Package 'asympDiag'

March 27, 2025

Type Package

Title Diagnostic Tools for Asymptotic Theory

Version 0.3.1

Description Leveraging Monte Carlo simulations, this package provides tools for diagnosing regression models. It implements a parametric bootstrap framework to compute statistics, generates diagnostic envelopes to assess goodness-of-fit, and evaluates type I error control for Wald tests. By simulating data under the assumption that the model is true, it helps to identify model mis-specifications and enhances the reliability of the model inferences.

License MIT + file LICENSE

URL https://github.com/Alvaro-Kothe/asympDiag

BugReports https://github.com/Alvaro-Kothe/asympDiag/issues

Imports cli

Suggests glmmTMB, lme4, MASS, Matrix, rlang, sandwich, survival, testthat (>= 3.0.0), vctrs, withr

Config/testthat/edition 3

Encoding UTF-8

RoxygenNote 7.3.2

NeedsCompilation no

Author Álvaro Kothe [aut, cre, cph] (<https://orcid.org/0000-0003-2114-3193>), Alexandre Patriota [aut]

Maintainer Álvaro Kothe <kothe65@gmail.com>

Repository CRAN

Date/Publication 2025-03-27 22:00:06 UTC

Contents

concat_pvalues									• •	 											2
envelope	•	•						•		 			•	•		•					3

concat_pvalues

22

envelope_residual	
get_converged	· · · · · · · · · · · · · · · · · · ·
get_fixef	
get_model_response	
get_refit	
get_vcov	
parametric_bootstrap	
plot.AD_envelope	
plot.AD_pvalues	
plot_cook	
plot_ecdf_pvalue	
plot_res_vs_linear_predictor	
refit_model	
select_covariates	
simulate_wald_pvalues	

Index

concat_pvalues

Concatenate AD_pvalues object

Description

Concatenate AD_pvalues object

Usage

concat_pvalues(ld_pvalues)

Arguments

ld_pvalues list of elements of class AD_pvalues.

Value

Object of class AD_pvalues

envelope

Description

Generates QQ-plot with simulated envelope residuals.

Usage

```
envelope(
    model,
    residual_fn = envelope_residual(model),
    alpha = 0.05,
    nsim = 100,
    responses = NULL,
    no_warnings = FALSE,
    no_messages = FALSE,
    converged_only = FALSE,
    show_progress = TRUE,
    plot.it = TRUE,
    refit_fn = NULL,
    ...
)
```

Arguments

model	A model to generate responses and compute the observed residuals.
residual_fn	A function to calculate model residuals. The default is envelope_residual() for an absolute residual.
alpha	The significance level for constructing the envelope bounds. Defaults to 0.05.
nsim	The number of simulations to perform for envelope construction. Defaults to 100.
responses	An optional list of values to be used as response variables to refit the model.
no_warnings	If TRUE, ignore simulations that threw warnings.
no_messages	If TRUE, ignore simulations that shown messages.
converged_only	Use p-values from converged models only.
show_progress	Display a progress bar for the simulation iteration.
plot.it	Logical. Generate envelope plot.
refit_fn	Function to refit the model with new responses. If NULL, defaults to get_refit(model, y,).
	Extra arguments to get_refit()
	<pre>residual_fn alpha nsim responses no_warnings no_messages converged_only show_progress plot.it refit_fn</pre>

,

Details

Simulates new responses using stats::simulate() and refits the model for each vector of new responses using get_refit(). The function then computes residuals for each simulation, sorts them, and constructs envelope bands and a median line based on the quantiles of these residuals.

refit_fn is a function that supposedly compute the refit of model. Use this method if the de-fault get_refit() doesn't work. If refit_fn is NULL, it's value is defined as function(y, ...) get_refit(model, y, ...).

Value

An object of class AD_envelope, which contains the following components:

observed A vector of observed quantiles from the model residuals.

outside A logical vector indicating whether each observation falls outside the constructed envelope bounds.

lower The lower bounds of the envelope for each observation.

med The median bounds of the envelope for each observation.

upper The upper bounds of the envelope for each observation.

