

see: An R Package for Visualizing Statistical Models

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Summary

easystats is a collection of packages that operate in synergy to provide a consistent and intuitive syntax when working with statistical models in the R programming language (R Core Team, 2021). Most easystats packages return comprehensive numeric summaries of model parameters and performance. The see package complements these numeric summaries with a host of functions and tools to produce a range of publication-ready visualizations for model parameters, predictions, and performance diagnostics. As a core pillar of easystats, the see package helps users to use visualization for more informative, communicable and well-rounded scientific reporting.

Statement of Need

The grammar of graphics (Wilkinson, 2012), largely due to its implementation in the ggplot2 package (Wickham, 2016), has become the dominant approach to visualization in R. Building a model visualization with ggplot2 is somewhat disconnected from the model fitting and evaluation process. Generally, this process entails:

- 1. Fitting a model.
- 2. Extracting desired results from the model (e.g., model parameters and intervals, model predictions, diagnostic statistics) and arranging them into a dataframe.
- 3. Passing the results dataframe to ggplot() and specifying the graphical parameters. For example:

```
library(ggplot2)
model <- lm(mpg ~ factor(cyl) * wt, data = mtcars)
results <- fortify(model)
ggplot(results) +
  geom_point(aes(x = wt, y = mpg, color = factor(cyl))) +
  geom_line(aes(x = wt, y = .fitted, color = `factor(cyl)`))</pre>
```

A number of packages have been developed to extend ggplot2 and assist with model visualization.¹ Some of these packages provide functions for additional geoms, annotations, or common visualization types without linking them to a specific statistical analysis or

¹For a sampling of these packages, visit https://exts.ggplot2.tidyverse.org/gallery/



fundamentally changing the ggplot2 workflow (e.g., ggrepel, ggalluvial, ggridges, ggdist, ggpubr, etc.). Other ggplot2 extensions provide functions to generate publication-ready visualizations for specific types of models (e.g., metaviz, tidymv, survminer). For example, the ggstatsplot package (Patil, 2021) offers visualizations for statistical analysis of one-way factorial designs, and the plotmm package (Waggoner, 2020) supports specific types of mixture model objects.

The aim of the see package is to produce visualizations for a wide variety of models and statistical analyses in a way that is tightly linked with the model fitting process and requires minimal interruption of users' workflow. see accomplishes this aim by providing a single plot() method for objects created by the other easystats packages, such as parameters tables, modelbased predictions, performance diagnostic tests, correlation matrices, and so on. The easystats packages compute numeric results for a wide range of statistical models, and the see package acts as a visual support to the entire easystats ecosystem. As such, visualizations corresponding to all stages of statistical analysis, from model fitting to diagnostics to reporting, can be easily created using see. see plots are compatible with other ggplot2 functions for further customization (e.g., labs() for a plot title). In addition, see provides several aesthetic utilities to embellish both easystats plots and other ggplot2 plots. The result is a package that minimizes the barrier to producing high-quality statistical visualizations in R.

The central goal of *easystats* is to make the task of doing statistics in R as easy as possible. This goal is realized through intuitive and consistent syntax, consistent and transparent argument names, comprehensive documentation, informative warnings and error messages, and smart functions with sensible default parameter values. The *see* package follows this philosophy by using a single access point—the generic plot() method—for visualization of all manner of statistical results supported by *easystats*.

Features

Below we present one or two plotting methods for each *easystats* package, but many other methods are available. Interested readers are encouraged to explore the range of examples on the package website, https://easystats.github.io/see/.



