

Package ‘pqrBayes’

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Type Package

Title Bayesian Penalized Quantile Regression

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Description Bayesian regularized quantile regression utilizing two major classes of shrinkage priors (the spike-and-slab priors and the horseshoe family of priors) leads to efficient Bayesian shrinkage estimation, variable selection and valid statistical inference. In this package, we have implemented robust Bayesian variable selection with spike-and-slab priors under high-dimensional linear regression models (Fan et al. (2024) <[doi:10.3390/e26090794](https://doi.org/10.3390/e26090794)> and Ren et al. (2023) <[doi:10.1111/biom.13670](https://doi.org/10.1111/biom.13670)>), and regularized quantile varying coefficient models (Zhou et al.(2023) <[doi:10.1016/j.csda.2023.107808](https://doi.org/10.1016/j.csda.2023.107808)>). In particular, valid robust Bayesian inferences under both models in the presence of heavy-tailed errors can be validated on finite samples. Additional models with spike-and-slab priors include robust Bayesian group LASSO and robust binary Bayesian LASSO (Fan and Wu (2025) <[doi:10.1002/sta4.70078](https://doi.org/10.1002/sta4.70078)>). Besides, robust sparse Bayesian regression with the horseshoe family of (horseshoe, horseshoe+ and regularized horseshoe) priors has also been implemented and yielded valid inference results under heavy-tailed model errors(Fan et al.(2025) <[doi:10.48550/arXiv.2507.10975](https://doi.org/10.48550/arXiv.2507.10975)>). The Markov chain Monte Carlo (MCMC) algorithms of the proposed and alternative models are implemented in C++.

Depends R (>= 3.5.0)

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Encoding UTF-8

URL <https://github.com/cenwu/pqrBayes>

BugReports <https://github.com/cenwu/pqrBayes/issues>

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LinkingTo Rcpp, RcppArmadillo

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pqrBayes-package	<i>Bayesian penalized quantile regression for the sparse linear model, binary LASSO, group LASSO and varying coefficient models based on spike-and-slab priors and/or the horseshoe family of priors</i>
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Description

In this package, we implement Bayesian penalized quantile regression for the sparse linear model, binary LASSO, group LASSO and quantile varying coefficient (VC) models. The point-mass spike-and-slab priors and horseshoe family of (horseshoe, horseshoe+ and regularized horseshoe) priors have been incorporated in the Bayesian hierarchical models to facilitate Bayesian shrinkage estimation, variable selection and statistical inference. For the spike-and-slab priors, the four default methods are Bayesian regularized quantile regression with spike-and-slab priors under the sparse linear (i.e. LASSO), binary LASSO, group LASSO and VC model, correspondingly. In addition to the default methods, users can also choose models without robustness and/or spike-and-slab priors. Furthermore, under sparse linear models, we have implemented the horseshoe family of (horseshoe, horseshoe+ and regularized horseshoe) priors and the three non-robust alternatives. Currently, the horseshoe family of priors is only implemented under the sparse linear model.

Details

The user friendly, integrated interface **pqrBayes()** allows users to flexibly choose fitting models by specifying the following parameters:

- robust:** whether to fit a robust sparse quantile regression model (the sparse linear model, binary LASSO, group LASSO or Varying Coefficient models) or their non-robust counterparts.
- prior:** specify which prior to use (the spike-and-slab prior, Laplace prior and the horseshoe family of priors).
- model:** whether to fit a sparse linear model, binary LASSO, group LASSO or a varying coefficient model.

The function **pqrBayes()** returns a **pqrBayes** object that stores the posterior estimates of regression coefficients.

