Package 'bridgesampling'

July 22, 2019

```
Type Package
Title Bridge Sampling for Marginal Likelihoods and Bayes Factors
Version 0.7-2
Depends R (>= 3.0.0)
Imports mynfast, Matrix, Brobdingnag, stringr, coda, parallel, scales,
      utils, methods
Suggests testthat, Rcpp, RcppEigen, R2jags, rjags, runjags, knitr,
      rmarkdown, R.rsp, BayesFactor, rstan, rstanarm, nimble,
      MCMCpack
Description Provides functions for estimating marginal likelihoods, Bayes
      factors, posterior model probabilities, and normalizing constants in general,
      via different versions of bridge sampling (Meng & Wong, 1996,
      <a href="http://www3.stat.sinica.edu.tw/statistica/j6n4/j6n43/j6n43.htm">http://www3.stat.sinica.edu.tw/statistica/j6n4/j6n43/j6n43.htm</a>).
License GPL (>= 2)
LazyData true
RoxygenNote 6.1.1
VignetteBuilder knitr, R.rsp
URL https://github.com/quentingronau/bridgesampling
NeedsCompilation no
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Repository CRAN
Date/Publication 2019-07-21 22:40:06 UTC
```

bf

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bf

Bayes Factor(s) from Marginal Likelihoods

Description

Generic function that computes Bayes factor(s) from marginal likelihoods. $bayes_factor()$ is simply an (S3 generic) alias for bf().

Usage

```
bf(x1, x2, log = FALSE, ...)
bayes_factor(x1, x2, log = FALSE, ...)
## Default S3 method:
bayes_factor(x1, x2, log = FALSE, ...)
## S3 method for class 'bridge'
bf(x1, x2, log = FALSE, ...)
## S3 method for class 'bridge_list'
bf(x1, x2, log = FALSE, ...)
## Default S3 method:
bf(x1, x2, log = FALSE, ...)
```

Arguments

x1	Object of class "bridge" or "bridge_list" as returned from bridge_sampler. Additionally, the default method assumes that x1 is a single numeric log marginal likelihood (e.g., from logml) and will throw an error otherwise.
x2	Object of class "bridge" or "bridge_list" as returned from bridge_sampler. Additionally, the default method assumes that x2 is a single numeric log marginal likelihood (e.g., from logml) and will throw an error otherwise.
log	Boolean. If TRUE, the function returns the log of the Bayes factor. Default is FALSE.
	currently not used here, but can be used by other methods.

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Details

Computes the Bayes factor (Kass & Raftery, 1995) in favor of the model associated with x1 over the model associated with x2.

Value

For the default method returns a list of class "bf_default" with components:

- bf: (scalar) value of the Bayes factor in favor of the model associated with x1 over the model associated with x2.
- log: Boolean which indicates whether bf corresponds to the log Bayes factor.

For the method for "bridge" objects returns a list of class "bf_bridge" with components:

- bf: (scalar) value of the Bayes factor in favor of the model associated with x1 over the model associated with x2.
- log: Boolean which indicates whether bf corresponds to the log Bayes factor.

For the method for "bridge_list" objects returns a list of class "bf_bridge_list" with components:

- bf: a numeric vector consisting of Bayes factors where each element gives the Bayes factor for one set of logmls in favor of the model associated with x1 over the model associated with x2. The length of this vector is given by the "bridge_list" element with the most repetitions. Elements with fewer repetitions will be recycled (with warning).
- bf_median_based: (scalar) value of the Bayes factor in favor of the model associated with x1 over the model associated with x2 that is based on the median values of the logml estimates.
- log: Boolean which indicates whether bf corresponds to the log Bayes factor.

Note

```
For examples, see bridge_sampler and the accompanying vignettes: vignette("bridgesampling_example_jags") vignette("bridgesampling_example_stan")
```

Author(s)

Quentin F. Gronau

References

Kass, R. E., & Raftery, A. E. (1995). Bayes factors. *Journal of the American Statistical Association*, 90(430), 773-795. http://dx.doi.org/10.1080/01621459.1995.10476572

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bridge-methods

Methods for bridge and bridge_list objects

Description

Methods defined for objects returned from the generic bridge_sampler function.

Usage

```
## S3 method for class 'bridge'
summary(object, na.rm = TRUE, ...)

## S3 method for class 'bridge_list'
summary(object, na.rm = TRUE, ...)

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

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

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

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

Arguments

```
object, x object of class bridge or bridge_list as returned from bridge_sampler.

na.rm logical. Should NA estimates in bridge_list objects be removed? Passed to error_measures.