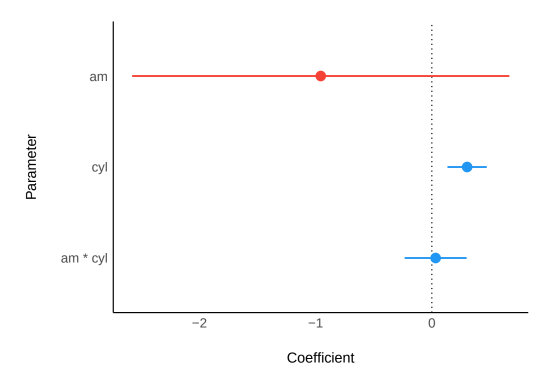
Visualizing Model Parameters

The parameters package converts summaries of regression model objects into dataframes (Lüdecke et al., 2020). The see package can take this transformed object and, for example, create a dot-and-whisker plot for the extracted regression estimates simply by passing the parameters class object to plot().

```
library(parameters)
library(see)

model <- lm(wt ~ am * cyl, data = mtcars)

plot(parameters(model))</pre>
```





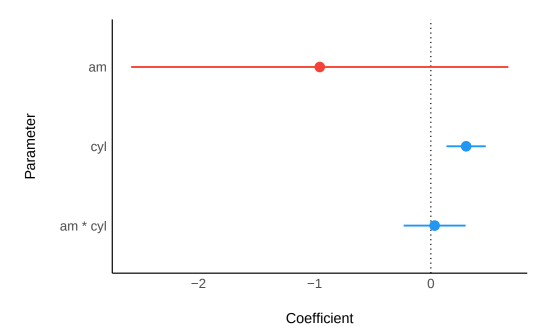
As see outputs objects of class ggplot, ggplot2 functions can be added as layers to the plot the same as with all other ggplot2 visualizations. For example, we might add a title using labs() from ggplot2.

```
library(parameters)
library(see)

model <- lm(wt ~ am * cyl, data = mtcars)

plot(parameters(model)) +
    ggplot2::labs(title = "A Dot-and-Whisker Plot")</pre>
```

A Dot-and-Whisker Plot





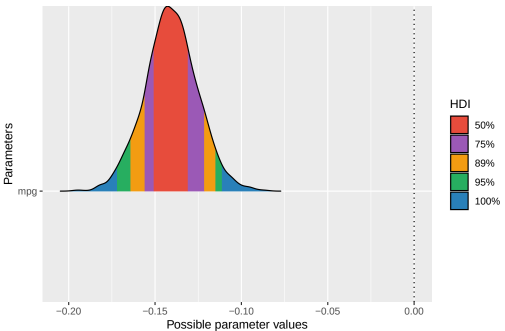
Similarly, for Bayesian regression model objects, which are handled by the bayestestR package (Makowski et al., 2019), the see package provides special plotting methods relevant for Bayesian models (e.g., Highest Density Interval, or HDI). Users can fit the model and pass the model results, extracted via bayestestR, to plot().

```
library(bayestestR)
library(rstanarm)
library(see)

model <- stan_glm(wt ~ mpg, data = mtcars, refresh = 0)
result <- hdi(model, ci = c(0.5, 0.75, 0.89, 0.95))

plot(result)</pre>
```

Highest Density Interval (HDI)





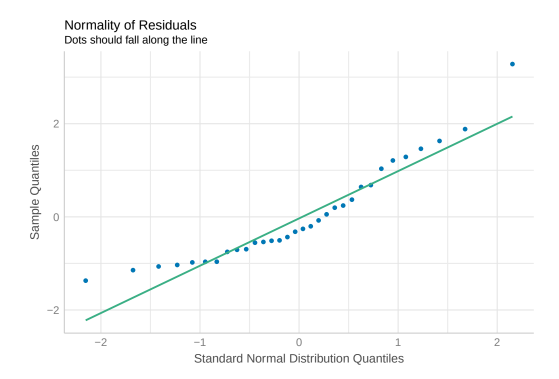
Visualizing Model Performance and Diagnostic Checks

The performance package is primarily concerned with checking regression model assumptions (Lüdecke et al., 2021). The see package offers tools to visualize these assumption checks, such as the normality of residuals. Users simply pass the fit model object to the relevant performance function (check_normality() in the example below). Then, this result can be passed to plot() to produce a ggplot2 visualization of the check on normality of the residuals.

```
library(performance)
library(see)

model <- lm(wt ~ mpg, data = mtcars)
check <- check_normality(model)

#> Warning: Non-normality of residuals detected (p = 0.016).
plot(check, type = "qq")
```





