

modelbased: An R package to make the most out of your statistical models through marginal means, marginal effects, and model predictions

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Statement of need

Applied statistics have historically focused on statistical tests (e.g., t-tests, correlation tests, and analyses of variances, ANOVAs), seen as most apt to provide researchers with interpretable answers to the questions they seek. These tests, however, typically rely on statistical models — the true underlying cornerstone of modern data science. The replication crisis (Camerer et al., 2018; Open Science Collaboration, 2015) and methodological (r)evolutions (Makowski & Waggoner, 2023) have underlined some of the issues with the traditional focus on statistical tests (e.g., the effacement of model assumptions, an emphasis on null-hypothesis testing, non-compatibility with more complex variance structures) and called for shifting the focus to the models themselves (Cumming, 2014).

In line with these efforts, new tools have been created to facilitate the direct usage and reporting of statistical models. For instance, the easystats collection of R packages (Lüdecke et al., 2023) has been developed to help researchers "tame, discipline, and harness" the power of statistical models. Within this framework, specific packages are dedicated to model parameters (the parameters package, Lüdecke et al., 2020), predictive performance (the performance package, Lüdecke et al., 2021) or effect importance (the effectsize package, Ben-Shachar et al., 2020).

But the models themselves pack even more usefulness!

The fundamental nature of these models—a statistical link between an outcome y and predictor variables X—enables the generation of predictions for any observed or unobserved combination of predictors. These predictions refer to expected values of the outcome for given levels of predictors of interest, making it possible to test and visualize the model's behaviour in a more meaningful and comprehensive way, and answering a broad range of research questions.

The two most popular R packages for extracting these quantities of interest from statistical models are emmeans (Lenth, 2024) and marginaleffects (Arel-Bundock et al., 2024). These packages pack an enormously rich set of features and cover (almost) all imaginable needs for post-hoc analysis of statistical models. Their power and flexibility can be intimidating for users not familiar with the underlying statistical concepts. The modelbased package, built on top of these two packages, aims to unleash this untapped potential by providing a unified interface to extract marginal means, marginal effects,



contrasts, comparisons, and model predictions from a wide range of statistical models. In line with the easystats' raison d'être, the modelbased package focuses on simplicity, flexibility, and user-friendliness to help researchers harness the full power of their models.

Key concepts

Answering research questions based on statistical models means describing the relationship between predictors of interest (also called *focal* predictors) and the outcome, as well as differences between observed groups in the sample. There are four key concepts in modelbased to achieve this: Predictions, and Marginal Means, Effects, and Contrasts.

Predictions

At a fundamental level, modelbased and similar packages leverage model predictions. These predictions can be of different types, depending on the model and the question at hand. For instance, predictions can be associated with confidence intervals (predict = "expectation") or prediction intervals (predict = "prediction"). The former corresponds to the uncertainty around the "relationship" (i.e., the conditional estimate, typically of the expectation (E[X]) according to a model's parameters), while the latter is typically larger and provides information about the range which individual observations might fall in. Moreover, for generalized linear models (GLMs), predictions can be made on the response scale (predict = "response") or the link scale (predict = "link"). This corresponds for instance to predictions in terms of probability (response scale) or log odds (link scale) for logistic regression models.

These different types of estimates can be obtained for observations in the original dataset - which is useful to assess the model's goodness-of-fit - for new data (typically a "data grid"), which is useful for visualization.

For convenience, the modelbased package includes four related functions, which mostly differ in their default arguments for data and predict¹:

- estimate_prediction(): original data, prediction intervals.
- estimate_expectation(): original data, confidence intervals.
- estimate_relation(): data grid, predictions on the response scale.
- estimate_link(): data grid, predictions on the link scale.

Marginal means, contrasts and effects

Means

The concept of "marginal" in this context refers to how non-focal predictors (i.e., those not of direct interest, for instance "adjustment" variables added to "control" for it) are treated. While predictions, as described above, fix by default non-focal variables at their reference level, marginal means compute the empirical or theoretical averages over them. These kind of predictions are a good representation of the sample, because they are not based on very specific characteristics. For example, predictions can be made for specific combinations of predictors, such as people with *high* income, while marginal means might calculate the expected outcome for an *average* observation (averaged over income).

¹These functions can become redundant if the defaults are changed. For instance, estimate_relation(..., predict = "link") is equivalent to estimate_link(...).



The modelbased package provides a simple and clear interface to extract marginal means via the estimate_means() function (with focal predictors specified using the by argument), which can be considered as the "marginal" pendant to estimate_relation().

Contrasts

The computation of these model-based quantities allow for the direct statistical comparison of the predicted outcomes across different groups or levels of a focal predictor, in the form of marginal contrasts. Rather than simply observing differences in marginal means, contrast analysis quantifies these differences and assesses their statistical significance, as well as its associated uncertainty (e.g., confidence intervals).