... further arguments, currently ignored.
```

Value

The summary methods return a data.frame which contains the log marginal likelihood plus the result returned from invoking error_measures.

The print methods simply print and return nothing.

bridge_sampler

Log Marginal Likelihood via Bridge Sampling

Description

Computes log marginal likelihood via bridge sampling.

Usage

```
bridge_sampler(samples, ...)
## S3 method for class 'stanfit'
bridge sampler(samples = NULL,
  stanfit_model = samples, repetitions = 1, method = "normal",
  cores = 1, use_neff = TRUE, maxiter = 1000, silent = FALSE,
  verbose = FALSE, ...)
## S3 method for class 'mcmc.list'
bridge sampler(samples = NULL,
  log_posterior = NULL, ..., data = NULL, lb = NULL, ub = NULL,
  repetitions = 1, param_types = rep("real", ncol(samples[[1]])),
  method = "normal", cores = 1, use_neff = TRUE, packages = NULL,
  varlist = NULL, envir = .GlobalEnv, rcppFile = NULL,
 maxiter = 1000, silent = FALSE, verbose = FALSE)
## S3 method for class 'mcmc'
bridge_sampler(samples = NULL, log_posterior = NULL,
  ..., data = NULL, lb = NULL, ub = NULL, repetitions = 1,
  method = "normal", cores = 1, use_neff = TRUE, packages = NULL,
  varlist = NULL, envir = .GlobalEnv, rcppFile = NULL,
 maxiter = 1000, param_types = rep("real", ncol(samples)),
  silent = FALSE, verbose = FALSE)
## S3 method for class 'matrix'
bridge_sampler(samples = NULL, log_posterior = NULL,
  ..., data = NULL, lb = NULL, ub = NULL, repetitions = 1,
 method = "normal", cores = 1, use_neff = TRUE, packages = NULL,
  varlist = NULL, envir = .GlobalEnv, rcppFile = NULL,
 maxiter = 1000, param_types = rep("real", ncol(samples)),
  silent = FALSE, verbose = FALSE)
## S3 method for class 'stanreg'
bridge_sampler(samples, repetitions = 1,
  method = "normal", cores = 1, use_neff = TRUE, maxiter = 1000,
  silent = FALSE, verbose = FALSE, ...)
## S3 method for class 'rjags'
```

Arguments

. . .

samples

an mcmc.list object, a fitted stanfit object, a stanreg object, an rjags object, a runjags object, or a matrix with posterior samples (colnames need to correspond to parameter names in 1b and ub) with posterior samples.

additional arguments passed to log_posterior. Ignored for the stanfit and stanreg methods.

stanfit_model

for the stanfit method, an additional object of class "stanfit" with the same model as samples, which will be used for evaluating the log_posterior (i.e., it does not need to contain any samples). The default is to use samples. In case samples was compiled in a different R session or on another computer with a different OS or setup, the samples model usually cannot be used for evaluation. In this case, one can compile the model on the current computer with iter = 0 and pass it here (this usually needs to be done before samples is loaded).

repetitions

number of repetitions.

method

either "normal" or "warp3".

cores

number of cores used for evaluating log_posterior and when calling mvn-fast functions. For evaluating log_posterior, on unix-like systems (where .Platform\$OS.type == "unix" evaluates to TRUE; e.g., Linux and Mac OS) forking via mclapply is used. Hence, elements needed for evaluation should be in the .GlobalEnv. For other systems (e.g., Windows) makeCluster is used and further arguments specified below will be used. mvnfast uses OpenMP.

use_neff

Boolean which determines whether the effective sample size is used in the optimal bridge function. Default is TRUE. If FALSE, the number of samples is used instead. If samples is a matrix, it is assumed that the matrix contains the samples of one chain in order. If samples come from more than one chain, we recommend to use an mcmc.list object for optimal performance.

maxiter maximum number of iterations for the iterative updating scheme. Default is 1,000 to avoid infinite loops. Boolean which determines whether to print the number of iterations of the upsilent dating scheme to the console. Default is FALSE. Boolean. Should internal debug information be printed to console? Default is verbose FALSE. log_posterior function or name of function that takes a parameter vector and the data as input and returns the log of the unnormalized posterior density (i.e., a scalar value). If the function name is passed, the function should exist in the .GlobalEnv. For special behavior if cores > 1 see Details. data data object which is used in log_posterior. named vector with lower bounds for parameters. lb named vector with upper bounds for parameters. ub character vector of length ncol(samples) with "real", "simplex" or param_types "circular". For all regular bounded or unbounded continuous parameters, this should just be "real". However, if there are parameters which lie on a simplex or on the circle, this should be noted here. Simplex parameters are parameters which are bounded below by zero and collectively sum to one, such as weights in a mixture model. For these, the stick-breaking transformation is performed as described in the Stan reference manual. The circular variables are given a numerical representation to which the normal distribution is most likely a good fit. Only possible to use with bridge_sampler.matrix. character vector with names of packages needed for evaluating log_posterior packages in parallel (only relevant if cores > 1 and .Platform\$OS.type != "unix"). varlist character vector with names of variables needed for evaluating log_posterior (only needed if cores > 1 and .Platform\$OS.type != "unix" as these objects will be exported to the nodes). These objects need to exist in envir. specifies the environment for varlist (only needed if cores > 1 and .Platform\$OS.type envir != "unix" as these objects will be exported to the nodes). Default is .GlobalEnv. in case cores > 1 and log_posterior is an Rcpp function, rcppFile rcppFile specifies the path to the cpp file (will be compiled on all cores).

