# Package 'RoBTT'

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**Title** Robust Bayesian T-Test

Version 1.3.1

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Description An implementation of Bayesian model-averaged t-tests that allows users to draw inferences about the presence versus absence of an effect, variance heterogeneity, and potential outliers. The 'RoBTT' package estimates ensembles of models created by combining competing hypotheses and applies Bayesian model averaging using posterior model probabilities. Users can obtain model-averaged posterior distributions and inclusion Bayes factors, accounting for uncertainty in the data-generating process (Maier et al., 2024, <doi:10.3758/s13423-024-02590-5>). The package also provides a truncated likelihood version of the model-averaged t-test, enabling users to exclude potential outliers without introducing bias (Godmann et al., 2024, <doi:10.31234/osf.io/j9f3s>). Users can specify a wide range of informative priors for all parameters of interest. The package offers convenient functions for summary, visualization, and fit diagnostics.

```
URL https://fbartos.github.io/RoBTT/
```

```
BugReports https://github.com/FBartos/RoBTT/issues
```

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RdMacros Rdpack

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## **Description**

RoBTT: Bayesian model-averaged t-test extends the Bayesian t-test by incorporating inference about heterogeneity of variances and outliers.

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#### User guide

See Maier et al. (2024) for details regarding the RoBTT methodology.

More details regarding customization of the Bayesian model-averaged t-test are provided in **Introduction to RoBTT** vignette. Please, use the "Issues" section in the GitHub repository to ask any further questions.

#### Author(s)

František Bartoš < f.bartos 96@gmail.com>

#### References

Maier M, Bartoš F, Quintana DS, van den Bergh D, Marsman M, Ly A, Wagenmakers E (2024). "Model-averaged Bayesian t-tests." *Psychonomic Bulletin & Review*. doi:10.3758/s13423024-025905.

#### See Also

Useful links:

- https://fbartos.github.io/RoBTT/
- Report bugs at https://github.com/FBartos/RoBTT/issues

check\_RoBTT

Check fitted 'RoBTT' object for errors and warnings

#### **Description**

Checks fitted 'RoBTT' object for warnings and errors and prints them to the console.

#### Usage

```
check_RoBTT(fit)
```

#### **Arguments**

fit

a fitted 'RoBTT' object.

#### Value

check\_RoBTT returns a vector of error and warning messages.

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check\_setup

Prints summary of "RoBTT" ensemble implied by the specified priors

#### Description

check\_setup prints summary of "RoBTT" ensemble implied by the specified prior distributions. It is useful for checking the ensemble configuration prior to fitting all of the models.

#### Usage

```
check_setup(
  prior_delta = prior(distribution = "cauchy", parameters = list(location = 0, scale = sqrt(2)/2)),
  prior_rho = prior(distribution = "beta", parameters = list(alpha = 1, beta = 1)),
  prior_nu = prior(distribution = "exp", parameters = list(rate = 1)),
  prior_delta_null = prior(distribution = "spike", parameters = list(location = 0)),
  prior_rho_null = prior(distribution = "spike", parameters = list(location = 0.5)),
  prior_nu_null = prior_none(),
  prior_mu = NULL,
  prior_sigma2 = NULL,
  truncation = NULL,
  models = FALSE,
  silent = FALSE
```

#### **Arguments**

prior\_delta prior distributions for the effect size delta parameter that will be treated as belonging to the alternative hypothesis. Defaults to prior(distribution = "Cauchy", parameters = list(location = 0, scale = sqrt(2)/2)). prior distributions for the precision allocation rho parameter that will be treated prior\_rho as belonging to the alternative hypothesis. Defaults to prior(distribution = "beta", parameters = list(alpha = 1, beta = 1)). prior distribution for the degrees of freedom + 2 nu parameter that will be treated prior\_nu as belonging to the alternative hypothesis. Defaults to prior(distribution = "exp", parameters = list(rate = 1)) if no truncation is specified. If truncation is specified, the default is NULL (i.e., use only normal likelihood). prior\_delta\_null prior distribution for the delta parameter that will be treated as belonging to the null hypothesis. Defaults to point distribution with location at 0 (prior(distribution = "point", parameters = list(location = 0))). prior\_rho\_null prior distribution for the rho parameter that will be treated as belonging to the null hypothesis. Defaults to point distribution with location at 0.5 (prior(distribution = "point", parameters = list(location = 0.5))). prior distribution for the nu parameter that will be treated as belonging to the prior\_nu\_null null hypothesis. Defaults to prior\_none ( (i.e., normal likelihood)).

