Usage

```
mcmc_trace(x, pars = character(), regex_pars = character(),
    transformations = list(), facet_args = list(), ..., n_warmup = 0,
    window = NULL, size = NULL, np = NULL, np_style = trace_style_np(),
    divergences = NULL)

mcmc_trace_highlight(x, pars = character(), regex_pars = character(),
    transformations = list(), facet_args = list(), ..., n_warmup = 0,
    window = NULL, size = NULL, alpha = 0.2, highlight = 1)

trace_style_np(div_color = "red", div_size = 0.25, div_alpha = 1)
```

#### **Arguments**

Х

A 3-D array, matrix, list of matrices, or data frame of MCMC draws. The MCMC-overview page provides details on how to specify each these allowed inputs.

pars

An optional character vector of parameter names. If neither pars nor regex\_pars is specified then the default is to use *all* parameters.

regex\_pars

An optional regular expression to use for parameter selection. Can be specified instead of pars or in addition to pars.

## transformations

Optionally, transformations to apply to parameters before plotting. If transformations is a function or a single string naming a function then that function will be used to transform all parameters. To apply transformations to particular parameters, the transformations argument can be a named list with length equal to the number of parameters to be transformed. Currently only univariate transformations of scalar parameters can be specified (multivariate transformations will be implemented in a future release). If transformations is a list, the name of each list element should be a parameter name and the content of each list element should be a function (or any item to match as a function via match. fun, e.g. a string naming a function). If a function is specified by its name as a string (e.g. "log"), then it can be used to construct a new parameter label for the appropriate parameter (e.g. "log(sigma)"). If a function itself is specified (e.g. log or function(x) log(x)) then "t" is used in the new parameter label to indicate that the parameter is transformed (e.g. "t(sigma)").

Note: due to partial argument matching transformations can be abbreviated for convenience in interactive use (e.g., transform, trans, etc.).

facet\_args

A named list of arguments (other than facets) passed to facet\_wrap or facet\_grid to control faceting.

... Currently ignored.

An integer; the number of warmup iterations included in x. The default is n\_warmup n\_warmup = 0, i.e. to assume no warmup iterations are included. If n\_warmup > 0 then the background for iterations 1: n\_warmup is shaded gray. window An integer vector of length two specifying the limits of a range of iterations to display. size An optional value to override the default line size (mcmc\_trace) or the default point size (mcmc\_trace\_highlight). np For models fit using NUTS (more generally, any symplectic integrator), an optional data frame providing NUTS diagnostic information. The data frame should be the object returned by nuts\_params or one with the same structure. If np is specified then tick marks are added to the bottom of the trace plot indicating within which iterations there was a divergence (if there were any). See the end of the **Examples** section, below. np\_style A call to the trace\_style\_np helper function to specify arguments controlling the appearance of tick marks representing divergences (if the np argument is specified). Deprecated. Use the np argument instead. divergences alpha For mcmc\_trace\_highlight, passed to geom\_point to control the transparency of the points for the chains not highlighted. highlight For mcmc\_trace\_highlight, an integer specifying one of the chains that will be more visible than the others in the plot.

div\_color, div\_size, div\_alpha

Optional arguments to the trace\_style\_np helper function that are eventually passed to geom\_rug if the np argument is also specified. They control the color, size, and transparency specifications for showing divergences in the plot. The default values are displayed in the **Usage** section above.

#### Value

A ggplot object that can be further customized using the **ggplot2** package.

#### **Plot Descriptions**

mcmc\_trace Standard trace plots of MCMC draws. For models fit using NUTS, the np argument can be used to also show divergences on the trace plot.

mcmc\_trace\_highlight Traces are plotted using points rather than lines and the opacity of all chains but one (specified by the highlight argument) is reduced.

#### See Also

 $\label{lem:combos} Other\ MCMC-combos,\ MCMC-diagnostics,\ MCMC-distributions,\ MCMC-intervals,\ MCMC-nuts,\ MCMC-overview,\ MCMC-parcoord,\ MCMC-recover,\ MCMC-scatterplots$ 

```
# some parameter draws to use for demonstration
x <- example_mcmc_draws(chains = 4, params = 6)</pre>
dim(x)
dimnames(x)
# trace plots of the betas
color_scheme_set("viridis")
mcmc_trace(x, regex_pars = "beta")
color_scheme_set("viridisA")
mcmc_trace(x, regex_pars = "beta")
color_scheme_set("viridisC")
mcmc_trace(x, regex_pars = "beta")
# mix color schemes
color_scheme_set("mix-blue-red")
mcmc_trace(x, regex_pars = "beta")
# use traditional ggplot discrete color scale
mcmc_trace(x, pars = c("alpha", "sigma")) +
ggplot2::scale_color_discrete()
# zoom in on a window of iterations, increase line size,
# add tick marks, move legend to the top, add gray background
color_scheme_set("viridisA")
mcmc_trace(x[,, 1:4], window = c(100, 130), size = 1) +
  panel_bg(fill = "gray90", color = NA) +
  legend_move("top")
## Not run:
# parse facet label text
color_scheme_set("purple")
p <- mcmc_trace(</pre>
 х,
  regex_pars = "beta\\[[1,3]\\]",
  facet_args = list(labeller = ggplot2::label_parsed)
p + facet_text(size = 15)
# mark first 100 draws as warmup
mcmc_trace(x, n_warmup = 100)
# plot as points, highlighting chain 2
color_scheme_set("brightblue")
mcmc_trace_highlight(x, pars = "sigma", highlight = 2, size = 2)
# for models fit using HMC/NUTS divergences can be displayed in the trace plot
library("rstanarm")
fit <- stan_glm(mpg ~ ., data = mtcars,</pre>
```