Details

Bridge sampling is implemented as described in Meng and Wong (1996, see equation 4.1) using the "optimal" bridge function. When method = "normal", the proposal distribution is a multivariate normal distribution with mean vector equal to the sample mean vector of samples and covariance matrix equal to the sample covariance matrix of samples. For a recent tutorial on bridge sampling, see Gronau et al. (in press).

When method = "warp3", the proposal distribution is a standard multivariate normal distribution and the posterior distribution is "warped" (Meng & Schilling, 2002) so that it has the same mean vector, covariance matrix, and skew as the samples. method = "warp3" takes approximately twice as long as method = "normal".

Note that for the matrix method, the lower and upper bound of a parameter cannot be a function of the bounds of another parameter. Furthermore, constraints that depend on multiple parameters

of the model are not supported. This usually excludes, for example, parameters that constitute a covariance matrix or sets of parameters that need to sum to one.

However, if the retransformations are part of the model itself and the log_posterior accepts parameters on the real line and performs the appropriate Jacobian adjustments, such as done for stanfit and stanreg objects, such constraints are obviously possible (i.e., we currently do not know of any parameter supported within Stan that does not work with the current implementation through a stanfit object).

Parallel Computation: On unix-like systems forking is used via mclapply. Hence elements needed for evaluation of log_posterior should be in the .GlobalEnv.

On other OSes (e.g., Windows), things can get more complicated. For normal parallel computation, the log_posterior function can be passed as both function and function name. If the latter, it needs to exist in the environment specified in the envir argument. For parallel computation when using an Rcpp function, log_posterior can only be passed as the function name (i.e., character). This function needs to result from calling sourceCpp on the file specified in rcppFile.

Due to the way rstan currently works, parallel computations with stanfit and stanneg objects only work with forking (i.e., NOT on Windows).

Value

if repetitions = 1, returns a list of class "bridge" with components:

- logml: estimate of log marginal likelihood.
- niter: number of iterations of the iterative updating scheme.
- method: bridge sampling method that was used to obtain the estimate.
- q11: log posterior evaluations for posterior samples.
- q12: log proposal evaluations for posterior samples.
- q21: log posterior evaluations for samples from proposal.
- q22: log proposal evaluations for samples from proposal.

if repetitions > 1, returns a list of class "bridge_list" with components:

- logml: numeric vector with estimates of log marginal likelihood.
- niter: numeric vector with number of iterations of the iterative updating scheme for each repetition.
- method: bridge sampling method that was used to obtain the estimates.
- repetitions: number of repetitions.

Warning

Note that the results depend strongly on the parameter priors. Therefore, it is strongly advised to think carefully about the priors before calculating marginal likelihoods. For example, the prior choices implemented in **rstanarm** or **brms** might not be optimal from a testing point of view. We recommend to use priors that have been chosen from a testing and not a purely estimation perspective.

Also note that for testing, the number of posterior samples usually needs to be substantially larger than for estimation.

Note

To be able to use a stanreg object for samples, the user crucially needs to have specified the diagnostic_file when fitting the model in **rstanarm**.

Author(s)

Quentin F. Gronau and Henrik Singmann. Parallel computing (i.e., cores > 1) and the stanfit method use code from rstan by Jiaqing Guo, Jonah Gabry, and Ben Goodrich. Ben Goodrich added the stanreg method. Kees Mulder added methods for simplex and circular variables.

References

Gronau, Q. F., Sarafoglou, A., Matzke, D., Ly, A., Boehm, U., Marsman, M., Leslie, D. S., Forster, J. J., Wagenmakers, E.-J., & Steingroever, H. (in press). A tutorial on bridge sampling. *Journal of Mathematical Psychology*. https://arxiv.org/abs/1703.05984 vignette("bridgesampling_tutorial")

Gronau, Q. F., Wagenmakers, E.-J., Heck, D. W., & Matzke, D. (2017). A simple method for comparing complex models: Bayesian model comparison for hierarchical multinomial processing tree models using Warp-III bridge sampling. Manuscript submitted for publication. https://psyarxiv.com/yxhfm

Meng, X.-L., & Wong, W. H. (1996). Simulating ratios of normalizing constants via a simple identity: A theoretical exploration. *Statistica Sinica*, 6, 831-860. http://www3.stat.sinica.edu.tw/statistica/j6n4/j6n43/j6n43.htm

Meng, X.-L., & Schilling, S. (2002). Warp bridge sampling. *Journal of Computational and Graphical Statistics*, 11(3), 552-586. http://dx.doi.org/10.1198/106186002457

Overstall, A. M., & Forster, J. J. (2010). Default Bayesian model determination methods for generalised linear mixed models. *Computational Statistics & Data Analysis*, 54, 3269-3288. http://dx.doi.org/10.1016/j.csda.2010.03.008

See Also

bf allows the user to calculate Bayes factors and post_prob allows the user to calculate posterior model probabilities from bridge sampling estimates. bridge-methods lists some additional methods that automatically invoke the error_measures function.

Examples