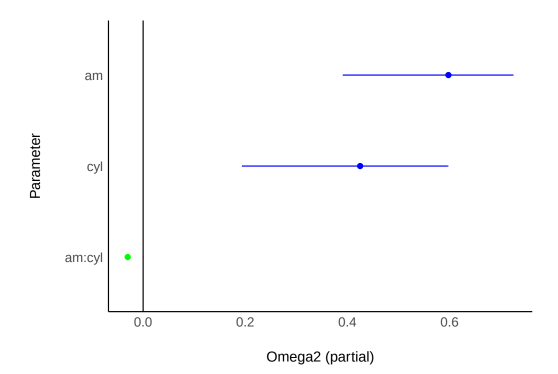
Visualizing Effect Sizes

The effectsize package computes a variety of effect size metrics for fitted models to assesses the practical importance of observed effects (Ben-Shachar et al., 2020). In conjunction with see, users are able to visualize the magnitude and uncertainty of effect sizes by passing the model object to the relevant effectsize function (omega_squared() in the following example), and then to plot().

```
library(effectsize)
library(see)

model <- aov(wt ~ am * cyl, data = mtcars)

plot(omega_squared(model))</pre>
```





Visualizing Marginal Effects

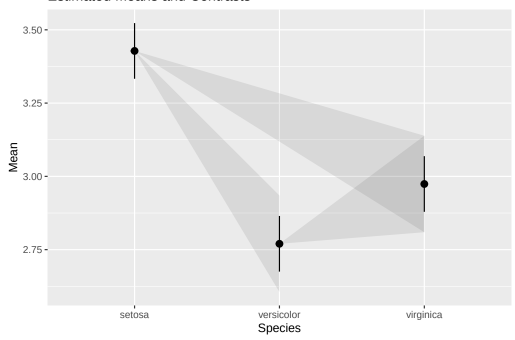
The *modelbased* package computes model-based estimates and predictions from fitted models (Makowski et al., 2020a). see provides methods to quickly visualize these model predictions. For example, to visualize group contrasts in a factorial design, users first pass their model object to the *modelbased* functions estimate_contrasts() and estimate_means(), then pass these *two* results objects to plot().

```
library(modelbased)
library(rstanarm)
library(see)

model <- lm(Sepal.Width ~ Species, data = iris)
contrasts <- estimate_contrasts(model)
means <- estimate_means(model)

plot(contrasts, means)</pre>
```

Estimated Means and Contrasts

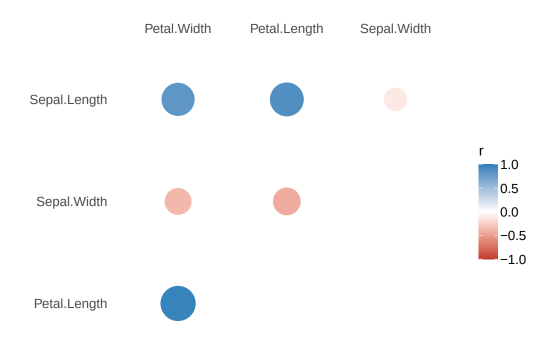




Visualizing Correlation Matrices

The correlation package provides a unified syntax and human-readable code to carry out many types of correlation analysis (Makowski et al., 2020b). A user can run summary(correlation(data)) to create a construct a correlation matrix for the variables in a dataframe. With see, this matrix can be passed to plot() to visualize these correlations in a corrgram.

```
library(correlation)
library(see)
results <- summary(correlation(iris))
plot(results)</pre>
```



Licensing and Availability

see is licensed under the GNU General Public License (v3.0), with all source code openly developed and stored at GitHub (https://github.com/easystats/see), along with a corresponding issue tracker for bug reporting and feature enhancements. In the spirit of honest and open science, we encourage requests, tips for fixes, feature updates, as well as general questions and concerns via direct interaction with contributors and developers.

Acknowledgments

see is part of the collaborative easystats ecosystem. Thus, we thank the members of easystats as well as the users.



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