See Also

get_refit, simulate, rstudent, plot.AD_envelope, parametric_bootstrap()

Examples

```
fit <- lm(mpg ~ cyl, data = mtcars)</pre>
envelope(fit)
# Use pearson residuals, and plot it agains the expected normal quantiles.
env_measures <- envelope(fit,</pre>
 residual_fn = function(x) residuals.lm(x, type = "pearson"), plot.it = FALSE
)
plot(env_measures, distribution = stats::qnorm, colors = c("gray", "black"))
## Using custom refit_fn
if (require("survival")) {
 fit <- survreg(Surv(futime, fustat) ~ ecog.ps + rx, ovarian,</pre>
    dist = "exponential"
 )
 fitted_rate <- 1 / fitted(fit)</pre>
 new_responses <- replicate(100, rexp(length(fitted_rate), fitted_rate), simplify = FALSE)</pre>
 refit_surv_ovarian <- function(.y) {</pre>
    survreg(Surv(.y, fustat) ~ ecog.ps + rx, ovarian, dist = "exponential")
 }
 env_measures <- envelope(fit,</pre>
    responses = new_responses,
    residual_fn = function(x) abs(residuals(x, type = "deviance")),
    refit_fn = refit_surv_ovarian, plot.it = FALSE
```

```
)
# Absolute residuals are best shown with halfnormal quantiles
plot(env_measures, distribution = function(p) qnorm((1 + p) / 2))
}
```

envelope_residual Recommended Residuals for Envelope Plots

Description

This function returns a function that computes residuals for envelope plots. These residuals are typically absolute values to be compared against the half-normal distribution.

Usage

```
envelope_residual(object, ...)
## Default S3 method:
envelope_residual(object, ...)
## S3 method for class 'glm'
envelope_residual(object, ...)
## S3 method for class 'lm'
envelope_residual(object, ...)
```

Arguments

object	An object for which model residuals can be extracted.
	Additional arguments passed to the residual function.

Details

For objects of class glm, the default residuals are:

- Deviance residuals, except for poisson and binomial families.
- For poisson and binomial families, the residuals are rstudent(), for the case deletion residual.

For objects of class lm, the default residuals are also rstudent().

For other classes, the default is stats::residuals(), meaning no specialized recommendation is currently provided.

Value

A function that computes residuals from an object

get_converged

Description

Retrieves if the model converged.

Usage

```
get_converged(object)
```

Default S3 method: get_converged(object)

Arguments

object A model to check for convergence.

Details

The default method retrieves object\$converged. For models of class merMod it verifies if the infinity norm of the Newton-Raphson step is less than 0.0001.

Value

A boolean value.

get_fixef

Description

Extracts the fixed effects coefficients from a model.

Get fixed effects

Usage

```
get_fixef(object)
## Default S3 method:
get_fixef(object)
## S3 method for class 'merMod'
```

get_fixef(object)

S3 method for class 'glmmTMB'
get_fixef(object)

Arguments

object A model object for which fixed effects are to be retrieved.

Details

By default it calls stats::coef(). If the model class is merMod calls fixef.

Value

A numeric vector of fixed effects coefficients.

get_model_response Extract the Response Variable from a Model Object

Description

This function is designed to extract the response variable from a fitted model object.

Usage

```
get_model_response(object)
```

Arguments

object A fitted model from which the response variable should be extracted.

Details

The default method attempts to create a model frame using model.frame(). Any error encountered during this process is caught and results in a NULL return value.

Value

The response variable extracted from the model. If it couldn't be extracted, the function returns NULL.

Examples

```
lm_model <- lm(mpg ~ wt + cyl, data = mtcars)
response <- get_model_response(lm_model)
print(response)</pre>
```

get_refit

Description

Refit an existing regression model using a new response variable.