```
-.5*t(samples.row) %*% samples.row
lb <- rep(-Inf, 2)</pre>
ub \leftarrow rep(Inf, 2)
names(lb) <- names(ub) <- colnames(samples)</pre>
bridge_result <- bridge_sampler(samples = samples, log_posterior = log_density,
                                 data = NULL, lb = lb, ub = ub, silent = TRUE)
# compare to analytical value
analytical <- log(2*pi)
print(cbind(bridge_result$logml, analytical))
## Not run:
## Example 2: Hierarchical Normal Model
# for a full description of the example, see
vignette("bridgesampling_example_jags")
library(R2jags)
### generate data ###
set.seed(12345)
mu <- 0
tau2 <- 0.5
sigma2 <- 1
n < -20
theta <- rnorm(n, mu, sqrt(tau2))</pre>
y <- rnorm(n, theta, sqrt(sigma2))
### set prior parameters
alpha <- 1
beta <- 1
mu0 <- 0
tau20 <- 1
### functions to get posterior samples ###
### H0: mu = 0
getSamplesModelH0 <- function(data, niter = 52000, nburnin = 2000, nchains = 3) {</pre>
  model <- "
   model {
     for (i in 1:n) {
        theta[i] ~ dnorm(0, invTau2)
```

```
y[i] ~ dnorm(theta[i], 1/sigma2)
      invTau2 ~ dgamma(alpha, beta)
      tau2 <- 1/invTau2
  s <- jags(data, parameters.to.save = c("theta", "invTau2"),
            model.file = textConnection(model),
            n.chains = nchains, n.iter = niter,
            n.burnin = nburnin, n.thin = 1)
  return(s)
### H1: mu != 0
getSamplesModelH1 <- function(data, niter = 52000, nburnin = 2000,</pre>
                               nchains = 3) {
  model <- "
    model {
     for (i in 1:n) {
       theta[i] ~ dnorm(mu, invTau2)
        y[i] ~ dnorm(theta[i], 1/sigma2)
      mu \sim dnorm(mu0, 1/tau20)
      invTau2 ~ dgamma(alpha, beta)
      tau2 <- 1/invTau2
  s <- jags(data, parameters.to.save = c("theta", "mu", "invTau2"),</pre>
            model.file = textConnection(model),
            n.chains = nchains, n.iter = niter,
            n.burnin = nburnin, n.thin = 1)
  return(s)
### get posterior samples ###
# create data lists for Jags
data_H0 \leftarrow list(y = y, n = length(y), alpha = alpha, beta = beta, sigma2 = sigma2)
data_H1 \leftarrow list(y = y, n = length(y), mu0 = mu0, tau20 = tau20, alpha = alpha,
                beta = beta, sigma2 = sigma2)
# fit models
samples_H0 <- getSamplesModelH0(data_H0)</pre>
samples_H1 <- getSamplesModelH1(data_H1)</pre>
### functions for evaluating the unnormalized posteriors on log scale ###
```