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prior\_mu

prior distribution for the grand mean parameter. Defaults to NULL which sets Jeffreys prior for the grand mean in case of no truncation or an unit Cauchy prior distributions for the grand mean in case of truncation (which greatly improves sampling efficiency).

prior\_sigma2

prior distribution for the grand variance parameter. Defaults to NULL which sets Jeffreys prior for the variance in case of no truncation or an exponential prior distribution for the variance in case of truncation (which greatly improves sampling efficiency).

truncation

an optional list specifying truncation applied to the data. Defaults to NULL, i.e., no truncation was applied and the full likelihood is applied. Alternative the truncation can be specified via a named list with:

- "x" where x is a vector of two values specifying the lower and upper truncation points common across the groups
- "x1" and "x2" where x1 is a vector of two values specifying the lower and upper truncation points for the first group and x2 is a vector of two values specifying the lower and upper truncation points for the second group.
- "sigma" where sigma corresponds to the number of standard deviations from the common mean where the truncation points should be set.
- "sigma1" and "sigma2" where sigma1 corresponds to the number of standard deviations from the mean of the first group where the truncation points should be set and sigma2 corresponds to the number of standard deviations from the mean of the second group where the truncation points should be set.

models should the models' details be printed.

silent do not print the results.

#### Value

check\_setup invisibly returns list of summary tables.

#### See Also

RoBTT()

diagnostics

Checks a fitted RoBTT object

#### **Description**

diagnostics creates visual checks of individual models convergence. Numerical overview of individual models can be obtained by summary(object, type = "models", diagnostics = TRUE), or even more detailed information by summary(object, type = "individual").

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#### Usage

```
diagnostics(
  fit,
  parameter,
  type,
  show_models = NULL,
  lags = 30,
  title = is.null(show_models) | length(show_models) > 1,
)
diagnostics_autocorrelation(
  fit,
  parameter = NULL,
  show_models = NULL,
  lags = 30,
  title = is.null(show_models) | length(show_models) > 1,
)
diagnostics_trace(
  fit,
  parameter = NULL,
  show_models = NULL,
  title = is.null(show_models) | length(show_models) > 1,
)
diagnostics_density(
  fit,
  parameter = NULL,
  show_models = NULL,
  title = is.null(show_models) | length(show_models) > 1,
)
```

#### **Arguments**

fit a fitted RoBTT object

parameter a parameter to be plotted. Either "delta", "rho", "nu", "mu", or "sigma".

type type of MCMC diagnostic to be plotted. Options are "chains" for the chains' trace plots, "autocorrelation" for autocorrelation of the chains, and "densities" for the overlaying densities of the individual chains. Can be abbreviated to first letters.

show\_models MCMC diagnostics of which models should be plotted. Defaults to NULL which

MCMC diagnostics of which models should be plotted. Defaults to NULL which plots MCMC diagnostics for a specified parameter for every model that is part of the ensemble.

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```
lags number of lags to be shown for type = "autocorrelation". Defaults to 30.
title whether the model number should be displayed in title. Defaults to TRUE when more than one model is selected.
additional arguments to be passed to par if plot_type = "base".
```

#### **Details**

The visualization functions are based on stan\_plot function and its color schemes.

#### Value

diagnostics returns either NULL if plot\_type = "base" or an object/list of objects (depending on the number of parameters to be plotted) of class 'ggplot2' if plot\_type = "ggplot2".

#### See Also

