

Package ‘bfp’

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Description Implements the Bayesian paradigm for fractional polynomial models under the assumption of normally distributed error terms, see Sabanes Bove, D. and Held, L. (2011) <doi:10.1007/s11222-010-9170-7>.

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as.data.frame.BayesMfp

Convert a BayesMfp object to a data frame

Description

Convert the [BayesMfp](#) object to a data frame with the saved models.

Usage

```
## S3 method for class 'BayesMfp'
as.data.frame(x, row.names = NULL, ..., freq = TRUE)
```

Arguments

x	valid BayesMfp object
row.names	optional rownames (default is to keep the names of the BayesMfp list)
freq	should empirical frequencies of the models in the sampling path be given? (default)
...	unused

Author(s)

Daniel Saban\`es Bov\`e

See Also[summary.BayesMfp](#)**Examples**

```
## generate a BayesMfp object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                method="exhaustive")

## get the models data frame
as.data.frame(test)
```

 BayesMfp

Bayesian model inference for multiple fractional polynomial models

Description

Bayesian model inference for multiple fractional polynomial models is conducted by means of either exhaustive model space evaluation or posterior model sampling.

Usage

```
BayesMfp(formula = formula(data), data = parent.frame(), family =
gaussian, priorSpecs = list(a = 4, modelPrior = "flat"), method =
c("ask", "exhaustive", "sampling"), subset = NULL, na.action = na.omit,
verbose = TRUE, nModels = NULL, nCache=1e9L, chainlength = 1e5L)
```

```
bfp(x, max = 2, scale = TRUE, rangeVals=NULL)
```

```
uc(x)
```

Arguments

formula	model formula
data	optional data.frame for model variables (defaults to the parent frame)
family	distribution and link: only gaussian("identity") supported at the moment

priorSpecs	prior specifications, see details
method	which method should be used to explore the posterior model space? (default: ask the user)
subset	optional subset expression
na.action	default is to skip rows with missing data, and no other option supported at the moment
verbose	should information on computation progress be given? (default)
nModels	how many best models should be saved? (default: 1% of the explored models or the chainlength, 1 would mean only the maximum a posteriori [MAP] model)
nCache	maximum number of best models to be cached at the same time during the model sampling (only has an effect if sampling has been chosen as method)
chainlength	length of the model sampling chain (only has an effect if sampling has been chosen as method)
x	variable
max	maximum degree for this FP (default: 2)
scale	use pre-transformation scaling to avoid numerical problems? (default)
rangeVals	extra numbers if the scaling should consider values in this range. Use this argument if you have test data with larger range than the training range.

Details

The formula is of the form $y \sim \text{bfp}(x_1, \text{max} = 4) + \text{uc}(x_2 + x_3)$, that is, the auxiliary functions `bfp` and `uc` must be used for defining the fractional polynomial and uncertain fixed form covariates terms, respectively. There must be an intercept, and no other fixed covariates are allowed. All `max` arguments of the `bfp` terms must be identical.

The prior specifications are a list:

a hyperparameter for hyper-g prior which must be greater than 3 and is recommended to be not greater than 4 (default is 4)

modelPrior choose if a flat model prior (default, "flat"), a model prior favoring sparse models explicitly ("sparse"), or a dependent model prior ("dependent") should be used.

If `method = "ask"`, the user is prompted with the maximum cardinality of the model space and can then decide whether to use posterior sampling or the exhaustive model space evaluation.

Note that if you specify only one FP term, the exhaustive model search must be done, due to the structure of the model sampling algorithm. However, in reality this will not be a problem as the model space will typically be very small.

Value

Returns an object of class `BayesMfp` that inherits from `list`. It is essentially a list of models. Each model is a list and has the following components:

powers a list of numeric vectors, where each vector contains the powers of the covariate that its name denotes.

ucTerms	an integer vector of the indices of uncertain fixed form covariates that are present in the model.
logM	log marginal likelihood
logP	log prior probability
posterior	normalized posterior probability, and if model sampling was done, the frequency of the model in the sampling algorithm
postExpectedg	posterior expected covariance factor g
postExpectedShrinkage	posterior expected shrinkage factor $t=g/(g + 1)$
R2	usual coefficient of determination for the linear model

Subsetting the object with `[.BayesMfp]` returns again a BayesMfp object with the same attributes, which are

numVisited	the number of models that have been visited (exhaustive search) or cached (model sampling)
inclusionProbs	BMA inclusion probabilities for all uncertain covariates
linearInclusionProbs	BMA probabilities for exactly linear inclusion of FP covariates
logNormConst	the (estimated) log normalizing constant $f(D)$
chainlength	length of the Markov chain, only present if method = "sampling"
call	the original call
formula	the formula by which the appropriate untransformed design matrix can be extracted
x	the shifted and scaled design matrix for the data
xCentered	the column-wise centered x
y	the response vector
yMean	the mean of the response values
SST	sum of squares total
indices	a list with components that describe the positions of uncertain covariate groups, fractional polynomial terms and fixed variables in the design matrix
termNames	a list of character vectors containing the names of uncertain covariate groups, fractional polynomial terms and fixed variables
shiftScaleMax	matrix with 4 columns containing preliminary transformation parameters, maximum degrees and cardinalities of the powersets of the fractional polynomial terms
priorSpecs	the utilized prior specifications
randomSeed	if a seed existed at function call (<code>get(".Random.seed", .GlobalEnv)</code>), it is saved here

Note

logNormConst may be unusable due to necessary conversion from long double to double!