References

- Fan, K., Subedi, S., Yang, G., Lu, X., Ren, J. and Wu, C. (2024). Is Seeing Believing? A Practitioner's Perspective on High-dimensional Statistical Inference in Cancer Genomics Studies. *Entropy*, 26(9).794 doi:[10.3390/e26090794](https://doi.org/10.3390/e26090794)
- Zhou, F., Ren, J., Ma, S. and Wu, C. (2023). The Bayesian regularized quantile varying coefficient model. *Computational Statistics & Data Analysis*, 107808 doi:[10.1016/j.csda.2023.107808](https://doi.org/10.1016/j.csda.2023.107808)
- Ren, J., Zhou, F., Li, X., Ma, S., Jiang, Y. and Wu, C. (2023). Robust Bayesian variable selection for gene-environment interactions. *Biometrics*, 79(2), 684-694 doi:[10.1111/biom.13670](https://doi.org/10.1111/biom.13670)
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- Zhou, F., Ren, J., Lu, X., Ma, S. and Wu, C. (2021). Gene–Environment Interaction: a Variable Selection Perspective. *Epistasis. Methods in Molecular Biology*. 2212:191–223 https://link.springer.com/protocol/10.1007/978-1-0716-0947-7_13
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- Ren, J., Zhou, F., Li, X., Wu, C. and Jiang, Y. (2019) spinBayes: Semi-Parametric Gene-Environment Interaction via Bayesian Variable Selection. R package version 0.1.0. <https://CRAN.R-project.org/package=spinBayes>
- Wu, C., Jiang, Y., Ren, J., Cui, Y. and Ma, S. (2018). Dissecting gene-environment interactions: A penalized robust approach accounting for hierarchical structures. *Statistics in Medicine*, 37:437–456 doi:[10.1002/sim.7518](https://doi.org/10.1002/sim.7518)

Wu, C., Shi, X., Cui, Y. and Ma, S. (2015). A penalized robust semiparametric approach for gene-environment interactions. *Statistics in Medicine*, 34 (30): 4016–4030 [doi:10.1002/sim.6609](#)

Wu, C., Cui, Y., and Ma, S. (2014). Integrative analysis of gene–environment interactions under a multi–response partially linear varying coefficient model. *Statistics in Medicine*, 33(28), 4988–4998 [doi:10.1002/sim.6287](#)

Wu, C., Zhong, P.S. and Cui, Y. (2018). Additive varying–coefficient model for nonlinear gene–environment interactions. *Statistical Applications in Genetics and Molecular Biology*, 17(2) [doi:10.1515/sagmb-20170008](#)

Wu, C., Zhong, P.S. and Cui, Y. (2013). High dimensional variable selection for gene-environment interactions. *Technical Report. Michigan State University.*

See Also

[pqrBayes](#)

coverage	95% empirical coverage probability for a pqrBayes object
----------	--

Description

Calculate 95% empirical coverage probabilities for regression coefficients under the sparse linear model, binary LASSO, group LASSO and VC models, respectively.

Usage

```
coverage(object,coefficient,u.grid=NULL,model="linear")
```

Arguments

object	the pqrBayes object.
coefficient	the vector of true regression coefficients under a sparse linear model, binary LASSO, group LASSO, or the matrix of true varying coefficients evaluated on the grid points under a varying coefficient model.
u.grid	the vector of grid points under a varying coefficient model. When assessing empirical coverage probabilities under a sparse linear model, binary LASSO or group LASSO, u.grid = NULL.
model	the model to be fitted. Users can also choose "linear" for a sparse linear model, "binary" for binary LASSO, "group" for group LASSO, and "VC" for a sparse varying coefficient model.

Value

c

See Also

[pqrBayes](#)

Examples

```
## The quantile regression model
data(data)
data = data$data_linear
g=data$g
y=data$y
e=data$e
coeff = data$coeff
fit1=pqrBayes(g,y,e,d = NULL,quant=0.5,model="linear")
coverage=coverage(fit1,coeff,model="linear")
```

data	<i>Simulated data under high-dimensional linear, binary, group LASSO and quantile varying coefficient models</i>
------	--

Description

Simulated data under high-dimensional linear, binary, group LASSO and quantile varying coefficient models

Format

The data_linear object consists of 4 components: g, y, e and coeff. coeff contains the true values of parameters used for generating the response variable y . The data_binary object consists of 4 components: g, y, e and coeff. coeff contains the true values of parameters used for generating the response variable y . The data_group object consists of 4 components: g, y, e and coeff. coeff contains the true values of parameters used for generating the response variable y . The data_varying object consists of five components: g, y, u, e and coeff. coeff contains the true values of parameters used for generating the response variable y .

Details

Generating Y using a sparse linear (quantile) regression model

The true data generating model under sparse linear regression:

$$Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \beta_3 X_{i3} + \epsilon_i,$$

where $\epsilon_i \sim N(0, 1)$, $\beta_0 = 0$, $\beta_1 = 1$, $\beta_2 = 1.5$ and $\beta_3 = 2$.