```
RoBTT(), summary.RoBTT()
```

#### **Examples**

```
## Not run:
# using the example data from Darwin
data("fertilization", package = "RoBTT")
fit <- RoBTT(</pre>
 x1
       = fertilization$Self,
 x2
          = fertilization$Crossed,
 prior_delta = prior("cauchy", list(0, 1/sqrt(2))),
 prior_rho = prior("beta", list(3, 3)),
 seed
             = 1,
             = 1,
 chains
             = 1000,
 warmup
             = 2000,
 iter
 control
             = set_control(adapt_delta = 0.95)
### ggplot2 version of all of the plots can be obtained by adding 'model_type = "ggplot"
# diagnostics function allows to visualize diagnostics of a fitted RoBTT object, for example,
# the trace plot for the mean parameter in each model model
diagnostics(fit, parameter = "delta", type = "chain")
# in order to show the trace plot only for the 11th model, add show_models parameter
diagnostics(fit, parameter = "delta", type = "chain", show_models = 11)
# furthermore, the autocorrelations
diagnostics(fit, parameter = "delta", type = "autocorrelation")
# and overlying densities for each plot can also be visualize
diagnostics(fit, parameter = "delta", type = "densities")
## End(Not run)
```

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fertilization

Height of 15 plant pairs collected by Darwin

# Description

The data set contains heights of self fertilized and cross fertilized plants as coded by Darwin.

# Usage

fertilization

#### **Format**

A data.frame with 2 columns and 15 observations.

#### Value

a data.frame.

interpret

Interprets results of a 'RoBTT' model.

# Description

interpret creates a brief textual summary of a fitted 'RoBTT' object.

#### Usage

```
interpret(object)
```

# Arguments

object

a fitted 'RoBTT' object

#### Value

interpret returns a character.

is.RoBTT

is.RoBTT

Reports whether x is a 'RoBTT' object

#### **Description**

Reports whether x is a 'RoBTT' object

# Usage

```
is.RoBTT(x)
```

#### **Arguments**

Χ

an object to test

#### Value

is.RoBTT returns a boolean.

plot.RoBTT

Plots a fitted 'RoBTT' object

# Description

plot.RoBTT allows to visualize different "RoBTT" object parameters in various ways. See type for the different model types.

# Usage

```
## S3 method for class 'RoBTT'
plot(
    x,
    parameter = "mu",
    transform_rho = FALSE,
    conditional = FALSE,
    plot_type = "base",
    prior = FALSE,
    dots_prior = NULL,
    ...
)
```

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#### **Arguments**

a fitted 'RoBTT' object Х parameter a parameter to be plotted. Defaults to "delta" (for the effect size). The additional options are "rho" (for the heterogeneity), "nu" (for the degrees of freedom). transform\_rho whether rho parameter should be translated into log standard deviation ratio conditional whether conditional estimates should be plotted. Defaults to FALSE which plots the model-averaged estimates. plot\_type whether to use a base plot "base" or ggplot2 "ggplot" for plotting. Defaults to "base". prior whether prior distribution should be added to figure. Defaults to FALSE. list of additional graphical arguments to be passed to the plotting function of the dots\_prior prior distribution. Supported arguments are 1wd, 1ty, col, and col. fill, to adjust the line thickness, line type, line color, and fill color of the prior distribution respectively. list of additional graphical arguments to be passed to the plotting function. Supported arguments are lwd, lty, col, col.fill, xlab, ylab, main, xlim, ylim to adjust the line thickness, line type, line color, fill color, x-label, y-label, title, x-axis range, and y-axis range respectively.

#### Value

plot.RoBTT returns either NULL if plot\_type = "base" or an object object of class 'ggplot2' if plot\_type = "ggplot2".

#### See Also

#### RoBTT()

#### **Examples**

```
## Not run:
data("fertilization", package = "RoBTT")
fit <- RoBTT(</pre>
 х1
           = fertilization$Self,
           = fertilization$Crossed,
 prior_delta = prior("cauchy", list(0, 1/sqrt(2))),
 prior_rho
             = prior("beta", list(3, 3)),
              = 1,
 seed
              = 1,
 chains
              = 1000,
 warmup
 iter
              = 2000,
 control
              = set_control(adapt_delta = 0.95)
)
# plot the model-averaged effect size estimate
plot(fit, parameter = "delta")
```

print.RoBTT

```
# plot prior and posterior of the conditional effect size estimate
plot(fit, parameter = "delta", conditional = TRUE, prior = TRUE)
## End(Not run)
```

print.RoBTT

Prints a fitted 'RoBTT' object

#### **Description**

Prints a fitted 'RoBTT' object

#### Usage

```
## S3 method for class 'RoBTT'
print(x, ...)
```

#### **Arguments**

x a fitted 'RoBTT' object.
... additional arguments.

#### Value

print.RoBTT invisibly returns the print statement.

#### See Also

RoBTT()

print.summary.RoBTT

Prints summary object for 'RoBTT' method

#### **Description**

Prints summary object for 'RoBTT' method

#### Usage

```
## S3 method for class 'summary.RoBTT' print(x, ...)
```

# Arguments

```
x a summary of a 'RoBTT' object
```

... additional arguments

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#### Value

print.summary.RoBTT invisibly returns the print statement.

#### See Also