In modelbased, this can be achieved using the estimate_contrasts() function. Focal predictors are specified in the contrast argument. As with the other functions, further stratification or grouping can be made with the by argument (for instance, to analyze the difference between two levels of a factor alongside the values of another interacting predictor).

Effects

Finally, the same approach can be used for the effects, i.e., the parameters of the model. While predictions and marginal means can be used to better understand the *relationship* of predictors with the outcome (e.g., to estimate "the average health score for a person at the age of sixty is 80 points"), marginal *effects* evaluate the (average) *effect* of a parameter (often called "slope"), telling you that "the average effect (i.e., relationship) of age on the health score is a decrease of 5 points per year".

For the simple case of linear regression without interaction terms, the regression coefficient (slope) equals the marginal effect. However, in even slightly more complex situations (e.g., with interactions or non-linear effects), the slope is not constant across the predictor's values, and estimating the *average* slope, or marginal effect, can become useful.

Again, the modelbased package has a simple function to do so, estimate_slopes(). This function calculates the *trend* or average effect, usually for numeric predictors. The trend argument specifies the focal predictors, while by allows for further grouping.

Marginalization Types

Until this point we have discussed marginal means and effects as being "averaged" over non-focal predictors, but there are actually various ways of doing so. The estimate_means(), estimate_contrasts(), and estimate_slopes() have an estimate argument that determines how predictions are averaged ("marginalized"). The options are:

- "typical" (default): Calculates predictions for a balanced data grid representing all combinations of focal predictor levels (specified in by). For non-focal numeric predictors, it uses the mean; for non-focal categorical predictors, it averages over all the levels. This represents a "typical" observation based on the data grid and is useful for comparing groups. It answers: "What would the average outcome be for a 'typical' observation?". This is the default approach when estimating marginal means using the emmeans package.
- "average": Calculates predictions for each observation in the sample and then averages these predictions within each group defined by the focal predictors. This reflects the sample's actual distribution of non-focal predictors, not a balanced grid. It answers: "What is the predicted value for an average observation in my data?".



• "population": "Clones" each observation, creating copies with all possible combinations of focal predictor levels. It then averages the predictions across these "counterfactual" observations (non-observed permutations) within each group. This extrapolates to a hypothetical broader population, considering "what if" scenarios. It answers: "What is the predicted response for the 'average' observation in a broader possible target population?". This approach entails more assumptions about the likelihood of different combinations, but can be more apt to generalize.

Setting estimate = "average" can be useful to calculate the average expected outcome from those observations from the sample at hand. For analyses emphasizing outcome differences between groups (e.g., when computing contrasts) and particularly when causal effects are being considered, it may be beneficial to model a hypothetical population not directly represented in the sample. This approach, known as G-computation (Chatton & Rohrer, 2024), is implemented by setting estimate = "population".

Group-level estimates for Mixed Models

The modelbased package also provides the estimate_grouplevel() function to conveniently extract parameters related to random factors, which typically correspond to group-level parameters (e.g., the intercept's or slope's value for each participant). These are known as BLUPs (Best Linear Unbiased Predictions) and can be estimated in two manners:

- "random" (default): Corresponds typically to the relative deviation of each individual group from their fixed effect. As such, a coefficient close to 0 means that the participants' effect is the same as the population-level effect
- "total": Returns the absolute individual-level effects, which typically corresponds to the sum of the relative random effect with its corresponding fixed effects.

Estimating these indices using mixed models can have important benefits over an empirical approach consisting of computing raw group means, of fitting individual models to all individuals separately. In particular, it is more resilient and robust to the presence of few or missing data, and naturally applies partial-pooling - aka "shrinkage", which combines information from the group and the overall population. This means that group estimates are "pulled" towards the population-level estimate if they are more uncertain (i.e., includes less observations), in essence giving more weight to more reliable estimates. Estimates shrinkage prevents overfitting and improves generalizability (Pan & Huang, 2014).

Examples

The iris dataset contains measures in centimeters of three different species of iris flowers (setosa, versicolor, and virginica, Anderson, 1936). Imagine the following linear model in which we predict those flowers' petal width (Petal.Width) from the interaction between their petal length (Petal.Length) and their Species.

```
library(easystats)
model <- lm(Petal.Width ~ Petal.Length * Species, data = iris)
parameters::parameters(model) |>
    print(select = "minimal")
```



