

# Package ‘DEBBI’

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**Title** Differential Evolution-Based Bayesian Inference

**Version** 0.1.0

**Description** Bayesian inference algorithms based on the population-based ``differential evolution'' (DE) algorithm. Users can obtain posterior mode (MAP) estimates via DEMAP, posterior samples via DEMCMC, and variational approximations via DEVI.

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**URL** <https://github.com/bmgaldo/DEBBI>

**BugReports** <https://github.com/bmgaldo/DEBBI>

**Encoding** UTF-8

**Imports** stats, doParallel, randtoolbox, numDeriv

**Depends** R (>= 3.5.0)

**RoxygenNote** 7.1.2

**NeedsCompilation** no

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`AlgoParamsDEMAP`*AlgoParamsDEMAP*

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## Description

get control parameters for DEMPAP function

## Usage

```
AlgoParamsDEMAP(
  n_params,
  n_chains = NULL,
  n_iter = 1000,
  init_sd = 0.01,
  init_center = 0,
  n_cores_use = 1,
  step_size = NULL,
  jitter_size = 1e-06,
  crossover_rate = 1,
  parallel_type = "none",
  return_trace = FALSE,
  thin = 1
)
```

## Arguments

<code>n_params</code>	number of free parameters estimated
<code>n_chains</code>	number of particle chains, $3 \times n\_params$ is the default value
<code>n_iter</code>	number of iterations to run the sampling algorithm, 1000 is default
<code>init_sd</code>	positive scalar or <code>n_params</code> -dimensional numeric vector, determines the standard deviation of the Gaussian initialization distribution
<code>init_center</code>	scalar or <code>n_params</code> -dimensional numeric vector that determines the mean of the Gaussian initialization distribution
<code>n_cores_use</code>	number of cores used when using parallelization.
<code>step_size</code>	positive scalar, jump size in DE crossover step, default is $2.38/\sqrt{2 \times n\_params}$ .
<code>jitter_size</code>	positive scalar, noise is added during crossover step from $\text{Uniform}(-\text{jitter\_size}, \text{jitter\_size})$ distribution. $1e-6$ is the default value.
<code>crossover_rate</code>	number on the interval $(0,1]$ . Determines the probability a parameter on a chain is updated on a given crossover step, sampled from a Bernoulli distribution.
<code>parallel_type</code>	string specifying parallelization type. 'none', 'FORK', or 'PSOCK' are valid values. 'none' is default value.
<code>return_trace</code>	logical, if true, function returns particle trajectories. This is helpful for diagnosing convergence or debugging model code. Function will return an iteration/thin $\$x\$ n\_chains \$x\$ n\_params$ array and the estimated ELBO of each particle in a iteration/thin $x n\_chains$ array.

**thin** positive integer, only every 'thin'-th iteration will be stored. Default value is 1. Increasing thin will reduce the memory required, while running chains for longer.

**Value**

list of control parameters for the DEMAP function

AlgoParamsDEMCMC

AlgoParamsDEMCMC

**Description**

AlgoParamsDEMCMC

**Usage**

```
AlgoParamsDEMCMC(
  n_params,
  n_chains = NULL,
  param_names = NULL,
  n_iter = 1000,
  init_sd = 0.01,
  init_center = 0,
  n_cores_use = 1,
  step_size = NULL,
  jitter_size = 1e-06,
  parallel_type = "none",
  burnin = 0,
  thin = 1
)
```

**Arguments**

<b>n_params</b>	number of free parameters estimated
<b>n_chains</b>	number of MCMC chains, $3*n_{params}$ is the default value
<b>param_names</b>	optional vector of parameter names
<b>n_iter</b>	number of iterations to run the sampling algorithm, 1000 is default
<b>init_sd</b>	positive scalar or $n_{params}$ -dimensional numeric vector, determines the standard deviation of the Gaussian initialization distribution
<b>init_center</b>	scalar or $n_{params}$ -dimensional numeric vector, determines the mean of the Gaussian initialization distribution
<b>n_cores_use</b>	number of cores used when using parallelization.
<b>step_size</b>	positive scalar, jump size in DE crossover step, default is $2.38/\sqrt{2*n_{params}}$ which is optimal for multivariate Gaussian target distribution (ter Braak, 2006)

jitter_size	positive scalar, noise is added during crossover step from Uniform(-jitter_size,jitter_size) distribution. 1e-6 is the default value.
parallel_type	string specifying parallelization type. 'none','FORK', or 'PSOCK' are valid values. 'none' is default value.
burnin	number of initial iterations to discard. Default value is 0.
thin	positive integer, only every 'thin'-th iteration will be stored. Default value is 1. Increasing thin will reduce the memory required, while running chains for longer.