```
RoBTT()
```

prior

Creates a prior distribution

#### **Description**

prior creates a prior distribution. The prior can be visualized by the plot function.

#### Usage

```
prior(
  distribution,
  parameters,
  truncation = list(lower = -Inf, upper = Inf),
  prior_weights = 1
)
```

#### **Arguments**

distribution

name of the prior distribution. The possible options are

"point" for a point density characterized by a location parameter.

"normal" for a normal distribution characterized by a mean and sd parameters.

- "lognormal" for a lognormal distribution characterized by a meanlog and sdlog parameters.
- "cauchy" for a Cauchy distribution characterized by a location and scale parameters. Internally converted into a generalized t-distribution with df =
- "t" for a generalized t-distribution characterized by a location, scale, and df parameters.
- "gamma" for a gamma distribution characterized by either shape and rate, or shape and scale parameters. The later is internally converted to the shape and rate parametrization
- "invgamma" for an inverse-gamma distribution characterized by a shape and scale parameters. The JAGS part uses a 1/gamma distribution with a shape and rate parameter.
- "beta" for a beta distribution characterized by an alpha and beta parameters.
- "exp" for an exponential distribution characterized by either rate or scale parameter. The later is internally converted to rate.
- "uniform" for a uniform distribution defined on a range from a to b

prior\_none 13

parameters list of appropriate parameters for a given distribution.

truncation list with two elements, lower and upper, that define the lower and upper trun-

cation of the distribution. Defaults to list(lower = -Inf), upper = Inf). The

truncation is automatically set to the bounds of the support.

prior\_weights prior odds associated with a given distribution. The value is passed into the

model fitting function, which creates models corresponding to all combinations of prior distributions for each of the model parameters and sets the model priors

odds to the product of its prior distributions.

#### Value

prior and prior\_none return an object of class 'prior'. A named list containing the distribution name, parameters, and prior weights.

#### See Also

```
plot.prior(), Normal, Lognormal, Cauchy, Beta, Exponential, LocationScaleT, InvGamma.
```

#### **Examples**

```
# create a standard normal prior distribution
p1 <- prior(distribution = "normal", parameters = list(mean = 1, sd = 1))
# create a half-normal standard normal prior distribution
p2 <- prior(distribution = "normal", parameters = list(mean = 1, sd = 1),
truncation = list(lower = 0, upper = Inf))
# the prior distribution can be visualized using the plot function
# (see ?plot.prior for all options)
plot(p1)</pre>
```

prior\_none

Creates a prior distribution

#### **Description**

prior creates a prior distribution. The prior can be visualized by the plot function.

#### Usage

```
prior_none(prior_weights = 1)
```

#### **Arguments**

prior\_weights

prior odds associated with a given distribution. The value is passed into the model fitting function, which creates models corresponding to all combinations of prior distributions for each of the model parameters and sets the model priors odds to the product of its prior distributions.

rho2logsdr

#### Value

prior and prior\_none return an object of class 'prior'. A named list containing the distribution name, parameters, and prior weights.

#### See Also

plot.prior(), Normal, Lognormal, Cauchy, Beta, Exponential, LocationScaleT, InvGamma.

#### **Examples**

```
# create a standard normal prior distribution
p1 <- prior(distribution = "normal", parameters = list(mean = 1, sd = 1))
# create a half-normal standard normal prior distribution
p2 <- prior(distribution = "normal", parameters = list(mean = 1, sd = 1),
truncation = list(lower = 0, upper = Inf))
# the prior distribution can be visualized using the plot function
# (see ?plot.prior for all options)
plot(p1)</pre>
```

rho2logsdr

rho to log standard deviation ratio transformations

# Description

A list containing the transformation function, inverse transformation function, and the jacobian function.

# Usage

rho2logsdr

#### **Format**

An object of class list of length 3.

#### Value

a list with the corresponding functions

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RoBTT

Estimate a Robust Bayesian T-Test

#### **Description**

RoBTT is used to estimate a robust Bayesian t-test or truncated Bayesian t-test (if truncation is used). The input either requires the vector of observations for each group, x1, x2, or the summary statistics (only if the normal likelihood models are used).

# Usage