Usage

```
get_refit(object, newresp, ...)
```

Arguments

object	A model.
newresp	the new response, may be a vector or a matrix.
	other arguments passed to refit or update.

Details

This generic method refit the object with the newresp. It shold maintain the object properties by default, but some aspects of the default fitting method may be overwritten with the

Value

An updated model object, refitted using the new response variable.

See Also

stats::update(), lme4::refit()

get_vcov

Get covariance matrix

Description

Retrieves the covariance matrix for the fixed effects.

Usage

```
get_vcov(object)
## Default S3 method:
get_vcov(object)
## S3 method for class 'glmmTMB'
get_vcov(object)
```

Arguments

object A model object for which the covariance matrix is to be retrieved.

Details

By default it calls stats::vcov() to retrieve the covariances.

Value

A covariance matrix of the model object.

Description

This function performs parametric bootstrap simulations by generating new response values based on the fitted model, refitting the model on these new responses, and computing a user-defined statistic for each simulated model.

Usage

```
parametric_bootstrap(
  model,
  statistic,
  nsim,
  responses = NULL,
  refit_fn = NULL,
  show_progress = TRUE,
  simplify = TRUE,
  stat_hc = NULL,
  show_message_count = TRUE,
  show_warning_count = TRUE,
  show_not_converged_count = TRUE,
  ...
)
```

model	A fitted model object that will be used to simulate responses.
statistic	A function that computes the desired statistic from the refitted model. It must take the refitted model as an argument.
nsim	The number of simulations to perform.
responses	An optional list of values to be used as response variables to refit the model.
refit_fn	Function to refit the model with new responses. If NULL, defaults to get_refit(model, y,).

show_progress	Display a progress bar for the simulation iteration.
simplify	logical or character string; should the result be simplified to a vector, matrix or higher dimensional array if possible? If occurs any errors during the refit procedure, the results will be a list, regardless of the value of this argument.
stat_hc	A function that verifies if the computed statistic is correct. It should return nothing, just throw errors to halt execution.
show_message_co	bunt
	Show total of captured messages from refit_fn as a message. It only shows if
	the number of messages is greater than 0.
show_warning_co	bunt
	Show total of captured warnings from refit_fn as a warning. It only shows if the number of warnings is greater than 0.
show_not_conver	ged_count
	Show total of models that didn't converge as a warning. It only shows if the number of models that didn't converged is greater than 0.
	Additional arguments to be passed to refit_fn.

Details

This function implements a parametric bootstrap procedure. It generates new response values from the fitted model, refits the model for each simulated response, and computes a user-defined statistic on the refitted model. The refit function can be customized through the refit_fn argument.

If show_progress is TRUE, the progress of the simulations will be displayed using a progress bar from the cli package.

Value

A list containing the following elements:

result A list of length nsim if simplify is FALSE. Otherwise an atomic vector or matrix or list of length nsim.

responses The list of simulated response values.

simulation_warning A logical vector indicating whether a warning occurred during model refitting for each simulation.

converged A logical vector indicating whether the model refit converged for each simulation.

Examples

```
model <- lm(mpg ~ wt + hp, data = mtcars)
statistic <- function(model) coef(model)[["wt"]]
bootstrap_results <- parametric_bootstrap(model, statistic, nsim = 100)
wt_coefs <- Reduce(c, bootstrap_results$result)
summary(wt_coefs)</pre>
```

Description

Plot AD_envelope

Usage

```
## S3 method for class 'AD_envelope'
plot(
    x,
    colors = getOption("asympDiag.plot.AD_envelope.colors"),
    xlab = "Expected quantiles",
    ylab = "Observed quantiles",
    distribution = function(p) stats::qnorm((1 + p)/2),
    ylim = base::range(c(x$observed, x$lower, x$upper), na.rm = TRUE, finite = TRUE),
    ...
)
```

Arguments

х	AD_envelope object, usually the result of envelope()
colors	Vector of length 2, with color for points outside and inside the envelope band, respectively.
xlab	The label for the x-axis.
ylab	The label for the y-axis.
distribution	quantile function for reference theoretical distribution.
ylim	the y limits of the plot.
	extra arguments passed to graphics::plot

Details

Create envelope plot. The expected quantile, by default, is from a half-normal distribution, as the default residuals from envelope_residual() are absolute. You may replace it with the distribution argument.