Generating Y using a sparse binary (quantile) regression model

The true data generating model under sparse linear regression:

$$\tilde{Y}_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \beta_3 X_{i3} + \epsilon_i,$$

where $\epsilon_i \sim N(0, 1)$, $\beta_0 = 0$, $\beta_1 = 0.22$, $\beta_2 = 0.18$ and $\beta_3 = 0.14$.

$Y_i = 1$ if $\tilde{Y}_i > 0$ and $Y_i = 0$ otherwise.

Generating Y using a high-dimensional group LASSO model

The true data generating model under a group LASSO model:

$$Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \beta_3 X_{i3} + \beta_7 X_{i7} + \beta_8 X_{i8} + \beta_9 X_{i9} + \epsilon_i,$$

where $\epsilon_i \sim N(0, 1)$, $\beta_0 = 0$, $\beta_1 = 0.6$, $\beta_2 = 0.7$, $\beta_3 = 0.8$, $\beta_7 = 0.65$, $\beta_8 = 0.75$ and $\beta_9 = 0.85$.

Generating Y using a (quantile) varying coefficient model

Data generation under sparse (quantile) VC model:

$$Y_i = \gamma_0(v_i) + \gamma_1(v_i)X_{i1} + \gamma_2(v_i)X_{i2} + \gamma_3(v_i)X_{i3} + \epsilon_i,$$

where $\epsilon_i \sim N(0, 1)$, $\gamma_0(v_i) = 1.5 \sin(0.2\pi * v_i)$, $\gamma_1(v_i) = 2 \exp(0.2v_i - 1) - 1.5$, $\gamma_2(v_i) = 2 - 2v_i$ and $\gamma_3(v_i) = -4 + (v_i - 2)^3/6$.

See Also

[pqrBayes](#)

Examples

```
data(data)
data = data$data_linear
g=data$g
dim(g)
y=data$y
coeff=data$coeff
print(coeff)
```

```
data = data$data_binary
g=data$g
dim(g)
y=data$y
coeff=data$coeff
print(coeff)
```

```
data = data$data_group
g=data$g
dim(g)
y=data$y
coeff=data$coeff
print(coeff)
```

```
data = data$data_varying
g=data$g
dim(g)
coeff=data$coeff
print(coeff)
```

estimation.pqrBayes	<i>Estimation and estimation accuracy for a pqrBayes object</i>
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Description

Calculate estimated regression coefficients with estimation accuracy from the sparse linear model, binary LASSO, group LASSO and quantile VC models, respectively.

Usage

```
estimation.pqrBayes(object,coefficient,u.grid=NULL,model="linear")
```

Arguments

object	an object of class 'pqrBayes'.
coefficient	the vector of quantile regression coefficients under a sparse linear model, binary LASSO and group LASSO or the matrix of true varying coefficients evaluated on the grid points under a varying coefficient model.
u.grid	the vector of grid points under a varying coefficient model. When fitting a sparse linear model, binary LASSO or group LASSO, u.grid = NULL.
model	the model to be fitted. Users can choose "linear" for a sparse linear model, "binary" for binary LASSO, "group" for group LASSO or "VC" for a varying coefficient model.

Value

an object of class 'pqrBayes.est' is returned, which is a list with components:

error	mean square error or integrated mean square errors and total integrated mean square error.
coeff.est	estimated values of the regression coefficients or the varying coefficients.

See Also

[pqrBayes](#)

Examples

```
## The quantile regression model
data(data)
data = data$data_linear
g=data$g
y=data$y
e=data$e
coeff = data$coeff
fit1=pqrBayes(g,y,e,d = NULL,quant=0.5,model="linear")
estimation=estimation.pqrBayes(fit1,coeff,model="linear")
```

pqrBayes	<i>fit Bayesian penalized quantile regression for the sparse linear model, binary LASSO, group LASSO, or varying coefficient model based on spike-and-slab priors and/or the horseshoe family of priors</i>
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Description

fit Bayesian penalized quantile regression for the sparse linear model, binary LASSO, group LASSO, or varying coefficient model based on spike-and-slab priors and/or the horseshoe family of priors