**Value**

list of control parameters for the DEMCMC function

AlgoParamsDEVI

*AlgoParamsDEVI***Description**

get control parameters for DEVI function

**Usage**

```
AlgoParamsDEVI(
  n_params,
  param_names = NULL,
  n_chains = NULL,
  n_iter = 1000,
  init_sd = 0.01,
  init_center = 0,
  n_cores_use = 1,
  step_size = NULL,
  jitter_size = 1e-06,
  parallel_type = "none",
  use_QMC = TRUE,
  purify = NULL,
  quasi_rand_seq = "halton",
  n_samples_ELBO = 10,
  LRVB_correction = TRUE,
  n_samples_LRVB = 25,
  neg_inf = -750,
  thin = 1,
  burnin = 0,
  return_trace = FALSE,
  crossover_rate = 1
)
```

## Arguments

n_params	number of free parameters estimated
param_names	optional vector of parameter names
n_chains	number of particle chains used for optimization, $3 * n\_params$ is the default value
n_iter	number of iterations to run the sampling algorithm, 1000 is default
init_sd	positive scalar or $n\_params$ -dimensional numeric vector, determines the standard deviation of the Gaussian initialization distribution
init_center	scalar or $n\_params$ -dimensional numeric vector, determines the mean of the Gaussian initialization distribution
n_cores_use	number of cores used when using parallelization.
step_size	positive scalar, jump size in DE crossover step, default is $2.38 / \sqrt{2 * n\_params}$ .
jitter_size	positive scalar, noise is added during crossover step from Uniform(-jitter_size,jitter_size) distribution. 1e-6 is the default value.
parallel_type	string specifying parallelization type. 'none', 'FORK', or 'PSOCK' are valid values. 'none' is default value.
use_QMC	logical, if true, a quasi-Monte Carlo estimator is used to estimate ELBO during optimization. default is TRUE.
purify	an integer, every 'purify'-th iteration, the Monte Carlo estimator of the ELBO is recalculated. This can help deal with noisy and outlier estimates of the ELBO. Default value is 25. If use_QMC is TRUE, purification is disabled as it is redundant.
quasi_rand_seq	type of low discrepancy sequence used for quasi Monte Carlo integration, either 'sobol' or 'halton'. LRVB correction always use QMC. Default is 'sobol'.
n_samples_ELBO	number of samples used for the Monte Carlo estimator of the ELBO (the objective function). default is 10.
LRVB_correction	logical, if true, LRVB covariance correction (Giordano, Brodderick, & Jordan 2018; Galdo, Bahg, & Turner 2020) is attempted.
n_samples_LRVB	number of samples used for LRVB correction. default is 25.
neg_inf	if density for a given value of theta is numerically 0 for q, this value is assigned for log density. This helps with numeric stability of algorithm. Default value is -750.
thin	positive integer, only every 'thin'-th iteration will be stored. Default value is 1. Increasing thin will reduce the memory required, while running algorithm for longer.
burnin	number of initial iterations to discard. Default value is 0.
return_trace	logical, if true, function returns particle trajectories. This is helpful for diagnosing convergence or debugging model code. Function will return an iteration/thin \$x\$ n_chains \$x\$ n_params array and the estimated ELBO of each particle in a iteration/thin x n_chains array.
crossover_rate	number on the interval (0,1]. Determines the probability a parameter on a chain is updated on a given crossover step, sampled from a Bernoulli distribution.

**Value**

list of control parameters for the DEVI function

DEMAP

*DEMAP***Description**

DE optimization for maximum a posteriori (MAP) estimation; his function tries to find the posterior mode.

**Usage**

```
DEMAP(LogPostLike, control_params = AlgoParamsDEMAP(), ...)
```

**Arguments**

- |                |  |
|----------------|--|
| LogPostLike    | function whose first argument is an n_params-dimensional model parameter vector and returns (scalar) sum of log prior density and log likelihood for the parameter vector. |
| control_params | control parameters for DE algorithm. see <a href="#">AlgoParamsDEMAP</a> function documentation for more details.  |
| ...            | additional arguments to pass LogPostLike   |

**Value**

list contain posterior samples from DEMCMC in a n\_iters\_per\_chain by n\_chains by n\_params array and the log likelihood of each sample in a n\_iters\_per\_chain by n\_chains array.