```
RoBTT(
  x1 = NULL,
  x2 = NULL
 mean1 = NULL,
 mean2 = NULL,
  sd1 = NULL,
  sd2 = NULL,
 N1 = NULL,
 N2 = NULL
  truncation = NULL,
 prior_delta = prior(distribution = "cauchy", parameters = list(location = 0, scale =
    sqrt(2)/2)),
 prior_rho = prior(distribution = "beta", parameters = list(alpha = 1, beta = 1)),
 prior_nu = if (is.null(truncation)) prior(distribution = "exp", parameters = list(rate
    = 1)),
 prior_delta_null = prior(distribution = "spike", parameters = list(location = 0)),
 prior_rho_null = prior(distribution = "spike", parameters = list(location = 0.5)),
  prior_nu_null = prior_none(),
  prior_mu = NULL,
  prior_sigma2 = NULL,
  chains = 4,
  iter = 10000,
  warmup = 5000,
  thin = 1,
  parallel = FALSE,
  control = set_control(),
  convergence_checks = set_convergence_checks(),
  save = "all",
  seed = NULL,
  silent = TRUE,
)
```

#### **Arguments**

x1

vector of observations of the first group

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x2	vector of observations of the second group
mean1	mean of the first group
mean2	mean of the first group
sd1	standard deviation of the first group
sd2	standard deviation of the first group
N1	sample size of the first group
N2	sample size of the first group
truncation	an optional list specifying truncation applied to the data. Defaults to NULL, i.e., no truncation was applied and the full likelihood is applied. Alternative the truncation can be specified via a named list with:
	"x" where x is a vector of two values specifying the lower and upper truncation points common across the groups
	"x1" <b>and</b> "x2" where x1 is a vector of two values specifying the lower and upper truncation points for the first group and x2 is a vector of two values specifying the lower and upper truncation points for the second group.
	"sigma" where sigma corresponds to the number of standard deviations from the common mean where the truncation points should be set.
	"sigma1" and "sigma2" where sigma1 corresponds to the number of standard deviations from the mean of the first group where the truncation points should be set and sigma2 corresponds to the number of standard deviations from the mean of the second group where the truncation points should be set.
prior_delta	prior distributions for the effect size delta parameter that will be treated as belonging to the alternative hypothesis. Defaults to prior(distribution = "Cauchy", parameters = list(location = 0, scale = sqrt(2)/2)).
prior_rho	prior distributions for the precision allocation rho parameter that will be treated as belonging to the alternative hypothesis. Defaults to prior(distribution = "beta", parameters = list(alpha = 1, beta = 1)).
prior_nu	prior distribution for the degrees of freedom + 2 nu parameter that will be treated as belonging to the alternative hypothesis. Defaults to prior(distribution = "exp", parameters = list(rate = 1)) if no truncation is specified. If truncation is specified, the default is NULL (i.e., use only normal likelihood).
prior_delta_nul	11
	prior distribution for the delta parameter that will be treated as belonging to the null hypothesis. Defaults to point distribution with location at 0 (prior(distribution = "point", parameters = list(location = 0))).
prior_rho_null	prior distribution for the rho parameter that will be treated as belonging to the null hypothesis. Defaults to point distribution with location at $0.5$ (prior(distribution = "point", parameters = list(location = $0.5$ )).
prior_nu_null	prior distribution for the nu parameter that will be treated as belonging to the null hypothesis. Defaults to prior_none ( (i.e., normal likelihood)).
prior_mu	prior distribution for the grand mean parameter. Defaults to NULL which sets Jeffreys prior for the grand mean in case of no truncation or an unit Cauchy prior distributions for the grand mean in case of truncation (which greatly improves sampling efficiency).

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prior\_sigma2 prior distribution for the grand variance parameter. Defaults to NULL which sets Jeffreys prior for the variance in case of no truncation or an exponential prior distribution for the variance in case of truncation (which greatly improves sampling efficiency). a number of chains of the MCMC algorithm. chains a number of sampling iterations of the MCMC algorithm. Defaults to 10000, iter with a minimum of 4000. warmup a number of warmup iterations of the MCMC algorithm. Defaults to 5000. a thinning of the chains of the MCMC algorithm. Defaults to 1. thin parallel whether the individual models should be fitted in parallel. Defaults to FALSE. The implementation is not completely stable and might cause a connection error. control allows to pass control settings with the set\_control() function. See ?set\_control for options and default settings. convergence\_checks automatic convergence checks to assess the fitted models, passed with set\_convergence\_checks() function. See ?set\_convergence\_checks for options and default settings. whether all models posterior distributions should be kept after obtaining a modelsave averaged result. Defaults to "all" which does not remove anything. Set to "min" to significantly reduce the size of final object, however, some model diagnostics and further manipulation with the object will not be possible.

silent whether all pri

terior mixing for reproducibility of results. Defaults to NULL - no seed is set. whether all print messages regarding the fitting process should be suppressed.

a seed to be set before model fitting, marginal likelihood computation, and pos-

Defaults to TRUE. Note that parallel = TRUE also suppresses all messages.

... additional arguments.

#### **Details**

seed

See Maier et al. (2024) for more details regarding the robust Bayesian t-test methodology and the corresponding vignette ("Introduction\_to\_RoBTT", package = "RoBTT")).

See Godmann et al. (2024) for more details regarding the truncated Bayesian t-test methodology and the corresponding vignette (vignette("Truncated\_t\_test", package = "RoBTT")).

Generic summary.RoBTT(), print.RoBTT(), and plot.RoBTT() functions are provided to facilitate manipulation with the ensemble.

#### Value

RoBTT returns an object of class "RoBTT".

#### References

Godmann HR, Bartoš F, Wagenmakers E (2024). "Truncating the Likelihood Allows Outlier Exclusion Without Overestimating the Evidence in the Bayes Factor t-Test." doi:10.31234/osf.io/j9f3s, PsyArxiv Preprint.

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Maier M, Bartoš F, Quintana DS, van den Bergh D, Marsman M, Ly A, Wagenmakers E (2024). "Model-averaged Bayesian t-tests." *Psychonomic Bulletin & Review*. doi:10.3758/s13423024-025905.

#### See Also