Value

No return value, called for side effects

plot.AD_pvalues

Description

This function creates several plots with the empirical cumulative distribution of the p-values obtained through simulation.

Usage

```
## S3 method for class 'AD_pvalues'
plot(
 х,
 which = seq_len(length(x$test_coefficients) + 1),
 caption = as.list(paste("ECDF of", c(names(x$test_coefficients), "all coefficients"))),
  ks_test = TRUE,
  signif = c(0.01, 0.05, 0.1),
  discrepancy_tol = 0.1,
  plot_uniform = TRUE,
  uniform_legend = TRUE,
  converged_only = FALSE,
  no_warnings = FALSE,
  no_messages = FALSE,
  ylab = "Empirical cumulative distribution",
  xlab = "p-value",
  ...,
 ask = prod(graphics::par("mfcol")) < length(which) && grDevices::dev.interactive()</pre>
)
```

х	AD_pvalues object, usually the result of simulate_wald_pvalues()
which	A vector specifying the indices of coefficients to plot. If index is bigger than the number of coefficients it plots the joint p_value.
caption	A character vector or a list with caption for each plot. If it's a list, the list index must match the coefficient index used by which. If it's a vector, it's values are used in order.
ks_test	If TRUE inserts Kolmogorov-Smirnov p-value in the graphic.
signif	Points to verify discrepancy.
discrepancy_to]	1
	Threshold to consider point discrepant.
plot_uniform	Logical. If TRUE, plot uniform distribution.
uniform_legend	Logical. If TRUE, a legend is added to the plot to distinguish between the p-value and $U(0, 1)$ curves. Defaults to TRUE.
converged_only	Use p-values from converged models only.

plot_cook

no_warnings	If TRUE, ignore simulations that threw warnings.
no_messages	If TRUE, ignore simulations that shown messages.
ylab	The label for the y-axis. Defaults to "Empirical cumulative distribution".
xlab	The label for the x-axis. Defaults to "p-value".
	extra arguments passed to graphics::plot
ask	Logical. If TRUE, the user is prompted before each plot. Defaults to TRUE if in an interactive session and the number of plots is greater than the available space; otherwise, FALSE.

Details

If the asymptotic approximation is valid the distribution of the p-values should be close to an uniform distribution. Discrepancies are highlighted, by default it verifies the significance on the most commonly used significance values are 0.01, 0.05 and 0.10.

The reported KS (Kolmogorov-Smirnov) test is the result of the "two-sided" stats::ks.test() function comparing the observed p-values distribution with the uniform. The test may reject the KS test due to few simulations, make sure that the lines shown in the plot are smooth before drawing any conclusions.

Value

A vector of joint p-values for all coefficients.

Examples

model <- lm(mpg ~ wt + hp, data = mtcars)
p_values_ld <- simulate_wald_pvalues(model, n_sim = 100)
plot(p_values_ld)</pre>

plot_cook

Plot Cook's distances

Description

Plot Cook's distances

Usage

```
plot_cook(
   model,
   n_highlights = 0,
   cut = FALSE,
   xlab = "Index",
   ylab = "Cook's distance",
   ...
)
```

Arguments

model	Model with cooks.distance() method
n_highlights	The number of observations with the highest Cook's distance to highlight on the plot. Defaults to 0 (no highlights).
cut	Logical. If TRUE, adds a cutoff line at the mean plus four times the standard deviation of Cook's distance. Defaults to FALSE.
xlab	The label for the x-axis. Defaults to "Index".
ylab	The label for the y-axis. Defaults to "Cook's distance".
	Further arguments for graphics::plot()

Value

An invisible object representing Cook's distance values.