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```
# next line to keep example fast and also ensure we get some divergences
                prior = hs(), iter = 400, adapt_delta = 0.8)
# extract draws using as.array (instead of as.matrix) to keep
# chains separate for trace plot
posterior <- as.array(fit)</pre>
# for stanfit and stanreg objects use nuts_params() to get the divergences
mcmc_trace(posterior, pars = "sigma", np = nuts_params(fit))
color_scheme_set("viridis")
mcmc_trace(
 posterior,
 pars = c("wt", "sigma"),
 size = 0.5,
 facet_args = list(nrow = 2),
 np = nuts_params(fit),
 np_style = trace_style_np(div_color = "black", div_size = 0.5)
)
color_scheme_set("viridis")
mcmc_trace(
 posterior,
 pars = c("wt", "sigma"),
 size = 0.8,
 facet_args = list(nrow = 2),
 divergences = nuts_params(fit),
 div_color = "black"
## End(Not run)
```

PPC-discrete

PPCs for discrete outcomes

#### **Description**

Many of the PPC functions in **bayesplot** can be used with discrete data. The small subset of these functions that can *only* be used if y and yrep are discrete are documented on this page. Currently these include rootograms for count outcomes and bar plots for ordinal, categorical, and multinomial outcomes. See the **Plot Descriptions** section below.

# Usage

```
ppc_bars(y, yrep, ..., prob = 0.9, width = 0.9, size = 1, fatten = 3,
    freq = TRUE)

ppc_bars_grouped(y, yrep, group, facet_args = list(), ..., prob = 0.9,
    width = 0.9, size = 1, fatten = 3, freq = TRUE)
```

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```
ppc_rootogram(y, yrep, style = c("standing", "hanging", "suspended"), ...,
prob = 0.9, size = 1)
```

# **Arguments**

у	A vector of observations. See <b>Details</b> .
yrep	An $S$ by $N$ matrix of draws from the posterior predictive distribution, where $S$ is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and $N$ is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See <b>Details</b> for additional instructions.
• • •	Currently unused.
prob	A value between 0 and 1 indicating the desired probability mass to include in the yrep intervals. Set prob=0 to remove the intervals. For ppc_rootogram these are intervals of the <i>square roots</i> of the expected counts.
width	For ppc_bars and ppc_bars_grouped, passed to geom_bar to control the bar width.
size, fatten	For ppc_bars and ppc_bars_grouped, size and fatten are passed to geom_pointrange to control the appearance of the yrep points and intervals. For ppc_rootogram size is passed to geom_line.
freq	For ppc_bars and ppc_bars_grouped, if TRUE (the default) the y-axis will display counts. Setting freq=FALSE will put proportions on the y-axis.
group	A grouping variable (a vector or factor) the same length as y. Each value in group is interpreted as the group level pertaining to the corresponding value of y.
facet_args	An optional list of arguments (other than facets) passed to facet_wrap to control faceting.
style	For ppc_rootogram, a string specifying the rootogram style. The options are "standing", "hanging", and "suspended". See the <b>Plot Descriptions</b> section, below, for details on the different styles.

#### **Details**

For all of these plots y and yrep must be non-negative integers, although they need not be integers in the strict sense of R's integer type.

# Value

A ggplot object that can be further customized using the **ggplot2** package.

# **Plot Descriptions**

ppc\_bars Bar plot of y with yrep medians and uncertainty intervals superimposed on the bars.

ppc\_bars\_grouped Same as ppc\_bars but a separate plot (facet) is generated for each level of a grouping variable.

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ppc\_rootogram Rootograms allow for diagnosing problems in count data models such as overdispersion or excess zeros. They consist of a histogram of y with the expected counts based on yrep overlaid as a line along with uncertainty intervals. The y-axis represents the square roots of the counts to approximately adjust for scale differences and thus ease comparison between observed and expected counts. Using the style argument, the histogram style can be adjusted to focus on different aspects of the data:

- Standing: basic histogram of observed counts with curve showing expected counts.
- Hanging: observed counts counts hanging from the curve representing expected counts.
- Suspended: histogram of the differences between expected and observed counts.

All of these are plotted on the square root scale. See Kleiber and Zeileis (2016) for advice on interpreting rootograms and selecting among the different styles.

#### References

Kleiber, C. and Zeileis, A. (2016). Visualizing count data regressions using rootograms. *The American Statistician*. 70(3): 296–303. https://arxiv.org/abs/1605.01311.

#### See Also

Other PPCs: PPC-distributions, PPC-errors, PPC-intervals, PPC-loo, PPC-overview, PPC-scatterplots, PPC-test-statistics

```
set.seed(9222017)
# bar plots
f <- function(N) {</pre>
 sample(1:4, size = N, replace = TRUE, prob = c(0.25, 0.4, 0.1, 0.25))
y < - f(100)
yrep <- t(replicate(500, f(100)))</pre>
dim(yrep)
group \leftarrow gl(2, 50, length = 100, labels = c("GroupA", "GroupB"))
color_scheme_set("mix-pink-blue")
ppc_bars(y, yrep)
# split by group, change interval width, and display proportion
# instead of count on y-axis
color_scheme_set("mix-blue-pink")
ppc_bars_grouped(y, yrep, group, prob = 0.5, freq = FALSE)
# rootograms for counts
y <- rpois(100, 20)
yrep <- matrix(rpois(10000, 20), ncol = 100)</pre>
color_scheme_set("brightblue")
ppc_rootogram(y, yrep)
ppc_rootogram(y, yrep, prob = 0)
```

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```
ppc_rootogram(y, yrep, style = "hanging", prob = 0.8)
ppc_rootogram(y, yrep, style = "suspended")
```

PPC-distributions

PPC distributions

## **Description**

Compare the empirical distribution of the data y to the distributions of simulated/replicated data yrep from the posterior predictive distribution. See the **Plot Descriptions** section, below, for details

#### Usage