```
log_posterior_H0 <- function(samples.row, data) {</pre>
  mu <- 0
  invTau2 <- samples.row[[ "invTau2" ]]</pre>
  theta <- samples.row[ paste0("theta[", seq_along(data$y), "]") ]</pre>
  sum(dnorm(data$y, theta, data$sigma2, log = TRUE)) +
    sum(dnorm(theta, mu, 1/sqrt(invTau2), log = TRUE)) +
    dgamma(invTau2, data$alpha, data$beta, log = TRUE)
}
log_posterior_H1 <- function(samples.row, data) {</pre>
  mu <- samples.row[[ "mu" ]]</pre>
  invTau2 <- samples.row[[ "invTau2" ]]</pre>
  theta <- samples.row[ paste0("theta[", seq_along(data$y), "]") ]</pre>
  sum(dnorm(data$y, theta, data$sigma2, log = TRUE)) +
    sum(dnorm(theta, mu, 1/sqrt(invTau2), log = TRUE)) +
    dnorm(mu, data\$mu0, sqrt(data\$tau20), log = TRUE) +
    dgamma(invTau2, data$alpha, data$beta, log = TRUE)
}
# specify parameter bounds HO
cn <- colnames(samples_H0$BUGSoutput$sims.matrix)</pre>
cn <- cn[cn != "deviance"]</pre>
lb_H0 <- rep(-Inf, length(cn))</pre>
ub_H0 <- rep(Inf, length(cn))</pre>
names(lb_H0) <- names(ub_H0) <- cn
lb_H0[[ "invTau2" ]] <- 0</pre>
# specify parameter bounds H1
cn <- colnames(samples_H1$BUGSoutput$sims.matrix)</pre>
cn <- cn[cn != "deviance"]</pre>
lb_H1 <- rep(-Inf, length(cn))</pre>
ub_H1 <- rep(Inf, length(cn))
names(lb_H1) <- names(ub_H1) <- cn
lb_H1[[ "invTau2" ]] <- 0</pre>
# compute log marginal likelihood via bridge sampling for HO
H0.bridge <- bridge_sampler(samples = samples_H0, data = data_H0,</pre>
                              log_posterior = log_posterior_H0, lb = lb_H0,
                              ub = ub_H0, silent = TRUE)
print(H0.bridge)
# compute log marginal likelihood via bridge sampling for H1
H1.bridge <- bridge_sampler(samples = samples_H1, data = data_H1,
                              log_posterior = log_posterior_H1, lb = lb_H1,
                              ub = ub_H1, silent = TRUE)
print(H1.bridge)
```

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```
# compute percentage error
print(error_measures(H0.bridge)$percentage)
print(error_measures(H1.bridge)$percentage)
# compute Bayes factor
BF01 <- bf(H0.bridge, H1.bridge)
print (BF01)
# compute posterior model probabilities (assuming equal prior model probabilities)
post1 <- post_prob(H0.bridge, H1.bridge)</pre>
print(post1)
# compute posterior model probabilities (using user-specified prior model probabilities)
post2 <- post_prob(H0.bridge, H1.bridge, prior_prob = c(.6, .4))</pre>
print(post2)
## End(Not run)
## Not run:
## -----
## Example 3: rstanarm
## -----
library(rstanarm)
# N.B.: remember to specify the diagnostic_file
fit_1 <- stan_glm(mpg ~ wt + qsec + am, data = mtcars,
                chains = 2, cores = 2, iter = 5000,
                diagnostic_file = file.path(tempdir(), "df.csv"))
bridge_1 <- bridge_sampler(fit_1)</pre>
fit_2 \leftarrow update(fit_1, formula = . \sim . + cyl)
bridge_2 <- bridge_sampler(fit_2, method = "warp3")</pre>
bf(bridge_1, bridge_2)
## End(Not run)
```

error_measures

Error Measures for Estimated Marginal Likelihood

Description

Computes error measures for estimated marginal likelihood.

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Usage

```
## S3 method for class 'bridge'
error_measures(bridge_object, ...)
## S3 method for class 'bridge_list'
error_measures(bridge_object, na.rm = TRUE, ...)
```

Arguments

```
an object of class "bridge" or "bridge_list" as returned from bridge_sampler.
... additional arguments (currently ignored).
na.rm a logical indicating whether missing values in logml estimates should be removed. Ignored for the bridge method.
```

Details

Computes error measures for marginal likelihood bridge sampling estimates. The approximate errors for a bridge_object of class "bridge" that has been obtained with method = "normal" and repetitions = 1 are based on Fruehwirth-Schnatter (2004). Not applicable in case the object of class "bridge" has been obtained with method = "warp3" and repetitions = 1. To assess the uncertainty of the estimate in this case, it is recommended to run the "warp3" procedure multiple times.

Value

If bridge_object is of class "bridge" and has been obtained with method = "normal" and repetitions = 1, returns a list with components:

- re2: approximate relative mean-squared error for marginal likelihood estimate.
- cv: approximate coefficient of variation for marginal likelihood estimate (assumes that bridge estimate is unbiased).
- percentage: approximate percentage error of marginal likelihood estimate.

If bridge_object is of class "bridge_list", returns a list with components:

- min: minimum of the log marginal likelihood estimates.
- max: maximum of the log marginal likelihood estimates.
- IQR: interquartile range of the log marginal likelihood estimates.

Note

```
For examples, see bridge_sampler and the accompanying vignettes: vignette("bridgesampling_example_jags") vignette("bridgesampling_example_stan")
```

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Author(s)

Quentin F. Gronau

References

Fruehwirth-Schnatter, S. (2004). Estimating marginal likelihoods for mixture and Markov switching models using bridge sampling techniques. *The Econometrics Journal*, 7, 143-167. http://dx.doi.org/10.1111/j.1368-423X.2004.00125.x

See Also

The summary methods for bridge and bridge_list objects automatically invoke this function, see bridge-methods.

ier

Standardized International Exchange Rate Changes from 1975 to 1986

Description

This data set contains the changes in monthly international exchange rates for pounds sterling from January 1975 to December 1986 obtained from West and Harrison (1997, pp. 612-615). Currencies tracked are US Dollar (column us_dollar), Canadian Dollar (column canadian_dollar), Japanese Yen (column yen), French Franc (column franc), Italian Lira (column lira), and the (West) German Mark (column mark). Each series has been standardized with respect to its sample mean and standard deviation.