Various methods for posterior summaries are available.

See Also

[BayesMfp Methods](#), [BmaSamples](#)

Examples

```
## generate some data
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

## run an exhaustive model space evaluation with a flat model prior and
## a uniform prior (a = 4) on the shrinkage factor t = g/(1 + g):
test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 method="exhaustive")

test

## now the same with a *dependent* model prior:
test2 <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 priorSpecs = list(a = 4, modelPrior = "dependent"),
                 method="exhaustive")

test2
```

BayesMfp Methods

Other methods for BayesMfp objects

Description

Print the object (print), get fitted values (fitted) and corresponding residuals (residuals).

Usage

```
## S3 method for class 'BayesMfp'
print(x, ...)
## S3 method for class 'BayesMfp'
fitted(object, design = getDesignMatrix(object), post =
getPosteriorParms(object, design = design), ...)
## S3 method for class 'BayesMfp'
residuals(object, ...)
```

Arguments

x valid [BayesMfp](#) object
object valid [BayesMfp](#) object, only the first model will be used.

design	design matrix of the first model in the object, which can be supplied by the caller if it is computed beforehand
post	posterior parameters of the normal-gamma distribution (defaults to the posterior expected mean, marginalized over the covariance factor g)
...	unused

Author(s)

Daniel Saban\`es Bov\`e

See Also

[BayesMfp](#), [BmaSamples](#) [Methods](#)

Examples

```
## generate a BayesMfp object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 method="exhaustive")

## the print method
test

## extract fitted values and corresponding residuals
fitted(test)
residuals(test)
```

bmaPredict

BMA prediction for new data points

Description

Make a Bayesian model averaged prediction for new data points, from those models saved in a [BayesMfp](#) object.

Usage

```
bmaPredict(BayesMfpObject, postProbs = posteriors(BayesMfpObject), newdata)
```

Arguments

`BayesMfpObject` [BayesMfp](#) object with the models over which the predictions should be averaged

`postProbs` vector of posterior probabilities, which are then normalized to the weights of the model average (defaults to the normalized posterior probability estimates)

`newdata` new covariate data as `data.frame`

Value

The predicted values as a vector.

Note

Note that this function is not an S3 predict method for [BmaSamples](#) objects, but a function working on [BayesMfp](#) objects (because we do not need BMA samples to do BMA point predictions).

Author(s)

Daniel Saban\`es Bov\`e

See Also

[BmaSamples Methods](#)

Examples

```
## generate a BayesMfp object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 method="exhaustive")

## predict new responses at (again random) covariates
bmaPredict(test,
            newdata = list(x1 = rnorm(n=15),
                           x2 = rbinom(n=15, size=5, prob=0.2) + 1,
                           x3 = rexp(n=15)))
```

BmaSamples *Bayesian model averaging over multiple fractional polynomial models*

Description

Draw samples from the Bayesian model average over the models in saved in a [BayesMfp](#)-object.

Usage

```
BmaSamples(object, sampleSize = length(object) * 10, postProbs =
posterior(object), gridList = list(), gridSize = 203, newdata=NULL,
verbose = TRUE, includeZeroSamples=FALSE)
```

Arguments

object	valid BayesMfp object containing the models over which to average
sampleSize	sample size (default is 10 times the number of models)
postProbs	vector of posterior probabilities (will be normalized within the function, defaults to the normalized posterior probabilities)
gridList	optional list of appropriately named grid vectors for FP evaluation, default is a length (gridSize - 2) grid per covariate additional to the observed values (two are at the minimum and maximum)
gridSize	see above (default: 203)
newdata	new covariate data.frame with exactly the names (and preferably ranges) as before (default: no new covariate data)
verbose	should information on sampling progress be printed? (default)
includeZeroSamples	should the function and coefficient samples include zero samples, from models where these covariates are not included at all? (default: FALSE, so the zero samples are not included)