```
summary.RoBTT(), prior()
```

#### **Examples**

```
## Not run:
# using the example data from Darwin
data("fertilization", package = "RoBTT")
fit <- RoBTT(</pre>
 x1
           = fertilization$Self,
           = fertilization$Crossed,
 prior_delta = prior("cauchy", list(0, 1/sqrt(2))),
 prior_rho = prior("beta", list(3, 3)),
 seed
              = 1,
              = 1,
 chains
 warmup
             = 1000,
 iter
              = 2000,
 control
             = set_control(adapt_delta = 0.95)
)
# summary can provide many details about the model
summary(fit)
## End(Not run)
```

RoBTT\_control

Convergence checks of the fitting process

### **Description**

Set values for the convergence checks of the fitting process.

#### Usage

```
set_convergence_checks(max_Rhat = 1.05, min_ESS = 500)
set_control(adapt_delta = 0.8, max_treedepth = 15, bridge_max_iter = 1000)
```

#### Arguments

```
max_Rhat maximum value of the R-hat diagnostic. Defaults to 1.05.
min_ESS minimum estimated sample size. Defaults to 500.
adapt_delta tuning parameter of HMC. Defaults to 0.80.
```

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```
max_treedepth tuning parameter of HMC. Defaults to 15.

bridge_max_iter

maximum number of iterations for the bridge_sampler function. Defaults to 10000
```

#### Value

set\_control returns a list of control settings and set\_convergence\_checks returns a list of convergence checks settings.

RoBTT\_options

Options for the 'RoBTT' package

#### **Description**

A placeholder object and functions for the 'RoBTT' package. (adapted from the runjags R package).

#### Usage

```
RoBTT.options(...)
RoBTT.get_option(name)
```

#### **Arguments**

... named option(s) to change - for a list of available options, see details below.

the name of the option to get the current value of - for a list of available options,

see details below.

#### Value

The current value of all available 'RoBTT' options (after applying any changes specified) is returned invisibly as a named list.

summary.RoBTT

Summarize fitted 'RoBTT' object

#### **Description**

summary. RoBTT creates summary tables for a RoBTT object.

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#### Usage

```
## S3 method for class 'RoBTT'
summary(
  object,
  type = "ensemble",
  conditional = FALSE,
  group_estimates = FALSE,
  probs = c(0.025, 0.975),
  logBF = FALSE,
  BF01 = FALSE,
  short_name = FALSE,
  remove_spike_0 = FALSE,
  ...
)
```

#### Arguments

object a fitted 'RoBTT' object

type whether to show the overall 'RoBTT' results ("ensemble"), an overview of the

individual models ("models"), an overview of the individual models MCMC diagnostics ("diagnostics"), or a detailed summary of the individual models

("individual"). Can be abbreviated to first letters.

conditional show the conditional estimates (assuming that the alternative is true). Defaults

to FALSE. Only available for type == "conditional".

group\_estimates

show the model-averaged mean and standard deviation estimates for each group.

probs quantiles of the posterior samples to be displayed. Defaults to c(.025, .975)

logBF show log of Bayes factors. Defaults to FALSE.