Usage

```
pqrBayes(
  g,
  y,
  e,
  d = NULL,
  quant = 0.5,
  iterations = 10000,
  burn.in = NULL,
  robust = TRUE,
  prior = "SS",
  model = "linear",
  hyper = NULL,
  debugging = FALSE
)
```

Arguments

<code>g</code>	the matrix of predictors (subject to selection). Users do not need to specify an intercept which will be automatically included.
<code>y</code>	the response variable.
<code>e</code>	a matrix of clinical covariates not subject to selection.
<code>d</code>	a positive integer denotes the group size. When fitting a sparse linear or varying coefficient model, <code>d = NULL</code> .
<code>quant</code>	the quantile level specified by users. The default value is 0.5.
<code>iterations</code>	the number of MCMC iterations. The default value is 10,000.
<code>burn.in</code>	the number of burn-in iterations. If <code>NULL</code> , the first half of MCMC iterations will be discarded as burn-ins.
<code>robust</code>	logical flag. If <code>TRUE</code> , robust methods are used. Otherwise, non-robust methods are used. The default value is <code>TRUE</code> .
<code>prior</code>	the type of prior used for variable selection. Users can specify "SS" for the spike-and-slab prior, "HS" for the horseshoe prior, "HS+" for the horseshoe plus prior, "RHS" for the regularized horseshoe prior and "Laplace" for the Laplace prior. The default value is "SS".

model	the model to be fitted. Users can specify "linear" for a sparse linear model, "binary" for binary LASSO, "group" for group LASSO and "VC" for a varying coefficient model.
hyper	a named list of hyper-parameters. The default value is NULL.
debugging	logical flag. If TRUE, progress will be output to the console and extra information will be returned. The default value is FALSE.

Details

The sparse linear quantile regression model described in "data" is:

$$Y_i = \sum_{k=1}^q E_{ik} \beta_k + \sum_{j=0}^p X_{ij} \gamma_j + \epsilon_i,$$

where β_k 's are the regression coefficients for clinical covariates and γ_j 's are the regression coefficients of $\mathbf{X}$.

The binary quantile regression model described in "data" is:

$$\tilde{Y}_i = \sum_{k=1}^q E_{ik} \beta_k + \sum_{j=0}^p X_{ij} \gamma_j + \epsilon_i,$$

where β_k 's are the regression coefficients for clinical covariates and γ_j 's are the regression coefficients of $\mathbf{X}$.

$Y_i = 1$ if $\tilde{Y}_i > 0$ and $Y_i = 0$ otherwise.

The group LASSO model described in "data" is:

$$Y_i = \sum_{k=1}^q E_{ik} \beta_k + X_{i0} \gamma_0 + \sum_{j=1}^m \mathbf{X}_{ij}^\top \boldsymbol{\gamma}_j + \epsilon_i,$$

where β_k 's are the regression coefficients for clinical covariates and $\boldsymbol{\gamma}_j = (\gamma_{j1}, \dots, \gamma_{jd})^\top$ is the vector of regression coefficients of the $n \times d$ matrix $\mathbf{X}_j$.

The quantile varying coefficient model described in "data" is:

$$Y_i = \sum_{k=1}^q E_{ik} \beta_k + \sum_{j=0}^p \gamma_j(V_i) X_{ij} + \epsilon_i,$$

where β_k 's are the regression coefficients for the clinical covariates, and γ_j 's are the varying intercept and varying coefficients for predictors (e.g. genetic factors), respectively.

Users can modify the hyper-parameters by providing a named list of hyper-parameters via the argument 'hyper'. The list can have the following named components

a0, b0: shape parameters of the Beta priors $(\pi^{a_0-1}(1-\pi)^{b_0-1})$ on π_0 .

c1, c2: the shape parameter and the rate parameter of the Gamma prior on ν .

Please check the references for more details about the prior distributions.