See Also

cooks.distance, plot

Examples

```
fit <- lm(mpg ~ cyl, data = mtcars)
plot_cook(fit)
plot_cook(fit, n_highlights = 2)</pre>
```

plot_ecdf_pvalue Plot Empirical Cumulative Distribution Function (ECDF) of p-values

Description

Plot Empirical Cumulative Distribution Function (ECDF) of p-values

Usage

```
plot_ecdf_pvalue(
  p_values,
  ks_test = TRUE,
  signif = c(0.01, 0.05, 0.1),
  discrepancy_tol = 0.1,
  plot_uniform = TRUE,
  uniform_legend = TRUE,
  main = "",
  ylab = "Empirical cumulative distribution",
  xlab = "p-value",
  ...
)
```

Arguments

p_values	vector of p-values
ks_test	If TRUE inserts Kolmogorov-Smirnov p-value in the graphic.
signif	Points to verify discrepancy.
discrepancy_to	1
	Threshold to consider point discrepant.
plot_uniform	Logical. If TRUE, plot uniform distribution.
uniform_legend	Logical. If TRUE, a legend is added to the plot to distinguish between the p-value and $U(0, 1)$ curves. Defaults to TRUE.
main	main caption passed to plot
ylab	The label for the y-axis. Defaults to "Empirical cumulative distribution".
xlab	The label for the x-axis. Defaults to "p-value".
	extra arguments passed to graphics::plot

Value

No return value, called for side effects

plot_res_vs_linear_predictor

Description

Plot Residuals against Linear Predictor

Usage

```
plot_res_vs_linear_predictor(
  model,
  residual_fn = stats::rstandard,
  xlab = "Linear Predictor",
  ylab = "Standardized deviance residuals",
  ...
)
```

model	Model with methods predict() or fitted()
residual_fn	A function to calculate model residuals. The default is $\texttt{stats}::\texttt{rstandard}$.
xlab	The label for the x-axis. Defaults to "Linear Predictor".
ylab	The label for the y-axis. Defaults to "Standardized deviance residuals".
	Extra arguments to residual_fn and plot().

Plot Residuals against Linear Predictor

Details

If the model was fitted using the glm() function, it will use the predict() method with type = link, otherwise, it will use the fitted() method.

Value

An invisible list containing the linear predictor (x) and standardized deviance residuals (y).

Examples

```
fit <- lm(mpg ~ cyl, data = mtcars)
plot_res_vs_linear_predictor(fit)
plot_res_vs_linear_predictor(fit, residual_fn = rstudent)
plot_res_vs_linear_predictor(fit, residual_fn = residuals)
glm_fit <- glm(cyl ~ mpg, family = poisson(), data = mtcars)
plot_res_vs_linear_predictor(glm_fit)
plot_res_vs_linear_predictor(glm_fit, type = "pearson")</pre>
```

refit_model Refit Model

Description

Refit a model with a new response.

Usage

refit_model(object, newresp, ...)

Arguments

object	A model.
newresp	the new response, may be a vector or a matrix.
	other arguments passed to refit or update.

Details

This function uses newresp to refit object replacing its old response variable. If the class is merMod it uses refit, otherwise uses stats::update().

The default method tries to update the model response using it's stats::model.frame(), if it errors it tries to update the model by inserting the newresp directly into the object formula.

select_covariates

Value

A model with same class as object.