```
ppc_data(y, yrep, group = NULL)

ppc_hist(y, yrep, ..., binwidth = NULL, breaks = NULL, freq = TRUE)

ppc_boxplot(y, yrep, ..., notch = TRUE, size = 0.5, alpha = 1)

ppc_freqpoly(y, yrep, ..., binwidth = NULL, freq = TRUE, size = 0.25, alpha = 1)

ppc_freqpoly_grouped(y, yrep, group, ..., binwidth = NULL, freq = TRUE, size = 0.25, alpha = 1)

ppc_dens(y, yrep, ..., trim = FALSE, size = 0.5, alpha = 1)

ppc_dens_overlay(y, yrep, ..., size = 0.25, alpha = 0.7, trim = FALSE, bw = "nrd0", adjust = 1, kernel = "gaussian", n_dens = 1024)

ppc_ecdf_overlay(y, yrep, ..., discrete = FALSE, pad = TRUE, size = 0.25, alpha = 0.7)

ppc_violin_grouped(y, yrep, group, ..., probs = c(0.1, 0.5, 0.9), size = 1, alpha = 1, y_draw = c("violin", "points", "both"), y_size = 1, y_alpha = 1, y_jitter = 0.1)
```

# **Arguments**

У

A vector of observations. See Details.

yrep

An S by N matrix of draws from the posterior predictive distribution, where S is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and N is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See **Details** for additional instructions.

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group A grouping variable (a vector or factor) the same length as y. Each value in

group is interpreted as the group level pertaining to the corresponding value of

у.

... Currently unused.

binwidth Passed to geom\_histogram to override the default binwidth.

breaks Passed to geom\_histogram as an alternative to binwidth.

freq For histograms, freq=TRUE (the default) puts count on the y-axis. Setting freq=FALSE

puts density on the y-axis. (For many plots the y-axis text is off by default. To view the count or density labels on the y-axis see the yaxis\_text convenience

function.)

notch A logical scalar passed to geom\_boxplot. Unlike for geom\_boxplot, the default

is notch=TRUE.

size, alpha Passed to the appropriate geom to control the appearance of the yrep distribu-

tions.

trim A logical scalar passed to geom\_density.

bw, adjust, kernel, n\_dens

Optional arguments passed to density to override default kernel density esti-

mation parameters. n\_dens defaults to 1024.

discrete For ppc\_ecdf\_overlay, should the data be treated as discrete? The default is

FALSE, in which case geom="line" is passed to stat\_ecdf. If discrete is set

to TRUE then geom="step" is used.

pad A logical scalar passed to stat\_ecdf.

probs A numeric vector passed to geom\_violin's draw\_quantiles argument to spec-

ify at which quantiles to draw horizontal lines. Set to NULL to remove the lines.

y\_draw For ppc\_violin\_grouped, a string specifying how to draw y: "violin" (de-

fault), "points" (jittered points), or "both".

y\_jitter, y\_size, y\_alpha

For ppc\_violin\_grouped, if y\_draw is "points" or "both" then y\_size, y\_alpha, and y\_jitter are passed to to the size, alpha, and width arguments of geom\_jitter to control the appearance of y points. The default of y\_jitter=NULL will let gg-

plot2 determine the amount of jitter.

## Details

For Binomial data, the plots will typically be most useful if y and yrep contain the "success" proportions (not discrete "success" or "failure" counts).

#### Value

A ggplot object that can be further customized using the **ggplot2** package. The \_data functions return the data that would have been drawn by the plotting function.

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#### **Plot Descriptions**

ppc\_hist, ppc\_freqpoly, ppc\_dens, ppc\_boxplot A separate histogram, shaded frequency polygon, smoothed kernel density estimate, or box and whiskers plot is displayed for y and each dataset (row) in yrep. For these plots yrep should therefore contain only a small number of rows. See the **Examples** section.

- ppc\_freqpoly\_grouped A separate frequency polygon is plotted for each level of a grouping variable for y and each dataset (row) in yrep. For this plot yrep should therefore contain only a small number of rows. See the **Examples** section.
- ppc\_dens\_overlay, ppc\_ecdf\_overlay Kernel density or empirical CDF estimates of each dataset (row) in yrep are overlaid, with the distribution of y itself on top (and in a darker shade). When using ppc\_ecdf\_overlay with discrete data set the discrete argument to TRUE for better results.
- ppc\_violin\_grouped The density estimate of yrep within each level of a grouping variable is plotted as a violin with horizontal lines at notable quantiles. y is overlaid on the plot either as a violin, points, or both, depending on the y\_draw argument.

#### References

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (Ch. 6)

#### See Also

Other PPCs: PPC-discrete, PPC-errors, PPC-intervals, PPC-loo, PPC-overview, PPC-scatterplots, PPC-test-statistics

```
color_scheme_set("brightblue")
y <- example_y_data()
yrep <- example_yrep_draws()
dim(yrep)
ppc_dens_overlay(y, yrep[1:25, ])

ppc_ecdf_overlay(y, yrep[sample(nrow(yrep), 25), ])

# for ppc_hist,dens,freqpoly,boxplot definitely use a subset yrep rows so
# only a few (instead of nrow(yrep)) histograms are plotted
ppc_hist(y, yrep[1:8, ])

color_scheme_set("red")
ppc_boxplot(y, yrep[1:8, ])

# wizard hat plot
color_scheme_set("blue")
ppc_dens(y, yrep[200:202, ])</pre>
```

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PPC-errors

PPC errors

#### **Description**

Various plots of predictive errors y - yrep. See the **Details** and **Plot Descriptions** sections, below.

# Usage

```
ppc_error_hist(y, yrep, ..., binwidth = NULL, breaks = NULL, freq = TRUE)

ppc_error_hist_grouped(y, yrep, group, ..., binwidth = NULL, breaks = NULL, freq = TRUE)

ppc_error_scatter(y, yrep, ..., size = 2.5, alpha = 0.8)

ppc_error_scatter_avg(y, yrep, ..., size = 2.5, alpha = 0.8)

ppc_error_scatter_avg_vs_x(y, yrep, x, ..., size = 2.5, alpha = 0.8)