Value

an object of class "pqrBayes" is returned, which is a list with components:

obj	a list of posterior samples from the MCMC and other parameters
coefficients	a list of posterior estimates of coefficients

References

- Fan, K., Subedi, S., Yang, G., Lu, X., Ren, J. and Wu, C. (2024). Is Seeing Believing? A Practitioner's Perspective on High-dimensional Statistical Inference in Cancer Genomics Studies. *Entropy*, 26(9).794 doi:[10.3390/e26090794](https://doi.org/10.3390/e26090794)
- Zhou, F., Ren, J., Ma, S. and Wu, C. (2023). The Bayesian regularized quantile varying coefficient model. *Computational Statistics & Data Analysis*, 107808 doi:[10.1016/j.csda.2023.107808](https://doi.org/10.1016/j.csda.2023.107808)
- Ren, J., Zhou, F., Li, X., Ma, S., Jiang, Y. and Wu, C. (2023). Robust Bayesian variable selection for gene-environment interactions. *Biometrics*, 79(2), 684-694 doi:[10.1111/biom.13670](https://doi.org/10.1111/biom.13670)
- Fan, K. and Wu, C. (2025). A New Robust Binary Bayesian LASSO. *Stat*, 14 (3), e70078 doi:[10.1002/sta4.70078](https://doi.org/10.1002/sta4.70078)
- Fan, K., Srijana, S., Dissanayake, V. and Wu, C. (2025). Robust Bayesian high-dimensional variable selection and inference with the horseshoe family of priors. *arXiv preprint* doi:[10.48550/arXiv.2507.10975](https://doi.org/10.48550/arXiv.2507.10975)
- Ren, J., Zhou, F., Li, X., Chen, Q., Zhang, H., Ma, S., Jiang, Y. and Wu, C. (2020) Semi-parametric Bayesian variable selection for gene-environment interactions. *Statistics in Medicine*, 39: 617– 638 doi:[10.1002/sim.8434](https://doi.org/10.1002/sim.8434)

Examples

```
## The sparse quantile (linear and binary) regression model
data(data)
data = data$data_linear
g=data$g
y=data$y
e=data$e
data(data)
data_1=data$data_binary
g_1=data_1$g
y_1=data_1$y
e_1=data_1$e

fit1=pqrBayes(g,y,e,d=NULL,quant=0.5,model="linear")
fit1_1=pqrBayes(g_1,y_1,e_1,d=NULL,quant=0.5,model="binary")

## Non-sparse example (LASSO)
fit2 <- pqrBayes(
  g = g, y = y, e = e, d = NULL,
  quant = 0.5,
  prior = "Laplace",
  model = "linear"
)
```

```
## Non-robust example (LASSO)
fit3 <- pqrBayes(
  g = g, y = y, e = e, d = NULL,
  quant = 0.5,
  robust = FALSE,
  model = "linear"
)

## The group LASSO model
data(data)
data = data$data_group
g=data$g
y=data$y
e=data$e

fit1=pqrBayes(g,y,e,d=3,quant=0.5,model="group")

## Non-sparse version
fit2 <- pqrBayes(
  g = g, y = y, e = e, d = 3,
  quant = 0.5,
  prior = "Laplace",
  model = "group"
)

## Non-robust version
fit3 <- pqrBayes(
  g = g, y = y, e = e, d = 3,
  quant = 0.5,
  robust = FALSE,
  model = "group"
)

## The quantile varying coefficient model
data(data)
data = data$data_varying
g=data$g
y=data$y
e=data$e
fit1=pqrBayes(g,y,e,quant=0.5,model="VC")

## Non-sparse example
fit2 <- pqrBayes(
  g = g, y = y, e = e,
  quant = 0.5,
  prior = "Laplace",
  model = "VC"
)
```

```
## Non-robust example
fit3 <- pqrBayes(
  g = g, y = y, e = e,
  quant = 0.5,
  robust = FALSE,
  model = "VC"
)
```

pqrBayes.select	Variable selection for a pqrBayes object
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Description

Variable selection for a pqrBayes object

Usage

```
pqrBayes.select(object, prior="SS", model="linear")
```

Arguments

- object a pqrBayes object.
- prior the prior used in the pqrBayes function. Users can choose "SS" for the spike-and-slab prior, "HS" for the horseshoe prior, "HS+" for the horseshoe plus prior, "RHS" for the regularized horseshoe prior and "Laplace" for the Laplace prior. The default value is "SS".
- model the model to be fitted. Users can also choose "linear" for a sparse linear model, "VC" for a varying coefficient model or "group" for group LASSO.

Details

For class ‘Sparse’, the median probability model (MPM) (Barbieri and Berger, 2004) is used to identify predictors that are significantly associated with the response variable. For class ‘NonSparse’, variable selection is based on 95% credible interval. Please check the references for more details about the variable selection.

