

# Package ‘emBayes’

May 8, 2026

**Type** Package

**Title** Robust Bayesian Variable Selection via Expectation-Maximization

**Version** 0.1.6

**Date** 2024-08-26

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**Description** Variable selection methods have been extensively developed for analyzing highdimensional omics data within both the frequentist and Bayesian frameworks. This package provides implementations of the spike-and-slab quantile (group) LASSO which have been developed along the line of Bayesian hierarchical models but deeply rooted in frequentist regularization methods by utilizing Expectation–Maximization (EM) algorithm. The spike-and-slab quantile LASSO can handle data irregularity in terms of skewness and outliers in response variables, compared to its non-robust alternative, the spike-and-slab LASSO, which has also been implemented in the package. In addition, procedures for fitting the spike-and-slab quantile group LASSO and its non-robust counterpart have been implemented in the form of quantile/least-square varying coefficient mixed effect models for high-dimensional longitudinal data. The core module of this package is developed in 'C++'.

**Depends** R (>= 4.2.0)

**License** GPL-2

**Encoding** UTF-8

**LazyData** true

**Imports** Rcpp, glmnet

**LinkingTo** Rcpp, RcppArmadillo

**RoxygenNote** 7.2.3

**NeedsCompilation** yes

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**Repository** CRAN

**Date/Publication** 2024-09-15 00:00:02 UTC

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emBayes-package	<i>Robust Bayesian Variable Selection via Expectation-Maximization</i>
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### Description

This package provides the implementation of the spike-and-slab quantile LASSO (ssQLASSO) and spike-and-slab quantile group LASSO varying coefficient mixed model (ssQVCM) which combines the strength of Bayesian robust variable selection and the Expectation-Maximization (EM) coordinate descent approach. The alternative methods spike-and-slab LASSO (ssLASSO) and spike-and-slab group LASSO varying coefficient mixed model (ssVCM) are also included in the package.

### Details

Two user friendly, integrated interface **cv.emBayes()** and **emBayes()** allows users to flexibly choose the variable selection method by specifying the following parameter:

- quant: to specify different quantiles when using robust methods.
- func: the model to perform variable selection. Four choices are available: "ssLASSO", "ssQLASSO", "ssVCM" and "ssQVCM".
- error: to specify the difference between expectations of likelihood of two consecutive iterations. It can be used to determine convergence.
- maxiter: to specify the maximum number of iterations.

Function `cv.emBayes()` returns cross-validation errors based on the check loss, least squares loss and Schwarz Information Criterion along with the corresponding optimal tuning parameters. Function `emBayes()` returns the estimated intercept, clinical coefficients, beta coefficients, scale parameter, probability parameter, number of iterations and expectation of likelihood at each iteration.

### References

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### See Also

[cv.emBayes](#) [emBayes](#)

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cv.emBayes

*k-folds cross-validation for 'emBayes'*

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### Description

This function performs cross-validation and returns the optimal values of the tuning parameters.

### Usage

```
cv.emBayes(
  y,
  clin = NULL,
  X,
  W = NULL,
  nt = NULL,
  group = NULL,
  quant,
  t0,
  t1,
```

```

    k,
    func,
    error = 0.01,
    maxiter = 100
  )

```

### Arguments

y	a vector of response variable.
clin	a matrix of clinical factors. It has default value NULL.
X	a matrix of genetic factors.
W	a matrix of random factors.
nt	a vector of number of repeated measurements for each subject. They can be same or different.
group	a vector of group sizes. They can be same or different.
quant	value of quantile.
t0	a user-supplied sequence of the spike scale $s_0$ .
t1	a user-supplied sequence of the slab scale $s_1$ .
k	number of folds for cross-validation.
func	methods to perform variable selection. Four choices are available. For non longitudinal analysis: "ssLASSO" and "ssQLASSO". For longitudinal varying-coefficient analysis: "ssVCM" and "ssQVCM".
error	cutoff value for determining convergence. The algorithm reaches convergence if the difference in the expected log-likelihood of two iterations is less than the value of error. The default value is 0.01.
maxiter	the maximum number of iterations that is used in the estimation algorithm. The default value is 200.