Usage

ier

Format

A matrix with 143 rows and 6 columns.

Source

West, M., Harrison, J. (1997). *Bayesian forecasting and dynamic models* (2nd ed.). Springer-Verlag, New York.

Lopes, H. F., West, M. (2004). Bayesian model assessment in factor analysis. *Statistica Sinica*, 14, 41-67. https://www.jstor.org/stable/24307179

ier

Examples

```
## Not run:
*****
# BAYESIAN FACTOR ANALYSIS (AS PROPOSED BY LOPES & WEST, 2004)
library(bridgesampling)
library(rstan)
cores <- 4
options (mc.cores = cores)
data("ier")
#-----
# plot data
#-----
currency <- colnames(ier)</pre>
label <- c("US Dollar", "Canadian Dollar", "Yen", "Franc", "Lira", "Mark")</pre>
op <- par(mfrow = c(3, 2), mar = c(6, 6, 3, 3))
for (i in seq_along(currency)) {
 plot(ier[,currency[i]], type = "l", col = "darkblue", axes = FALSE,
     ylim = c(-4, 4), ylab = "", xlab = "", lwd = 2)
 axis(1, at = 0:12*12, labels = 1975:1987, cex.axis = 1.7)
 axis(2, at = pretty(c(-4, 4)), las = 1, cex.axis = 1.7)
 mtext("Year", 1, cex = 1.5, line = 3.2)
 mtext("Exchange Rate Changes", 2, cex = 1.4, line = 3.2)
 mtext(label[i], 3, cex = 1.6, line = .1)
par(op)
# stan model
#-----
model_code <-
"data {
 int<lower=1> T; // number of observations
 int<lower=1> m; // number of variables
 int<lower=1> k; // number of factors
 matrix[T,m] Y; // data matrix
transformed data {
 int<lower = 1> r;
 vector[m] zeros;
 r = m * k - k * (k - 1) / 2; // number of non-zero factor loadings
 zeros = rep_vector(0.0, m);
```

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```
parameters {
 real beta_lower[r - k]; // lower-diagonal elements of beta
 real<lower = 0> beta_diag [k]; // diagonal elements of beta
 vector<lower = 0>[m] sigma2; // residual variances
transformed parameters {
 matrix[m,k] beta;
  cov_matrix[m] Omega;
 // construct lower-triangular factor loadings matrix
   int index_lower = 1;
   for (j in 1:k) {
     for (i in 1:m) {
       if (i == j) {
        beta[j,j] = beta_diag[j];
       } else if (i >= j) {
        beta[i,j] = beta_lower[index_lower];
         index_lower = index_lower + 1;
       } else {
         beta[i,j] = 0.0;
     }
   }
  Omega = beta * beta' + diag_matrix(sigma2);
model {
  // priors
  target += normal_lpdf(beta_diag \mid 0, 1) - k * normal_lccdf(0 \mid 0, 1);
  target += normal_lpdf(beta_lower | 0, 1);
  target += inv_gamma_lpdf(sigma2 | 2.2 / 2.0, 0.1 / 2.0);
  // likelihood
  for(t in 1:T) {
   target += multi_normal_lpdf(Y[t] | zeros, Omega);
} "
# compile model
model <- stan_model(model_code = model_code)</pre>
#-----
# fit models and compute log marginal likelihoods
#-----
# function for generating starting values
init_fun <- function(nchains, k, m) {</pre>
 r < -m * k - k * (k - 1) / 2
  out <- vector("list", nchains)</pre>
  for (i in seq_len(nchains)) {
   beta_lower <- array(runif(r - k, 0.05, 1), dim = r - k)
```

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```
beta_diag <- array(runif(k, .05, 1), dim = k)
    sigma2 \leftarrow array(runif(m, .05, 1.5), dim = m)
    out[[i]] <- list(beta_lower = beta_lower,</pre>
                     beta_diag = beta_diag,
                      sigma2 = sigma2)
  return(out)
set.seed(1)
stanfit <- bridge <- vector("list", 3)
for (k in 1:3) {
  stanfit[[k]] <- sampling(model,</pre>
                            data = list(Y = ier, T = nrow(ier),
                                        m = ncol(ier), k = k),
                            iter = 11000, warmup = 1000, chains = 4,
                            init = init_fun(nchains = 4, k = k, m = ncol(ier)),
                            cores = cores, seed = 1)
  bridge[[k]] <- bridge_sampler(stanfit[[k]], method = "warp3",</pre>
                                 repetitions = 10, cores = cores)
}
# example output
summary(bridge[[2]])
# compute posterior model probabilities
pp <- post_prob(bridge[[1]], bridge[[2]], bridge[[3]],</pre>
          model_names = c("k = 1", "k = 2", "k = 3"))
рp
op <- par(mar = c(6, 6, 3, 3))
boxplot(pp, axes = FALSE,
     ylim = c(0, 1), ylab = "",
     xlab = "")
axis(1, at = 1:3, labels = colnames(pp), cex.axis = 1.7)
axis(2, cex.axis = 1.1)
mtext("Posterior Model Probability", 2, cex = 1.5, line = 3.2)
mtext("Number of Factors", 1, cex = 1.4, line = 3.2)
par(op)
## End(Not run)
```