ppc_error_binned(y, yrep, ..., size = 1, alpha = 0.25)
```

## **Arguments**

У

A vector of observations. See **Details**.

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yrep An S by N matrix of draws from the posterior predictive distribution, where S is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and N is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See **Details** for additional instructions.

... Currently unused.

binwidth Passed to geom\_histogram to override the default binwidth.

breaks Passed to geom\_histogram as an alternative to binwidth.

freq For histograms, freq=TRUE (the default) puts count on the y-axis. Setting freq=FALSE

puts density on the y-axis. (For many plots the y-axis text is off by default. To view the count or density labels on the y-axis see the yaxis\_text convenience

function.)

group A grouping variable (a vector or factor) the same length as y. Each value in

group is interpreted as the group level pertaining to the corresponding value of

у.

size, alpha For scatterplots, arguments passed to geom\_point to control the appearance of

the points. For the binned error plot, arguments controlling the size of the outline

and opacity of the shaded region indicating the 2-SE bounds.

A numeric vector the same length as y to use as the x-axis variable.

#### **Details**

All of these functions (aside from the \*\_scatter\_avg functions) compute and plot predictive errors for each row of the matrix yrep, so it is usually a good idea for yrep to contain only a small number of draws (rows). See **Examples**, below.

For binomial and Bernoulli data the ppc\_error\_binned function can be used to generate binned error plots. Bernoulli data can be input as a vector of 0s and 1s, whereas for binomial data y and yrep should contain "success" proportions (not counts). See the **Examples** section, below.

#### Value

A ggplot object that can be further customized using the **ggplot2** package.

# Plot descriptions

- ppc\_error\_hist A separate histogram is plotted for the predictive errors computed from y and each dataset (row) in yrep. For this plot yrep should have only a small number of rows.
- ppc\_error\_hist\_grouped Like ppc\_error\_hist, except errors are computed within levels of a grouping variable. The number of histograms is therefore equal to the product of the number of rows in yrep and the number of groups (unique values of group).
- ppc\_error\_scatter A separate scatterplot is displayed for y vs. the predictive errors computed from y and each dataset (row) in yrep. For this plot yrep should have only a small number of rows.
- ppc\_error\_scatter\_avg A single scatterplot of y vs. the average of the errors computed from y and each dataset (row) in yrep. For each individual data point y[n] the average error is the average of the errors for y[n] computed over the draws from the posterior predictive distribution.

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ppc\_error\_scatter\_avg\_vs\_x Same as ppc\_error\_scatter\_avg, except the average is plotted on the y-axis and a a predictor variable x is plotted on the x-axis.

ppc\_error\_binned Intended for use with binomial data. A separate binned error plot (similar to binnedplot) is generated for each dataset (row) in yrep. For this plot y and yrep should contain proportions rather than counts, and yrep should have only a small number of rows.

#### References

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (Ch. 6)

#### See Also

Other PPCs: PPC-discrete, PPC-distributions, PPC-intervals, PPC-loo, PPC-overview, PPC-scatterplots, PPC-test-statistics

```
y <- example_y_data()</pre>
yrep <- example_yrep_draws()</pre>
ppc_error_hist(y, yrep[1:3, ])
# errors within groups
group <- example_group_data()</pre>
(p1 <- ppc_error_hist_grouped(y, yrep[1:3, ], group))</pre>
p1 + yaxis_text() # defaults to showing counts on y-axis
table(group) # more obs in GroupB, can set freq=FALSE to show density on y-axis
(p2 <- ppc_error_hist_grouped(y, yrep[1:3, ], group, freq = FALSE))</pre>
p2 + yaxis_text()
# scatterplots
ppc_error_scatter(y, yrep[10:14, ])
ppc_error_scatter_avg(y, yrep)
x <- example_x_data()</pre>
ppc_error_scatter_avg_vs_x(y, yrep, x)
# ppc_error_binned with binomial model from rstanarm
## Not run:
library(rstanarm)
example("example_model", package = "rstanarm")
formula(example_model)
# get observed proportion of "successes"
y <- example_model$y # matrix of "success" and "failure" counts
trials <- rowSums(y)</pre>
y_prop <- y[, 1] / trials # proportions</pre>
# get predicted success proportions
yrep <- posterior_predict(example_model)</pre>
```

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```
yrep_prop <- sweep(yrep, 2, trials, "/")
ppc_error_binned(y_prop, yrep_prop[1:6, ])
## End(Not run)</pre>
```

PPC-intervals

PPC intervals

#### **Description**

Medians and central interval estimates of yrep with y overlaid. See the **Plot Descriptions** section, below.

## Usage

```
ppc_intervals(y, yrep, x = NULL, ..., prob = 0.5, prob_outer = 0.9,
    size = 1, fatten = 3)

ppc_intervals_grouped(y, yrep, x = NULL, group, facet_args = list(), ...,
    prob = 0.5, prob_outer = 0.9, size = 1, fatten = 3)

ppc_ribbon(y, yrep, x = NULL, ..., prob = 0.5, prob_outer = 0.9,
    alpha = 0.33, size = 0.25)

ppc_ribbon_grouped(y, yrep, x = NULL, group, facet_args = list(), ...,
    prob = 0.5, prob_outer = 0.9, alpha = 0.33, size = 0.25)

ppc_intervals_data(y, yrep, x = NULL, group = NULL, prob = 0.5,
    prob_outer = 0.9, ...)

ppc_ribbon_data(y, yrep, x = NULL, group = NULL, prob = 0.5,
    prob_outer = 0.9, ...)
```

#### **Arguments**

У

A vector of observations. See **Details**.

yrep

An S by N matrix of draws from the posterior predictive distribution, where S is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and N is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See **Details** for additional instructions.

Х

A numeric vector the same length as y to use as the x-axis variable. For example, x could be a predictor variable from a regression model, a time variable for timeseries models, etc. If x is missing or NULL, then 1:length(y) is used for the x-axis.

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Currently unused.

prob, prob\_outer

Values between 0 and 1 indicating the desired probability mass to include in the inner and outer intervals. The defaults are prob=0.5 and prob\_outer=0.9.

group

A grouping variable (a vector or factor) the same length as y. Each value in group is interpreted as the group level pertaining to the corresponding value of y.

facet\_args

An optional list of arguments (other than facets) passed to facet\_wrap to control faceting.

alpha, size, fatten

Arguments passed to geoms. For ribbon plots alpha and size are passed to geom\_ribbon. For interval plots size and fatten are passed to geom\_pointrange.

#### Value

A ggplot object that can be further customized using the **ggplot2** package. The \_data functions return the data that would have been drawn by the plotting function.

#### **Plot Descriptions**

ppc\_intervals, ppc\_ribbon 100\*prob% central intervals for yrep at each x value. ppc\_intervals plots intervals as vertical bars with points indicating yrep medians and darker points indicating observed y values. ppc\_ribbon plots a ribbon of connected intervals with a line through the median of yrep and a darker line connecting observed y values. In both cases an optional x variable can also be specified for the x-axis variable.