**Examples**

```
# simulate from model
dataExample <- matrix(stats::rnorm(100, c(-1, 1), c(1, 1)), nrow = 50, ncol = 2, byrow = TRUE)

# list parameter names
param_names_example <- c("mu_1", "mu_2")

# log posterior likelihood function = log likelihood + log prior | returns a scalar
LogPostLikeExample <- function(x, data, param_names) {
  out <- 0

  names(x) <- param_names

  # log prior
  out <- out + sum(dnorm(x["mu_1"], 0, sd = 1, log = TRUE))
  out <- out + sum(dnorm(x["mu_2"], 0, sd = 1, log = TRUE))

  # log likelihoods
```

```

out <- out + sum(dnorm(data[, 1], x["mu_1"], sd = 1, log = TRUE))
out <- out + sum(dnorm(data[, 2], x["mu_2"], sd = 1, log = TRUE))

return(out)
}

# Get map estimates
DEMAP(
  LogPostLike = LogPostLikeExample,
  control_params = AlgoParamsDEMAP(
    n_params = length(param_names_example),
    n_iter = 1000,
    n_chains = 12
  ),
  data = dataExample,
  param_names = param_names_example
)

```

DEMCMC

*DEMCMC*

## Description

Sample from posterior using Differential Evolution Markov Chain Monte Carlo

## Usage

```
DEMCMC(LogPostLike, control_params = AlgoParamsDEMCMC(), ...)
```

## Arguments

- `LogPostLike` function whose first argument is an `n_params`-dimensional model parameter vector and returns (scalar) sum of log prior density and log likelihood for the parameter vector.
- `control_params` control parameters for DEMCMC algorithm. see [AlgoParamsDEMCMC](#) function documentation for more details. You must specify '`n_params`' here.
- `...` additional arguments to pass `LogPostLike`

## Value

list contain posterior samples from DEMCMC in a '`n_samples_per_chain`' by '`n_chains`' by `n_params` array and the log posterior likelihood of each sample in a '`n_samples_per_chain`' by '`n_chains`' array.

## Examples

```
# simulate from model
dataExample <- matrix(stats::rnorm(100, c(-1, 1), c(1, 1)), nrow = 50, ncol = 2, byrow = TRUE)
#
# list parameter names
param_names_example <- c("mu_1", "mu_2")

# log posterior likelihood function = log likelihood + log prior | returns a scalar
LogPostLikeExample <- function(x, data, param_names) {
  out <- 0

  names(x) <- param_names

  # log prior
  out <- out + sum(dnorm(x["mu_1"], 0, sd = 1, log = TRUE))
  out <- out + sum(dnorm(x["mu_2"], 0, sd = 1, log = TRUE))

  # log likelihoods
  out <- out + sum(dnorm(data[, 1], x["mu_1"], sd = 1, log = TRUE))
  out <- out + sum(dnorm(data[, 2], x["mu_2"], sd = 1, log = TRUE))

  return(out)
}

# Sample from posterior
DEMCMC(
  LogPostLike = LogPostLikeExample,
  control_params = AlgoParamsDEMCMC(
    n_params = length(param_names_example),
    n_iter = 1000,
    n_chains = 12
  ),
  data = dataExample,
  param_names = param_names_example
)
```

## Description

DE optimization for mean-field variational inference. Minimizes the KL divergence (maximizes the ELBO) between  $q(\theta|all\lambda)$  and the target posterior  $p(\theta|data)$ . For a tutorial on variational inference check out Galdo, Bahg, & Turner 2020.

## Usage

```
DEVI(LogPostLike, control_params = AlgoParamsDEVI(), ...)
```

## Arguments

LogPostLike function whose first argument is an n\_params-dimensional model parameter vector and returns (scalar) sum of log prior density and log likelihood for the parameter vector.  
 control\_params control parameters for DE algorithm. see [AlgoParamsDEVI](#) function documentation for more details.  
 ... additional arguments to pass LogPostLike

## Value

list contain mean in a n\_iters\_per\_chain by n\_chains by 2\*n\_params\_model array and the ELBO of each sample in a n\_iters\_per\_chain by n\_chains array.

## Examples

```

# simulate from model
dataExample <- matrix(stats::rnorm(100, c(-1, 1), c(1, 1)), nrow = 50, ncol = 2, byrow = TRUE)
## list parameter names
param_names_example <- c("mu_1", "mu_2")

# log posterior likelihood function = log likelihood + log prior | returns a scalar
LogPostLikeExample <- function(x, data, param_names) {
  out <- 0

  names(x) <- param_names

  # log prior
  out <- out + sum(dnorm(x["mu_1"], 0, sd = 1, log = TRUE))
  out <- out + sum(dnorm(x["mu_2"], 0, sd = 1, log = TRUE))

  # log likelihoods
  out <- out + sum(dnorm(data[, 1], x["mu_1"], sd = 1, log = TRUE))
  out <- out + sum(dnorm(data[, 2], x["mu_2"], sd = 1, log = TRUE))

  return(out)
}

# Get variational approximation
DEVI(
  LogPostLike = LogPostLikeExample,
  control_params = AlgoParamsDEVI(
    n_params = length(param_names_example),
    n_iter = 200,
    n_chains = 12
  ),
  data = dataExample,
  param_names = param_names_example
)

```

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