### Details

When performing cross-validation for emBayes, function `cv.emBayes` returns two sets of optimal tuning parameters and their corresponding cross-validation error matrices. The spike scale parameter  $CL.s0$  and the slab scale parameter  $CL.s1$  are obtained based on the quantile check loss. The spike scale parameter  $SL.s0$  and the slab scale parameter  $SL.s1$  are obtained based on the least squares loss. The spike scale parameter  $SIC.s0$  and the slab scale parameter  $SIC.s1$  are obtained based on the Schwarz Information Criterion (SIC). Corresponding error matrices  $CL.CV$ ,  $SL.CV$  and  $SIC.CV$  can also be obtained from the output.

Schwarz Information Criterion has the following form:

$$SIC = \log \sum_{i=1}^n L(y_i - \hat{y}_i) + \frac{\log n}{2n} edf$$

where  $L(\cdot)$  is the check loss and  $edf$  is the number of close to zero residuals ( $\leq 0.001$ ). For non-robust method "ssLASSO", one should use least squares loss for tuning selection. For robust method "ssQLASSO", one can either use quantile check loss or SIC for tuning selection. We suggest using SIC, since it has been extensively utilized for tuning selection in high-dimensional quantile regression, as documented in numerous literature sources.

**Value**

A list with components:

CL.s0	the optimal spike scale under check loss.
CL.s1	the optimal slab scale under check loss.
SL.s0	the optimal slab scale under least squares loss.
SL.s1	the optimal slab scale under least squares loss.
SIC.s0	the optimal slab scale under SIC.
SIC.s1	the optimal slab scale under SIC.
CL.CV	cross-validation error matrix under check loss.
SL.CV	cross-validation error matrix under least squares loss.
SIC.CV	cross-validation error matrix under SIC.

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data	<i>simulated gene expression example data</i>
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**Description**

Simulated gene expression data for demonstrating the usage of emBayes.

**Usage**

```
data(data)
```

**Format**

The data file consists of five components: y, clin, X, quant, coef and clin.coef. The coefficients and clinical coefficients are the true values of parameters used for generating response y. They can be used for performance evaluation.

**Details****The data model for generating response**

Let  $y_i$  be the response of the  $i$ -th subject ( $1 \leq i \leq n$ ). We have  $z_i = (1, z_{i1}, \dots, z_{iq})^\top$  being a  $(q + 1)$ -dimensional vector of which the last  $q$  components indicate clinical factors and  $x_i = (x_{i1}, \dots, x_{ip})^\top$  denoting a  $p$ -dimensional vector of genetic factors. The linear quantile regression model for the  $\tau$ -th quantile ( $0 < \tau < 1$ ) is:

$$y_i = z_i^\top \alpha + x_i^\top \beta + \epsilon_i$$

where  $\alpha = (\alpha_0, \dots, \alpha_q)^\top$  contains the intercept and the regression coefficients for the clinical covariates.  $\beta = (\beta_1, \dots, \beta_p)^\top$  are the regression coefficients and random error  $\epsilon_i = (\epsilon_{i1}, \dots, \epsilon_{in})^\top$  is set to follow a T2 distribution and has value 0 at its  $\tau$ -th quantile.

**See Also**

[emBayes](#)

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emBayes

*fit a model with given tuning parameters*


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## Description

This function performs penalized variable selection based on spike-and-slab quantile LASSO (ssQLASSO), spike-and-slab LASSO (ssLASSO), spike-and-slab quantile group LASSO varying coefficient mixed model (ssQVCM) and spike-and-slab group LASSO varying coefficient mixed model (ssVCM). Typical usage is to first obtain the optimal spike scale and slab scale using cross-validation, then specify them in the 'emBayes' function.