BF01 show Bayes factors in support of the null hypotheses. Defaults to FALSE.

short\_name whether priors names should be shortened to the first (couple) of letters. Defaults

to FALSE.

remove\_spike\_0 whether spike prior distributions with location at zero should be omitted from

the summary. Defaults to FALSE.

... additional arguments

#### Value

summary.RoBTT returns a list of tables of class 'BayesTools\_table'.

#### See Also

RoBTT()

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#### **Examples**

```
## Not run:
# using the example data from Darwin
data("fertilization", package = "RoBTT")
fit <- RoBTT(</pre>
 x1
          = fertilization$Self,
 x2
          = fertilization$Crossed,
 prior_delta = prior("cauchy", list(0, 1/sqrt(2))),
 prior_rho = prior("beta", list(3, 3)),
 seed
             = 1,
 chains
             = 1.
 warmup
             = 1000,
 iter
             = 2000,
 control = set_control(adapt_delta = 0.95)
)
# summary can provide many details about the model
summary(fit)
# estimates from the conditional models can be obtained with
summary(fit, conditional = TRUE)
# overview of the models and their prior and posterior probability, marginal likelihood,
# and inclusion Bayes factor can be obtained with
summary(fit, type = "models")
# diagnostics overview, containing the maximum R-hat, minimum ESS, maximum MCMC error, and
# maximum MCMC error / sd across parameters for each individual model can be obtained with
summary(fit, type = "diagnostics")
# summary of individual models and their parameters can be further obtained by
summary(fit, type = "individual")
## End(Not run)
```

update.RoBTT

Updates a fitted RoBTT object

#### Description

update. RoBTT can be used to

- 1. change the prior odds of fitted models by specifying a vector prior\_weights of the same length as the fitted models,
- 2. refitting models that failed to converge with updated settings of control parameters,
- 3. or changing the convergence criteria and recalculating the ensemble results by specifying new control argument and setting refit\_failed == FALSE.

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#### Usage

```
## S3 method for class 'RoBTT'
update(
  object,
  refit_failed = TRUE,
  prior_weights = NULL,
  chains = NULL,
  iter = NULL,
 warmup = NULL,
  thin = NULL,
  parallel = NULL,
  control = NULL,
  convergence_checks = NULL,
  save = "all",
  seed = NULL,
  silent = TRUE,
)
```

#### **Arguments**

object a fitted RoBTT object

refit\_failed whether failed models should be refitted. Relevant only prior\_weights are not

supplied. Defaults to TRUE.

prior\_weights either a single value specifying prior model weight of a newly specified model

using priors argument, or a vector of the same length as already fitted models to

update their prior weights.

chains a number of chains of the MCMC algorithm.

iter a number of sampling iterations of the MCMC algorithm. Defaults to 10000,

with a minimum of 4000.

warmup a number of warmup iterations of the MCMC algorithm. Defaults to 5000.

thin a thinning of the chains of the MCMC algorithm. Defaults to 1.

parallel whether the individual models should be fitted in parallel. Defaults to FALSE.

The implementation is not completely stable and might cause a connection error.

control allows to pass control settings with the set\_control() function. See ?set\_control

for options and default settings.

convergence\_checks

automatic convergence checks to assess the fitted models, passed with set\_convergence\_checks()

function. See ?set\_convergence\_checks for options and default settings.

save whether all models posterior distributions should be kept after obtaining a model-

averaged result. Defaults to "all" which does not remove anything. Set to "min" to significantly reduce the size of final object, however, some model di-

agnostics and further manipulation with the object will not be possible.

seed a seed to be set before model fitting, marginal likelihood computation, and pos-

terior mixing for reproducibility of results. Defaults to NULL - no seed is set.

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silent whether all print messages regarding the fitting process should be suppressed.

Defaults to TRUE. Note that parallel = TRUE also suppresses all messages.

... additional arguments.

# **Details**

See RoBTT() for more details.

#### Value

RoBTT returns an object of class 'RoBTT'.

#### See Also

RoBTT(), summary.RoBTT(), prior(), check\_setup()

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