post_prob

Description

Generic function that returns log marginal likelihood from bridge objects. For objects of class "bridge_list", which contains multiple log marginal likelihoods, fun is performed on the vector and its result returned.

Usage

```
logml(x, ...)
## S3 method for class 'bridge'
logml(x, ...)
## S3 method for class 'bridge_list'
logml(x, fun = median, ...)
```

Arguments

```
x Object of class "bridge" or "bridge_list" as returned from bridge_sampler.
... Further arguments passed to fun.
fun Function which returns a scalar value and is applied to the logml vector of "bridge_list" objects. Default is median.
```

Value

scalar numeric

post_prob

Posterior Model Probabilities from Marginal Likelihoods

Description

Generic function that computes posterior model probabilities from marginal likelihoods.

Usage

```
post_prob(x, ..., prior_prob = NULL, model_names = NULL)
## S3 method for class 'bridge'
post_prob(x, ..., prior_prob = NULL,
    model_names = NULL)
## S3 method for class 'bridge_list'
post_prob(x, ..., prior_prob = NULL,
    model_names = NULL)
## Default S3 method:
post_prob(x, ..., prior_prob = NULL,
    model_names = NULL)
```

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Arguments

Х	Object of class "bridge" or "bridge_list" as returned from bridge_sampler. Additionally, the default method assumes that all passed objects are numeric log marginal likelihoods (e.g., from logml) and will throw an error otherwise.
• • •	further objects of class "bridge" or "bridge_list" as returned from bridge_sampler. Or numeric values for the default method.
prior_prob	numeric vector with prior model probabilities. If omitted, a uniform prior is used (i.e., all models are equally likely a priori). The default NULL corresponds to equal prior model weights.
model_names	If NULL (the default) will use model names derived from deparsing the call. Otherwise will use the passed values as model names.

Value

For the default method and the method for "bridge" objects, a named numeric vector with posterior model probabilities (i.e., which sum to one).

For the method for "bridge_list" objects, a matrix consisting of posterior model probabilities where each row sums to one and gives the model probabilities for one set of logmls. The (named) columns correspond to the models and the number of rows is given by the "bridge_list" element with the most repetitions. Elements with fewer repetitions will be recycled (with warning).

Note

```
For realistic examples, see bridge_sampler and the accompanying vignettes: vignette("bridgesampling_example_jags") vignette("bridgesampling_example_stan")
```

Author(s)

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Examples

```
post_prob(c(H0$logml, H1$logml, H2$logml))
post_prob(H0$logml, c(H1$logml, H2$logml))
post_prob(H0$logml, c(H1$logml, H2$logml), model_names = c("H0", "H1", "H2"))
### with bridge list elements:
HOL < -structure(list(logml = c(-20.8088381186739, -20.8072772698116,
-20.808454454621, -20.8083419072281, -20.8087870541247, -20.8084887398113,
-20.8086023582344, -20.8079083169745, -20.8083048489095, -20.8090050811436
), niter = c(4, 4, 4, 4, 4, 4, 4, 4, 4, 4), method = "normal",
    repetitions = 10), .Names = c("logml", "niter", "method",
"repetitions"), class = "bridge_list")
H1L \leftarrow structure(list(logml = c(-17.961665507006, -17.9611290723151,
-17.9607509604499, -17.9608629535992, -17.9602093576442, -17.9600223300432,
-17.9610157118017, -17.9615557696561, -17.9608437034849, -17.9606743200309
), niter = c(4, 4, 4, 4, 4, 4, 4, 4, 3, 4), method = "normal",
    repetitions = 10), .Names = c("logml", "niter", "method",
"repetitions"), class = "bridge_list")
post_prob(H1L, H0L)
post_prob(H1L, H0L, H0) # last element recycled with warning.
```

turtles

Turtles Data from Janzen, Tucker, and Paukstis (2000)

Description

This data set contains information about 244 newborn turtles from 31 different clutches. For each turtle, the data set includes information about survival status (column y; 0 = died, 1 = survived), birth weight in grams (column x), and clutch (family) membership (column clutch; an integer between one and 31). The clutches have been ordered according to mean birth weight.

Usage

turtles

Format

A data.frame with 244 rows and 3 variables.