select_covariates Select covariates

Description

Select covariates

Usage

```
select_covariates(
   model,
   threshold = 0.15,
   direction = c("both", "backward", "forward"),
   addable_coefs = names(get_fixef(model)),
   measure_fn = function(x) summary(x)[["coefficients"]][, 4],
   measure_one_at_time = FALSE,
   minimize_only = FALSE,
   max_steps = 1000,
   return_step_results = FALSE,
   do_not_remove = c("(Intercept)"),
   ...
)
```

	model	A model with stats::update(), stats::coef() methods.
	threshold	Value threshold to remove variable. It can be a fixed value or a function. The variable is removed if measure_fn(model) > threshold and added if measure_fn(model) <= threshold.
	direction	The direction of variable selection. Options include "backward", "forward", or "both". Defaults to "both".
	addable_coefs	A vector of coefficients that can be added during forward selection. Defaults to all coefficients in the model.
	measure_fn	Function with model as argument and returns values to be used by threshold. It can also compare two models, where during forward step it calls measure_fn(candidate_model, current_selected_model) and during backward step it calls measure_fn(current_selected_model, candidate_model). Defaults to the p-value from the summary of the coefficients.
<pre>measure_one_at_time</pre>		time
		Boolean indicating to apply measure_fn to each variable individually during forward and backward steps. Set this option to TRUE if measure_fn returns an atomic value, for example if measure_fn is AIC.

minimize_only	Logical indicating that during backward model update it should minimize the measure_fn instead of maximize it.	
max_steps	The maximum number of steps for the variable selection process. Defaults to 1000.	
return_step_results		
	Logical. If TRUE, the function returns a list containing the final fitted model and a log of the selection steps. Defaults to FALSE.	
do_not_remove	A character vector specifying variables that should not be removed during back- ward selection. Defaults to "(Intercept)".	
	Extra arguments to stats::update().	

Value

A fitted model with selected covariates based on the variable selection process. If return_step_results is TRUE, a list containing the final fitted model and a log of the selection steps is returned.

Examples

```
model <- lm(mpg ~ ., data = mtcars)</pre>
select_covariates(model)
## measure_fn with two parameters
lrt <- function(model1, model2) {</pre>
  lrt_stat <- 2 * (logLik(model1)[1L] - logLik(model2)[1L])</pre>
  return(1 - pchisq(lrt_stat, 1))
}
select_covariates(model, measure_fn = lrt)
## AICc selection
AICc <- function(model) {</pre>
  loglike <- logLik(model)</pre>
  df <- attr(loglike, "df")</pre>
  nobs <- attr(loglike, "nobs")</pre>
  aic <- -2 * as.numeric(loglike) + 2 * df</pre>
  aicc <- aic + (2 * (df^2) + 2 * df) / (nobs - df - 1)
  return(aicc)
}
selection <- select_covariates(model,</pre>
  measure_fn = AICc,
  threshold = AICc,
  measure_one_at_time = TRUE,
  minimize_only = TRUE,
  direction = "both",
  data = mtcars
)
```

simulate_wald_pvalues Generate Wald test P-Values with Monte Carlo Simulations

Description

This function performs Monte Carlo simulations to generate p-values for model coefficients by refitting the model with new simulated responses and computing the Wald test statistics for each simulation. It's standard behavior verify if the type I error from Wald tests are under control, considering the provided model as "true", i.e., the model assumptions are valid. It supports univariate and joint Wald tests, using chi-squared distributions to calculate p-values.

Usage

```
simulate_wald_pvalues(
   model,
   nsim = 1000,
   responses = NULL,
   test_coefficients = NULL,
   refit_fn = NULL,
   coef_fn = get_fixef,
   vcov_fn = get_vcov,
   show_progress = TRUE,
   plot.it = TRUE,
   ...
)
```

model	A fitted model object that will be used to simulate responses.		
nsim	The number of simulations to perform.		
responses	An optional list of values to be used as response variables to refit the model.		
test_coefficien	est_coefficients		
	Numeric vector. A vector with values to be used to compute the test statistic. It should be the coefficients that was used to compute the fitted values of the response. If NULL defaults to coef_fn(model).		
refit_fn	Function to refit the model with new responses. If NULL, defaults to get_refit(model, y,).		
coef_fn	Function that retrieves the coefficients of the model.		
vcov_fn	Function that computes the variance-covariance matrix for the models adjusted in the simulations.		
show_progress	Display a progress bar for the simulation iteration.		
plot.it	Logical. Generate ecdf plot for joint Wald test.		
	Additional arguments to be passed to refit_fn.		