Depending on the number of observations and the variability in the predictions at different values of x, one or the other of these plots may be easier to read than the other.

ppc\_intervals\_grouped, ppc\_ribbon\_grouped Same as ppc\_intervals and ppc\_ribbon, respectively, but a separate plot (facet) is generated for each level of a grouping variable.

# References

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (Ch. 6)

#### See Also

 $Other\ PPCs:\ PPC-discrete,\ PPC-distributions,\ PPC-errors,\ PPC-loo,\ PPC-overview,\ PPC-scatterplots,\ PPC-test-statistics$ 

```
y <- rnorm(50)
yrep <- matrix(rnorm(5000, 0, 2), ncol = 50)

color_scheme_set("brightblue")
ppc_ribbon(y, yrep)
ppc_intervals(y, yrep)</pre>
```

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```
# change x axis to y values (instead of indices) and add x = y line
ppc_intervals(y, yrep, x = y) + abline_01()
color_scheme_set("teal")
year <- 1950:1999
ppc_ribbon(y, yrep, x = year, alpha = 0, size = 0.75) + ggplot2::xlab("Year")
color_scheme_set("pink")
year < - rep(2000:2009, each = 5)
group \leftarrow gl(5, 1, length = 50, labels = LETTERS[1:5])
ppc_ribbon_grouped(y, yrep, x = year, group) +
  ggplot2::scale_x_continuous(breaks = pretty)
ppc_ribbon_grouped(
y, yrep, x = year, group,
 facet_args = list(scales = "fixed"),
 alpha = 1,
size = 2
) +
xaxis_text(FALSE) +
 xaxis_ticks(FALSE) +
 panel_bg(fill = "gray20")
ppc_dat <- ppc_intervals_data(y, yrep, x = year, prob = 0.5)</pre>
ppc_group_dat <- ppc_intervals_data(y, yrep, x = year, group = group, prob = 0.5)</pre>
## Not run:
library("rstanarm")
fit <- stan_glmer(mpg ~ wt + (1|cyl), data = mtcars)</pre>
yrep <- posterior_predict(fit)</pre>
color_scheme_set("purple")
with(mtcars, ppc_intervals(mpg, yrep, x = wt, prob = 0.5)) +
 panel_bg(fill="gray90", color = NA) +
grid_lines(color = "white")
ppc_intervals_grouped(y = mtcars$mpg, yrep, prob = 0.8,
                      x = mtcars$wt, group = mtcars$cyl)
color_scheme_set("gray")
ppc_intervals(mtcars$mpg, yrep, prob = 0.5) +
 ggplot2::scale_x_continuous(
  labels = rownames(mtcars),
  breaks = 1:nrow(mtcars)
 xaxis_text(angle = -70, vjust = 1, hjust = 0)
## End(Not run)
```

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PPC-loo

LOO predictive checks

## **Description**

Leave-One-Out (LOO) predictive checks. See the **Plot Descriptions** section below for details.

# Usage

```
ppc_loo_pit_overlay(y, yrep, lw, pit, samples = 100, ..., size = 0.25,
    alpha = 0.7, trim = FALSE, bw = "nrd0", adjust = 1,
    kernel = "gaussian", n_dens = 1024)

ppc_loo_pit_qq(y, yrep, lw, pit, compare = c("uniform", "normal"), ...,
    size = 2, alpha = 1)

ppc_loo_pit(y, yrep, lw, pit, compare = c("uniform", "normal"), ...,
    size = 2, alpha = 1)

ppc_loo_intervals(y, yrep, psis_object, subset = NULL, intervals = NULL,
    ..., prob = 0.5, prob_outer = 0.9, size = 1, fatten = 3,
    order = c("index", "median"))