Source

Janzen, F. J., Tucker, J. K., & Paukstis, G. L. (2000). Experimental analysis of an early life-history stage: Selection on size of hatchling turtles. *Ecology*, 81(8), 2290-2304. http://doi.org/10.2307/177115

Overstall, A. M., & Forster, J. J. (2010). Default Bayesian model determination methods for generalised linear mixed models. *Computational Statistics & Data Analysis*, 54, 3269-3288. http://dx.doi.org/10.1016/j.csda.2010.03.008

Sinharay, S., & Stern, H. S. (2005). An empirical comparison of methods for computing Bayes factors in generalized linear mixed models. *Journal of Computational and Graphical Statistics*, 14(2), 415-435. http://dx.doi.org/10.1198/106186005X47471

Examples

```
## Not run:
# BAYESIAN GENERALIZED LINEAR MIXED MODEL (PROBIT REGRESSION)
library (bridgesampling)
library(rstan)
data("turtles")
#-----
# plot data
# reproduce Figure 1 from Sinharay & Stern (2005)
xticks <- pretty(turtles$clutch)</pre>
yticks <- pretty(turtles$x)</pre>
plot(1, type = "n", axes = FALSE, ylab = "", xlab = "", xlim = range(xticks),
   ylim = range(yticks))
points(turtles$clutch, turtles$x, pch = ifelse(turtles$y, 21, 4), cex = 1.3,
     col = ifelse(turtles$y, "black", "darkred"), bg = "grey", lwd = 1.3)
axis(1, cex.axis = 1.4)
mtext("Clutch Identifier", side = 1, line = 2.9, cex = 1.8)
axis(2, las = 1, cex.axis = 1.4)
mtext("Birth Weight (Grams)", side = 2, line = 2.6, cex = 1.8)
# Analysis: Natural Selection Study (compute same BF as Sinharay & Stern, 2005)
### HO (model without random intercepts) ###
H0_code <-
"data {
 int<lower = 1> N;
 int<lower = 0, upper = 1> y[N];
 real<lower = 0 > x[N];
parameters {
 real alpha0_raw;
 real alpha1_raw;
transformed parameters {
 real alpha0 = sqrt(10.0) * alpha0_raw;
 real alpha1 = sqrt(10.0) * alpha1_raw;
```

```
model {
  // priors
  target += normal_lpdf(alpha0_raw | 0, 1);
  target += normal_lpdf(alpha1_raw | 0, 1);
  // likelihood
  for (i in 1:N) {
    target += bernoulli_lpmf(y[i] | Phi(alpha0 + alpha1 * x[i]));
} "
### H1 (model with random intercepts) ###
H1_code <-
"data {
  int<lower = 1> N;
  int<lower = 0, upper = 1 > y[N];
  real<lower = 0 > x[N];
 int<lower = 1> C;
  int<lower = 1, upper = C> clutch[N];
parameters {
 real alpha0_raw;
  real alpha1_raw;
  vector[C] b_raw;
  real<lower = 0> sigma2;
transformed parameters {
  vector[C] b;
  real<lower = 0> sigma = sqrt(sigma2);
  real alpha0 = sqrt(10.0) * alpha0_raw;
  real alpha1 = sqrt(10.0) * alpha1_raw;
  b = sigma * b_raw;
}
model {
  // priors
  target += -2 * log(1 + sigma2); // p(sigma2) = 1 / (1 + sigma2) ^ 2
  target += normal_lpdf(alpha0_raw | 0, 1);
  target += normal_lpdf(alpha1_raw | 0, 1);
  // random effects
  target += normal_lpdf(b_raw | 0, 1);
  // likelihood
  for (i in 1:N) {
    target += bernoulli_lpmf(y[i] | Phi(alpha0 + alpha1 * x[i] + b[clutch[i]]));
} "
set.seed(1)
### fit models ###
stanfit_H0 <- stan(model_code = H0_code,</pre>
                   data = list(y = turtles$y, x = turtles$x, N = nrow(turtles)),
```

```
iter = 15500, warmup = 500, chains = 4, seed = 1)
stanfit_H1 <- stan(model_code = H1_code,</pre>
                   data = list(y = turtles$y, x = turtles$x, N = nrow(turtles),
                                C = max(turtles$clutch), clutch = turtles$clutch),
                   iter = 15500, warmup = 500, chains = 4, seed = 1)
set.seed(1)
### compute (log) marginal likelihoods ###
bridge_H0 <- bridge_sampler(stanfit_H0)</pre>
bridge_H1 <- bridge_sampler(stanfit_H1)</pre>
### compute approximate percentage errors ###
error_measures(bridge_H0)$percentage
error_measures(bridge_H1)$percentage
### summary ###
summary(bridge_H0)
summary(bridge_H1)
### compute Bayes factor ("true" value: BF01 = 1.273) ###
bf(bridge_H0, bridge_H1)
## End(Not run)
```