Details

If responses is provided, the function refits the model with these new response vectors. Otherwise, it generates new responses with stats::simulate().

For each new response calls get_refit() to generate a new model with the new response. It gets the fixed effects and the variance and covariance matrix with get_fixef() and get_vcov().

Each simulated model is refitted using the specified refit_fn (or the default refit function) and the fixed effects coefficients and variance-covariance matrix are extracted using coef_fn and vcov_fn, respectively. The univariate Wald test is computed from the Wald statistic for each coefficient, while the joint Wald test uses the inverse variance-covariance matrix to compute a Wald statistic for the test_coefficients. P-values are calculated from a chi-squared distribution with appropriate degrees of freedom.

Value

An object of class AD_pvalues, which contains the following components:

test_coefficients Vector of coefficients being tested.

pvalues_matrix Matrix of p-values where each column corresponds to a simulation and each row corresponds to a coefficient.

pvalues_joint Vector containing the joint p-values obtained from each simulation.

simulation_fixef List of fixed effect coefficient estimates from each simulation.

simulation_vcov List of covariance matrices estimated from each simulation.

simulation_warning Vector of boolean indicating if a simulation threw a warning.

converged Logical vector indicating whether model refitting converged for each simulation.

responses Simulated responses used for refitting the model.

See Also

plot.AD_pvalues() for plotting.

Examples

```
# from help("glm")
counts <- c(18, 17, 15, 20, 10, 20, 25, 13, 12)
outcome <- gl(3, 1, 9)
treatment <- gl(3, 3)
model <- glm(counts ~ outcome + treatment, family = poisson())
new_responses <- replicate(100, MASS::rnegbin(fitted.values(model), theta = 4.5), simplify = FALSE)
simulate_wald_pvalues(model, responses = new_responses, nsim = 100)
## Using custom refit_fn
if (require("survival")) {
    fit <- survreg(Surv(futime, fustat) ~ ecog.ps + rx, ovarian,
        dist = "exponential"
    )
    fitted_rate <- 1 / fitted(fit)
    new_responses <- replicate(100, rexp(length(fitted_rate), fitted_rate), simplify = FALSE)</pre>
```

20

```
refit_surv_ovarian <- function(.y) {
   survreg(Surv(.y, fustat) ~ ecog.ps + rx, ovarian, dist = "exponential")
   simulate_wald_pvalues(fit, responses = new_responses, refit_fn = refit_surv_ovarian)
}</pre>
```

Index

concat_pvalues, 2 cooks.distance, 14 cooks.distance(), 14 envelope, 3 envelope(), 11 envelope_residual, 5 envelope_residual(), 3, 11 fitted(), 15, 16 get_converged, 6 get_fixef, 6 get_fixef(), 20 get_model_response, 7 get_refit, 4, 8 get_refit(), 3, 4, 20 get_vcov, 8 get_vcov(), 20 glm(), 16 graphics::plot, 11, 13, 15 graphics::plot(), 14 lme4::refit(), 8 model.frame(),7 parametric_bootstrap, 9 parametric_bootstrap(), 4 plot, *14*, *15* plot(), 15 plot.AD_envelope, 4, 11 plot.AD_pvalues, 12 plot.AD_pvalues(), 20 plot_cook, 13 plot_ecdf_pvalue, 14 plot_res_vs_linear_predictor, 15 predict(), 15, 16 refit_model, 16

refit_model, 10 rstudent, 4 rstudent(), 5

```
select_covariates, 17
simulate, 4
simulate_wald_pvalues, 19
simulate_wald_pvalues(), 12
stats::coef(), 7, 17
stats::ks.test(), 13
stats::model.frame(), 16
stats::residuals(), 5
stats::simulate(), 4, 20
stats::update(), 8, 16–18
stats::vcov(), 9
```