ppc_loo_ribbon(y, yrep, lw, psis_object, subset = NULL, intervals = NULL,
    ..., prob = 0.5, prob_outer = 0.9, alpha = 0.33, size = 0.25)
```

# **Arguments**

У	A vector of observations. See <b>Details</b> .
yrep	An $S$ by $N$ matrix of draws from the posterior predictive distribution, where $S$ is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and $N$ is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See <b>Details</b> for additional instructions.
lw	A matrix of (smoothed) log weights with the same dimensions as yrep. See psis and the associated weights method and the <b>Examples</b> section, below.
pit	For ppc_loo_pit_overlay and ppc_loo_pit_qq, optionally a vector of pre- computed PIT values that can be specified instead of y, yrep, and lw (these are all ignored if pit is specified). If not specified the PIT values are computed internally before plotting.
samples	For ppc_loo_pit_overlay, the number of data sets (each the same size as y) to simulate from the standard uniform distribution. The default is 100. The density estimate of each dataset is plotted as a thin line in the plot, with the density estimate of the LOO PITs overlaid as a thicker dark line.
	Currently unused.

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alpha, size, fatten

Arguments passed to code geoms to control plot aesthetics. For ppc\_loo\_pit\_qq and ppc\_loo\_pit\_overlay, size and alpha are passed to geom\_point and geom\_density, respectively. For ppc\_loo\_intervals, size and fatten are passed to geom\_pointrange. For ppc\_loo\_ribbon, alpha and size are passed to geom\_ribbon.

trim Passed to stat\_density.

bw, adjust, kernel, n\_dens

Optional arguments passed to density to override default kernel density estimation parameters. n\_dens defaults to 1024.

For ppc\_loo\_pit\_qq, a string that can be either "uniform" or "normal". If compare "uniform" (the default) the Q-Q plot compares computed PIT values to the standard uniform distribution. If compare="normal", the Q-Q plot compares

standardized PIT values to the standard normal distribution.

psis\_object If using **loo** version 2.0.0 or greater, an object returned by the psis function

(or by the loo function with argument save\_psis set to TRUE).

For ppc\_loo\_intervals and ppc\_loo\_ribbon, an optional integer vector in-

dicating which observations in y (and yrep) to include. Dropping observations from y and yrep manually before passing them to the plotting function will not work because the dimensions will not match up with the dimensions of psis\_object, but if all of y and yrep are passed along with subset then bayesplot can do the subsetting internally for y, yrep and psis\_object. See

the Examples section for a demonstration.

For ppc\_loo\_intervals and ppc\_loo\_ribbon, optionally a matrix of precom-

puted LOO predictive intervals that can be specified instead of yrep and lw (these are both ignored if intervals is specified). If not specified the intervals are computed internally before plotting. If specified, intervals must be a matrix with number of rows equal to the number of data points and five columns in the following order: lower outer interval, lower inner interval, median (50%),

upper inner interval and upper outer interval (column names are ignored).

prob, prob\_outer

Values between 0 and 1 indicating the desired probability mass to include in the

inner and outer intervals. The defaults are prob=0.5 and prob\_outer=0.9.

For ppc\_loo\_intervals, a string indicating how to arrange the plotted intervals. The default ("index") is to plot them in the order of the observations. The alternative ("median") arranges them by median value from smallest (left) to

largest (right).

#### Value

A ggplot object that can be further customized using the **ggplot2** package.

#### **Plot Descriptions**

ppc\_loo\_pit\_qq,ppc\_loo\_pit\_overlay The calibration of marginal predictions can be assessed using probability integral transformation (PIT) checks. LOO improves the check by avoiding the double use of data. See the section on marginal predictive checks in Gelman et al. (2013,

subset

intervals

order

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p. 152–153) and section 5 of Gabry et al. (2018) for an example of using **bayesplot** for these checks.

The LOO PIT values are asymptotically uniform (for continuous data) if the model is calibrated. The ppc\_loo\_pit\_overlay function creates a plot comparing the density of the LOO PITs (thick line) to the density estimates of many simulated data sets from the standard uniform distribution (thin lines). See Gabry et al. (2018) for an example of interpreting the shape of the miscalibration that can be observed in these plots.

The ppc\_loo\_pit\_qq function provides an alternative visualization of the miscalibration with a quantile-quantile (Q-Q) plot comparing the LOO PITs to the standard uniform distribution. Comparing to the uniform is not good for extreme probabilities close to 0 and 1, so it can sometimes be useful to set the compare argument to "normal", which will produce a Q-Q plot comparing standardized PIT values to the standard normal distribution that can help see the (mis)calibration better for the extreme values. However, in most cases we have found that the overlaid density plot (ppc\_loo\_pit\_overlay) function will provided a clearer picture of calibration problems that the Q-Q plot.

ppc\_loo\_intervals, ppc\_loo\_ribbon Similar to ppc\_intervals and ppc\_ribbon but the intervals are for the LOO predictive distribution.

#### References

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (p. 152–153)

Gabry, J., Simpson, D., Vehtari, A., Betancourt, M., and Gelman, A. (2018). Visualization in Bayesian workflow. *Journal of the Royal Statistical Society Series A*, accepted for publication. arXiv preprint: http://arxiv.org/abs/1709.01449.

Vehtari, A., Gelman, A., and Gabry, J. (2017). Practical Bayesian model evaluation using leave-one-out cross-validation and WAIC. *Statistics and Computing*. 27(5), 1413–1432. doi:10.1007/s11222-016-9696-4. arXiv preprint: http://arxiv.org/abs/1507.04544/

#### See Also

 $Other\ PPCs:\ PPC-discrete,\ PPC-distributions,\ PPC-errors,\ PPC-intervals,\ PPC-overview,\ PPC-scatterplots,\ PPC-test-statistics$ 

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```
y <- radon$log_radon
yrep <- posterior_predict(fit)</pre>
loo1 <- loo(fit, save_psis = TRUE, cores = 2)</pre>
psis1 <- loo1$psis_object</pre>
lw <- weights(psis1)</pre>
# marginal predictive check using LOO probability integral transform
color_scheme_set("orange")
ppc_loo_pit_overlay(y, yrep, lw = lw)
ppc_loo_pit_qq(y, yrep, lw = lw)
ppc_loo_pit_qq(y, yrep, lw = lw, compare = "normal")
# loo predictive intervals vs observations
keep_obs <- 1:50
ppc_loo_intervals(y, yrep, psis_object = psis1, subset = keep_obs)
color_scheme_set("gray")
ppc_loo_intervals(y, yrep, psis_object = psis1, subset = keep_obs,
                   order = "median")
## End(Not run)
```

PPC-overview

Graphical posterior predictive checking

#### **Description**

The **bayesplot** PPC module provides various plotting functions for creating graphical displays comparing observed data to simulated data from the posterior predictive distribution. See below for a brief discussion of the ideas behind posterior predictive checking, a description of the structure of this package, and tips on providing an interface to **bayesplot** from another package.

#### **Details**

The idea behind posterior predictive checking is simple: if a model is a good fit then we should be able to use it to generate data that looks a lot like the data we observed.

**Posterior predictive distribution:** To generate the data used for posterior predictive checks we simulate from the *posterior predictive distribution*. The posterior predictive distribution is the distribution of the outcome variable implied by a model after using the observed data y (a vector of outcome values), and typically predictors X, to update our beliefs about the unknown parameters  $\theta$  in the model. For each draw of the parameters  $\theta$  from the posterior distribution  $p(\theta \mid y, X)$  we generate an entire vector of outcomes. The result is an  $S \times N$  matrix of simulations, where S is the the size of the posterior sample (number of draws from the posterior distribution) and N is the number of data points in y. That is, each row of the matrix is an individual "replicated" dataset of N observations.

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**Notation:** When simulating from the posterior predictive distribution we can use either the same values of the predictors X that we used when fitting the model or new observations of those predictors. When we use the same values of X we denote the resulting simulations by  $y^{rep}$  as they can be thought of as *replications* of the outcome y rather than predictions for future observations. This corresponds to the notation from Gelman et. al. (2013) and is the notation used throughout the documentation for this package.

**Graphical posterior predictive checking:** Using the datasets  $y^{rep}$  drawn from the posterior predictive distribution, the functions in the **bayesplot** package produce various graphical displays comparing the observed data y to the replications. For a more thorough discussion of posterior predictive checking see Chapter 6 of Gelman et. al. (2013).

### **PPC** plotting functions

The plotting functions for posterior predictive checking in this package are organized into several categories, each with its own documentation:

**Distributions** Histograms, kernel density estimates, boxplots, and other plots comparing the empirical distribution of the observed data y to the distributions of individual replicated datasets (rows) in yrep.