Value

an object of class ‘select’ is returned, which includes the indices of the selected predictors (e.g. genetic factors).

References

Ren, J., Zhou, F., Li, X., Ma, S., Jiang, Y. and Wu, C. (2023). Robust Bayesian variable selection for gene-environment interactions. *Biometrics*, 79(2), 684-694 [doi:10.1111/biom.13670](#)

Barbieri, M.M. and Berger, J.O. (2004). Optimal predictive model selection. *Ann. Statist.*, 32(3):870–897

See Also[pqrBayes](#)**Examples**

```
## The sparse quantile regression model
data(data)
data = data$data_linear
g=data$g
y=data$y
e=data$e

fit1=pqrBayes(g,y,e,d = NULL,quant=0.5,model="linear")
select=pqrBayes.select(obj = fit1,prior = "SS",model="linear")

## The quantile varying coefficient model
data(data)
data = data$data_varying
g=data$g
y=data$y
e=data$e
fit1=pqrBayes(g,y,e,d = NULL,quant=0.5,model="VC")
select=pqrBayes.select(obj = fit1,prior = "SS",model="VC")
select

## Non-sparse example with VC model
fit2 <- pqrBayes(
  g = g, y = y, e = e, d = NULL,
  quant = 0.5,
  prior= "Laplace",
  model = "VC"
)

select <- pqrBayes.select(obj = fit2, prior = "SS", model = "VC")
select
```

predict_pqrBayes*Make predictions from a pqrBayes object*

Description

Make predictions from a pqrBayes object

Usage

predict_pqrBayes(object, g.new, u.new, e.new, y.new, quant, model, ...)

Arguments

<code>object</code>	a pqrBayes object.
<code>g.new</code>	a matrix of new predictors (e.g. genetic factors) at which predictions are to be made. When being applied to the sparse linear, binary LASSO or group LASSO, <code>g.new = g</code> .
<code>u.new</code>	a vector of new environmental factor at which predictions are to be made. When being applied to the sparse linear model, binary LASSO or group LASSO, <code>u.new = e</code> .
<code>e.new</code>	a vector or matrix of new clinical covariates at which predictions are to be made.
<code>y.new</code>	a vector of the response of new observations. When being applied to the sparse linear model, binary LASSO or group LASSO, <code>y.new = y</code> .
<code>quant</code>	the quantile level. The default is 0.5.
<code>model</code>	the model to be fitted. The default is "VC" for a quantile varying coefficient model. Users can also specify "linear" for a sparse linear model, "binary" for binary LASSO and "group" for a group LASSO.
<code>...</code>	other predict arguments

Details

`g.new` (`u.new`) must have the same number of columns as `g` (`u`) used for fitting the model. By default, the clinical covariates are NULL unless provided. The predictions are made based on the posterior estimates of coefficients in the pqrBayes object.

Value

an object of class 'pqrBayes.pred' is returned, which is a list with components:

<code>error</code>	prediction error.
<code>y.pred</code>	predicted values of the new observations.

See Also

[pqrBayes](#)

Examples

```
## The quantile regression model
data(data)
data = data$data_linear
g=data$g
y=data$y
e=data$e
fit1=pqrBayes(g,y,e,d = NULL,quant=0.5,model="linear")
prediction=predict_pqrBayes(fit1,g,u.new=e,e.new = NULL, y.new = y,model="linear")
```

print.pqrBayes	<i>print a pqrBayes result</i>
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Description

Print a pqrBayes result

Usage

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

Arguments

x	pqrBayes result.
digits	significant digits in printout.
...	other print arguments.

Value

No return value, called for side effects.

See Also

[pqrBayes](#)

print.pqrBayes.pred	<i>print a pqrBayes.pred object</i>
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Description

Print a summary of a pqrBayes.pred object

Usage

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

Arguments

x	pqrBayes.pred object.
digits	significant digits in printout.
...	other print arguments

Value

No return value, called for side effects.

See Also

[predict_pqrBayes](#)

`print.pqrBayes.select` *print a select.pqrBayes object*

Description

Print a summary of a `select.pqrBayes` object

Usage

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

Arguments

<code>x</code>	<code>pqrBayes.select</code> object.
<code>digits</code>	significant digits in printout.
<code>...</code>	other print arguments

Value

No return value, called for side effects.

See Also

[pqrBayes.select](#)

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