## Usage

```
emBayes(
  y,
  clin = NULL,
  X,
  W = NULL,
  nt = NULL,
  group = NULL,
  quant,
  s0,
  s1,
  func,
  error = 0.01,
  maxiter = 100
)
```

## Arguments

y	a vector of response variable.
clin	a matrix of clinical factors. It has default value NULL.
X	a matrix of genetic factors.
W	a matrix of random factors.
nt	a vector of number of repeated measurements for each subject. They can be same or different.
group	a vector of group sizes. They can be same or different.
quant	value of quantile.
s0	value of the spike scale $s_0$ .
s1	value of the slab scale $s_1$ .
func	methods to perform variable selection. Four choices are available. For non longitudinal analysis: "ssLASSO" and "ssQLASSO". For longitudinal varying-coefficient analysis: "ssVCM" and "ssQVCM".

error	cutoff value for determining convergence. The algorithm reaches convergence if the difference in the expected log-likelihood of two iterations is less than the value of error. The default value is 0.01.
maxiter	the maximum number of iterations that is used in the estimation algorithm. The default value is 200.

## Details

The current version of emBayes supports four types of methods: "ssLASSO", "ssQLASSO", "ssVCM" and "ssQVCM".

- **ssLASSO:** spike-and-slab LASSO fits a Bayesian linear regression through the EM algorithm.
- **ssQLASSO:** spike-and-slab quantile LASSO fits a Bayesian quantile regression (based on asymmetric Laplace distribution) through the EM algorithm.
- **ssVCM:** spike-and-slab group LASSO varying coefficient mixed model fits a Bayesian linear mixed model through the EM algorithm.
- **ssQVCM:** spike-and-slab quantile group LASSO varying coefficient mixed model fits a Bayesian quantile mixed model through the EM algorithm.

Users can choose the desired method by setting `func="ssLASSO"`, `"ssQLASSO"`, `"ssVCM"` or `"ssQVCM"`.

## Value

A list with components:

alpha	a vector containing the estimated intercept and clinical coefficients.
intercept	value of the estimated intercept.
clin.coe	a vector of estimated clinical coefficients.
r	a vector of estimated random effect coefficients.
beta	a vector of estimated beta coefficients.
sigma	value of estimated asymmetric Laplace distribution scale parameter $\sigma$ .
theta	value of estimated probability parameter $\theta$ .
iter	value of number of iterations.
ll	a vector of expectation of likelihood at each iteration.

## Examples

```
data(data)
##load the clinical factors, genetic factors, response and quantile data
clin=data$clin
X=data$X
y=data$y
quant=data$quant

##generate tuning vectors of desired range
t0 <- seq(0.01,0.015,length.out=2)
```

```

t1 <- seq(0.1,0.5,length.out=2)

##perform cross-validation and obtain tuning parameters based on check loss
CV <- cv.emBayes(y,clin,X,W=NULL,nt=NULL,group=NULL,quant,t0,t1,k=5,
func="ssQLASSO",error=0.01,maxiter=200)
s0 <- CV$CL.s0
s1 <- CV$CL.s1

##perform BQLSS under optimal tuning and calculate value of TP and FP for selecting beta
EM <- emBayes(y,clin,X,W=NULL,nt=NULL,group=NULL,quant,s0,s1,func="ssQLASSO",
error=0.01,maxiter=200)
fit <- EM$beta
coef <- data$coef
tp <- sum(fit[coef!=0]!=0)
fp <- sum(fit[coef==0]!=0)
list(tp=tp,fp=fp)

```

---

```
print.cv.emBayes      print an cv.emBayes result
```

---

### Description

Print a summary of an 'cv.emBayes' result

### Usage

```
## S3 method for class 'cv.emBayes'
print(x, digits = max(3, getOption("digits") - 3), ...)
```

### Arguments

x	cv.emBayes result
digits	significant digits in printout.
...	other print arguments

### Value

Print a list of output from a cv.emBayes object.

### See Also

[cv.emBayes](#)

---

`print.emBayes`      *print an emBayes result*

---

### **Description**

Print a summary of an 'emBayes' result

### **Usage**

```
## S3 method for class 'emBayes'  
print(x, digits = max(3, getOption("digits") - 3), ...)
```

### **Arguments**

<code>x</code>	emBayes result
<code>digits</code>	significant digits in printout.
<code>...</code>	other print arguments

### **Value**

Print a list of output from a emBayes object.

### **See Also**

[emBayes](#)

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