**Test statistics** The distribution of a test statistic, or a pair of test statistics, over the replicated datasets (rows) in yrep compared to value of the statistic(s) computed from y.

**Intervals** Interval estimates of yrep with y overlaid. The x-axis variable can be optionally specified by the user (e.g. to plot against against a predictor variable or over time).

**Predictive errors** Plots of predictive errors (y -yrep) computed from y and replicated datasets (rows) in yrep. For binomial models binned error plots are also available.

**Scatterplots** Scatterplots (and similar visualizations) of the observed data y vs. individual replicated datasets (rows) in yrep, or vs. the average value of the distributions of each data point (columns) in yrep.

**Plots for discrete outcomes** PPC functions that can only be used if y and yrep are discrete. For example, rootograms for count outcomes and bar plots for ordinal, categorical, and multinomial outcomes.

**LOO predictive checks** PPC functions for predictive checks based on (approximate) leave-one-out (LOO) cross-validation.

# Providing an interface for posterior predictive checking from another package

In addition to the various plotting functions, the **bayesplot** package provides the S3 generic pp\_check. Authors of R packages for Bayesian inference are encouraged to define pp\_check methods for the fitted model objects created by their packages. See the package vignettes for more details and a simple example, and see the **rstanarm** and **brms** packages for full examples of pp\_check methods.

#### References

Gabry, J., Simpson, D., Vehtari, A., Betancourt, M., and Gelman, A. (2018). Visualization in Bayesian workflow. *Journal of the Royal Statistical Society Series A*, accepted for publication. arXiv preprint: http://arxiv.org/abs/1709.01449.

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (Ch. 6)

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#### See Also

 $Other\ PPCs:\ PPC-discrete,\ PPC-distributions,\ PPC-errors,\ PPC-intervals,\ PPC-loo,\ PPC-scatterplots,\ PPC-test-statistics$ 

PPC-scatterplots PPC scatterplots

# **Description**

Scatterplots of the observed data y vs. simulated/replicated data yrep from the posterior predictive distribution. See the **Plot Descriptions** and **Details** sections, below.

# Usage

```
ppc_scatter(y, yrep, ..., size = 2.5, alpha = 0.8)

ppc_scatter_avg(y, yrep, ..., size = 2.5, alpha = 0.8)

ppc_scatter_avg_grouped(y, yrep, group, ..., size = 2.5, alpha = 0.8)
```

## **Arguments**

У	A vector of observations. See <b>Details</b> .
yrep	An $S$ by $N$ matrix of draws from the posterior predictive distribution, where $S$ is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and $N$ is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See <b>Details</b> for additional instructions.
	Currently unused.
size, alpha	Arguments passed to geom_point to control the appearance of the points.
group	A grouping variable (a vector or factor) the same length as y. Each value in group is interpreted as the group level pertaining to the corresponding value of y.

# **Details**

For Binomial data, the plots will typically be most useful if y and yrep contain the "success" proportions (not discrete "success" or "failure" counts).

#### Value

A ggplot object that can be further customized using the ggplot2 package.

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# **Plot Descriptions**

```
ppc_scatter For each dataset (row) in yrep a scatterplot is generated showing y against that row of yrep. For this plot yrep should only contain a small number of rows.
```

ppc\_scatter\_avg A scatterplot of y against the average values of yrep, i.e., the points (mean(yrep[,n]), y[n]), where each yrep[,n] is a vector of length equal to the number of posterior draws.

ppc\_scatter\_avg\_grouped The same as ppc\_scatter\_avg, but a separate plot is generated for each level of a grouping variable.

#### References

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (Ch. 6)

#### See Also

Other PPCs: PPC-discrete, PPC-distributions, PPC-errors, PPC-intervals, PPC-loo, PPC-overview, PPC-test-statistics

#### **Examples**

```
y <- example_y_data()
yrep <- example_yrep_draws()
p1 <- ppc_scatter_avg(y, yrep)
p1
p2 <- ppc_scatter(y, yrep[20:23, ], alpha = 0.5, size = 1.5)
p2

# give x and y axes the same limits
lims <- ggplot2::lims(x = c(0, 160), y = c(0, 160))
p1 + lims
p2 + lims
group <- example_group_data()
ppc_scatter_avg_grouped(y, yrep, group, alpha = 0.7) + lims</pre>
```

# **Description**

The distribution of a test statistic T(yrep), or a pair of test statistics, over the simulated datasets in yrep, compared to the observed value T(y) computed from the data y. See the **Plot Descriptions** and **Details** sections, below.

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#### Usage

```
ppc_stat(y, yrep, stat = "mean", ..., binwidth = NULL, breaks = NULL,
  freq = TRUE)

ppc_stat_grouped(y, yrep, group, stat = "mean", ..., facet_args = list(),
  binwidth = NULL, breaks = NULL, freq = TRUE)

ppc_stat_freqpoly_grouped(y, yrep, group, stat = "mean", ...,
  facet_args = list(), binwidth = NULL, freq = TRUE)

ppc_stat_2d(y, yrep, stat = c("mean", "sd"), ..., size = 2.5, alpha = 0.7)
```

# **Arguments**

У	A vector of observations. See <b>Details</b> .
yrep	An $S$ by $N$ matrix of draws from the posterior predictive distribution, where $S$ is the size of the posterior sample (or subset of the posterior sample used to generate yrep) and $N$ is the number of observations (the length of y). The columns of yrep should be in the same order as the data points in y for the plots to make sense. See <b>Details</b> for additional instructions.
stat	A single function or a string naming a function, except for ppc_stat_2d which requires a vector of exactly two functions or function names. In all cases the function(s) should take a vector input and return a scalar test statistic. If specified as a string (or strings) then the legend will display function names. If specified as a function (or functions) then generic naming is used in the legend.
	Currently unused.
binwidth	Passed to geom_histogram to override the default binwidth.
breaks	Passed to geom_histogram as an alternative to binwidth.
freq	For histograms, freq=TRUE (the default) puts count on the y-axis. Setting freq=FALSE puts density on the y-axis. (For many plots the y-axis text is off by default. To view the count or density labels on the y-axis see the yaxis_text convenience function.)
group	A grouping variable (a vector or factor) the same length as y. Each value in

#### **Details**

facet\_args

size, alpha

For Binomial data, the plots will typically be most useful if y and yrep contain the "success" proportions (not discrete "success" or "failure" counts).

group is interpreted as the group level pertaining to the corresponding value of

Arguments passed to geom\_point to control the appearance of scatterplot points.