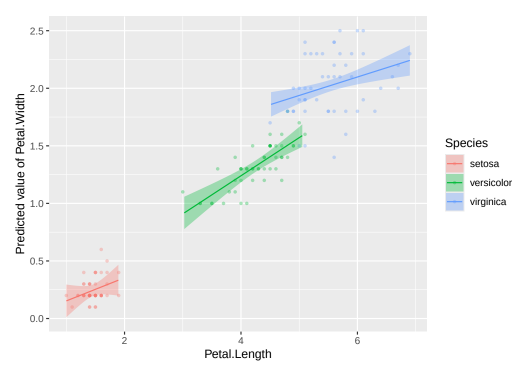


Figure 1: Scatter plot of petal length by pelal width, grouped by species

#>	Parameter		Coefficient			95% CI		p
#>								
#>	(Intercept)		-0.05		[-0.47,	0.38]		0.823
#>	Petal Length	-	0.20		[-0.09,	0.49]	1	0.170
#>	Species [versicolor]	-	-0.04		[-0.66,	0.59]	1	0.909
#>	Species [virginica]	-	1.18		[0.52,	1.84]	1	< .001
#>	Petal Length × Species [versicolor]	-	0.13		[-0.18,	0.44]	1	0.405
#>	Petal Length × Species [virginica]		-0.04	1	[-0.34,	0.26]	1	0.789

#>

#> Uncertainty intervals (equal-tailed) and p-values (two-tailed) computed
#> using a Wald t-distribution approximation.

The model's **parameters** can be challenging to interpret and do not offer us all the insights that this model actually contains.

Visualize relationship

The *modelbased* provides some basic plotting capabilities, which means that most outputs can be visualized using the plot() function. We can start by easily visualizing the relationship between our response variable and our predictors (Figure 1).

```
estimate_relation(model, by = c("Petal.Length", "Species"), length = 100) |>
plot(show_data = TRUE)
```

But what is the average value of Petal. Width for each species?



Marginal Means

The marginal means can be computed, which are the mean predictions for each level of a categorical predictor, *averaged across* all levels of other predictors (Petal.Length in this case).

However, are these different species significantly different from each other?

Marginal Contrasts

We can estimate all the pairwise contrasts between the levels of the Species factor.

```
estimate_contrasts(model, contrast = "Species")
#> Marginal Contrasts Analysis
#>
                                                         95% CI | t(144) |
#> Level1
                           | Difference |
                                             SE |
              | Level2
                                                                                 p
#> versicolor | setosa
                                   0.45 | 0.34 | [-0.22, 1.12] |
                                                                    1.34 | 0.183
                                   1.03 | 0.35 | [ 0.35, 1.72] |
#> virginica | setosa
                           1
                                                                    2.97 | 0.003
#> virginica
                                   0.58 | 0.09 | [ 0.39, 0.76] |
             | versicolor |
                                                                    6.18 \mid < .001
#>
#> Variable predicted: Petal.Width
#> Predictors contrasted: Species
#> Predictors averaged: Petal.Length (3.8)
#> p-values are uncorrected.
```

As we can see, the average difference between *versicolor* and *setosa* is not significant.

Marginal Slopes

Similarly, we can compute the marginal effect of Petal.Length (i.e., the "slope") for each species.

```
estimate_slopes(model, trend = "Petal.Length", by = "Species")
```



This shows that there is a significant positive relationship between Petal.Length and Petal.Width for all species but *setosa*.

Marginal Contrasts of Slopes

Finally, we can even compute the contrasts between the slopes of Petal.Length for each species.

```
estimate_contrasts(model, contrast = "Petal.Length", by = "Species")
```

```
#> Marginal Contrasts Analysis
#>
#> Level1
                          | Difference |
                                           SE I
                                                       95% CI | t(144) |
             | Level2
#> versicolor | setosa
                                0.13 | 0.16 | [-0.18, 0.44] |
                               -0.04 | 0.15 | [-0.34, 0.26] | -0.27 | 0.789
#> virginica | setosa
                         #> virginica | versicolor |
                               -0.17 | 0.07 | [-0.31, -0.03] | -2.41 | 0.017
#>
#> Variable predicted: Petal.Width
#> Predictors contrasted: Petal.Length
#> Predictors averaged: Petal.Length (3.8)
#> p-values are uncorrected.
```

The effect of Petal.Length on Petal.Width is significantly stronger in *virginica* compared to *versicolor*.

Conclusion

The modelbased package provides a simple and intuitive interface to extract and visualize important information contained within statistical models.

Declarations

Funding information

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Competing Interests

The authors declare no conflict of interest

Availability of data and materials (data transparency)

All data used in this paper uses data included with base R.

Code availability

The modelbased package is available at the official website (https://easystats.github.io/modelbased), on CRAN (https://cran.r-project.org/package=modelbased), and on the R-Universe (https://easystats.r-universe.dev/modelbased). The source code is available on GitHub (https://github.com/easystats/modelbased), and the package can be installed from CRAN with install.packages("modelbased") or from R-Universe with install.packages("modelbased", repos = "https://easystats.r-universe.dev").

Contributions

DM: Writing- Original draft preparation, Writing- Reviewing and Editing, Software. MSB-S, BMW, IP, RT, and DL: Writing- Reviewing and Editing, Software.

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