A named list of arguments (other than facets) passed to facet\_wrap or facet\_grid

# Value

A ggplot object that can be further customized using the ggplot2 package.

to control faceting.

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## **Plot Descriptions**

ppc\_stat A histogram of the distribution of a test statistic computed by applying stat to each dataset (row) in yrep. The value of the statistic in the observed data, stat(y), is overlaid as a vertical line.

ppc\_stat\_grouped,ppc\_stat\_freqpoly\_grouped The same as ppc\_stat, but a separate plot is generated for each level of a grouping variable. In the case of ppc\_stat\_freqpoly\_grouped the plots are frequency polygons rather than histograms.

ppc\_stat\_2d A scatterplot showing the joint distribution of two test statistics computed over the datasets (rows) in yrep. The value of the statistics in the observed data is overlaid as large point.

#### References

Gelman, A., Carlin, J. B., Stern, H. S., Dunson, D. B., Vehtari, A., and Rubin, D. B. (2013). *Bayesian Data Analysis*. Chapman & Hall/CRC Press, London, third edition. (Ch. 6)

#### See Also

Other PPCs: PPC-discrete, PPC-distributions, PPC-errors, PPC-intervals, PPC-loo, PPC-overview, PPC-scatterplots

```
y <- example_y_data()</pre>
yrep <- example_yrep_draws()</pre>
ppc_stat(y, yrep)
ppc_stat(y, yrep, stat = "sd") + legend_none()
ppc_stat_2d(y, yrep)
ppc_stat_2d(y, yrep, stat = c("median", "mean")) + legend_move("bottom")
color_scheme_set("teal")
group <- example_group_data()</pre>
ppc_stat_grouped(y, yrep, group)
color_scheme_set("mix-red-blue")
ppc_stat_freqpoly_grouped(y, yrep, group, facet_args = list(nrow = 2))
# use your own function to compute test statistics
color_scheme_set("brightblue")
q25 <- function(y) quantile(y, 0.25)</pre>
ppc_stat(y, yrep, stat = "q25") # legend includes function name
# can define the function in the 'stat' argument but then
# the legend doesn't include a function name
ppc_stat(y, yrep, stat = function(y) quantile(y, 0.25))
```

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pp_check	Posterior predictive checks (S3 generic and default method)

#### **Description**

S3 generic with simple default method. The intent is to provide a generic so authors of other R packages who wish to provide interfaces to the functions in **bayesplot** will be encouraged to include pp\_check methods in their package, preserving the same naming conventions for posterior predictive checking across many R packages for Bayesian inference. This is for the convenience of both users and developers. See the **Details** and **Examples** sections, below, and the package vignettes for examples of defining pp\_check methods.

# Usage

```
pp_check(object, ...)
## Default S3 method:
pp_check(object, yrep, fun, ...)
```

## **Arguments**

object	Typically a fitted model object. The default method, however, takes object to be a y (outcome) vector.
• • •	For the generic, arguments passed to individual methods. For the default method, these are additional arguments to pass to fun.
yrep	For the default method, a yrep matrix passed to fun.
fun	For the default method, the plotting function to call. Can be any of the PPC functions. The "ppc_" prefix can optionally be dropped if fun is specified as a string.

#### **Details**

A package that creates fitted model objects of class "foo" can include a method pp\_check.foo that prepares the appropriate inputs (y, yrep, etc.) for the **bayesplot** functions. The pp\_check.foo method may, for example, let the user choose between various plots, calling the functions from **bayesplot** internally as needed. See **Examples**, below, and the package vignettes.

# Value

The exact form of the value returned by pp\_check may vary by the class of object, but for consistency we encourage authors of methods to return the ggplot object created by one of **bayesplot**'s plotting functions. The default method returns the object returned by fun.

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#### **Examples**

```
# default method
y <- example_y_data()</pre>
yrep <- example_yrep_draws()</pre>
pp_check(y, yrep[1:50,], ppc_dens_overlay)
g <- example_group_data()</pre>
pp_check(y, yrep, fun = "stat_grouped", group = g, stat = "median")
# defining a method
x \leftarrow list(y = rnorm(50), yrep = matrix(rnorm(5000), nrow = 100, ncol = 50))
class(x) <- "foo"
pp_check.foo <- function(object, ..., type = c("multiple", "overlaid")) {</pre>
  y <- object[["y"]]</pre>
  yrep <- object[["yrep"]]</pre>
  switch(match.arg(type),
         multiple = ppc_hist(y, yrep[1:min(8, nrow(yrep)),, drop = FALSE]),
         overlaid = ppc_dens_overlay(y, yrep))
}
pp_check(x)
pp_check(x, type = "overlaid")
```

theme\_default

Default bayesplot plotting theme

#### **Description**

The theme\_default function returns the default ggplot theme used by the **bayesplot** plotting functions. See bayesplot\_theme\_set for details on setting and updating the plotting theme.

# Usage

```
theme_default(base_size = getOption("bayesplot.base_size", 12),
  base_family = getOption("bayesplot.base_family", "serif"))
```

# **Arguments**

```
base_size, base_family
```

Base font size and family (passed to theme\_bw). It is possible to set "bayesplot.base\_size" and "bayesplot.base\_family" via options to change the defaults, which are 12 and "serif", respectively.

#### Value

A ggplot theme object.

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# See Also

bayesplot-helpers for a variety of convenience functions, many of which provide shortcuts for tweaking theme elements after creating a plot.

bayesplot-colors to set or view the color scheme used for plotting.

```
class(theme_default())
bayesplot_theme_set() # defaults to setting theme_default()
x <- example_mcmc_draws()
mcmc_hist(x)

# change the default font size and family for bayesplots
bayesplot_theme_set(theme_default(base_size = 8, base_family = "sans"))
mcmc_hist(x)
mcmc_areas(x, regex_pars = "beta")

# change back
bayesplot_theme_set()
mcmc_areas(x, regex_pars = "beta")</pre>
```

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