Value

Return an object of class `BmaSamples`, which is a list with various elements that describe the `BayesMfp` object over which was averaged, model frequencies in the samples, the samples themselves etc:

priorSpecs	the utilized prior specifications
termNames	a list of character vectors containing the names of uncertain covariate groups, fractional polynomial terms and fixed variables
shiftScaleMax	matrix with 4 columns containing preliminary transformation parameters, maximum degrees and cardinalities of the powersets of the fractional polynomial terms
y	the response vector

x	the shifted and scaled design matrix for the data
randomSeed	if a seed existed at function call (<code>get(".Random.seed", .GlobalEnv)</code>), it is saved here
modelFreqs	The table of model frequencies in the BMA sample
modelData	data frame containing the normalized posterior probabilities of the models in the underlying <code>BayesMfp</code> object, corresponding log marginal likelihoods, model prior probabilities, posterior expected covariance and shrinkage factors, coefficients of determination, powers and inclusions, and finally model average weights and relative frequencies in the BMA sample.
sampleSize	sample size
sigma2	BMA samples of the regression variance
shrinkage	BMA samples of the shrinkage factor
fixed	samples of the intercept
bfp	named list of the FP function samples, where each element contains one FP covariate and is a matrix (samples x grid), with the following attributes: whereObsVals where in the scaled grid are the originally observed covariate values? (integer vector of the indexes) scaledGrid numeric vector with the positions of the scaled grid points, corresponding to the columns of the samples matrix counter how often has this covariate been included in the BMA sample? (identical to the number of rows in the samples matrix)
uc	named list of the uncertain fixed form covariates, where each element contains the coefficient samples of one group: in a matrix with the attribute counter as number of samples in the rows, and the columns are appropriately named to correspond to the single design variables.
fitted	fitted values of all models in object, in a matrix with layout models x observations.
predictions	samples from the predictive distribution at the covariates given in <code>newdata</code>
predictMeans	means of the predictive distribution at the covariates given in <code>newdata</code>

See Also

[BmaSamples Methods](#), [BayesMfp](#)

Examples

```
## construct a BayesMfp object
set.seed(19)

x1 <- rnorm (n=15)
x2 <- rbinom (n=15, size=20, prob=0.5)
x3 <- rexp (n=15)

y <- rt (n=15, df=2)
```

```
test <- BayesMfp (y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 200, method="exhaustive")

## now draw samples from the Bayesian model average
testBma <- BmaSamples (test)
testBma

## We can also draw predictive samples for new data points, but then
## we need to supply the new data to BmaSamples:
newdata <- data.frame(x1 = rnorm(15),
                     x2 = rbinom(n=15, size=5, prob=0.2) + 1,
                     x3 = rexp(n=15))
testBma <- BmaSamples(test, newdata=newdata)
predict(testBma)

## test that inclusion of zero samples works
testBma <- BmaSamples (test, includeZeroSamples=TRUE)
testBma
```

BmaSamples Methods *Other methods for BmaSamples objects*

Description

Print the object (print), get fitted values (fitted) and corresponding residuals (residuals).

Usage

```
## S3 method for class 'BmaSamples'
print(x, ...)
## S3 method for class 'BmaSamples'
fitted(object, ...)
## S3 method for class 'BmaSamples'
residuals(object, ...)
```

Arguments

x	valid BmaSamples object
object	valid BmaSamples object
...	unused

Author(s)

Daniel Saban\`es Bov\`e

See Also

[predict.BmaSamples](#), [summary.BmaSamples](#)

Examples

```
## construct a BayesMfp object
set.seed(19)

x1 <- rnorm (n=15)
x2 <- rbinom (n=15, size=20, prob=0.5)
x3 <- rexp (n=15)

y <- rt (n=15, df=2)

test <- BayesMfp (y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 200, method="exhaustive")

## now draw samples from the Bayesian model average
testBma <- BmaSamples (test)

## the print method:
testBma

## the fitted method:
fitted(testBma)

## and the corresponding residuals:
residuals(testBma)
```

empiricalHpd

Construct an empirical HPD interval from samples

Description

Construct an empirical highest posterior density (HPD) interval from samples which have been drawn from the distribution of a quantity of interest.

Usage

```
empiricalHpd(theta, level)
```

Arguments

theta	the vector of samples
level	the credible level

Value

A vector with the estimated lower and upper bounds of the HPD interval.

Author(s)

Daniel Saban\`es Bov\`e

Examples

```
## draw standard normal variates
test <- rnorm(n=1000)

## estimate the 95% HPD interval with these samples:
empiricalHpd(theta=test, level=0.95)

## compare with true HPD:
qnorm(p=c(0.025, 0.975))
```

Extract.BayesMfp	<i>Extract method for BayesMfp objects</i>
------------------	--

Description

Extract a subset of models from a [BayesMfp](#) object.

Usage

```
## S3 method for class 'BayesMfp'
x[...]
```

Arguments

x	valid BayesMfp object
...	transports the indexes of the models

Author(s)

Daniel Saban\`es Bov\`e

See Also

[BayesMfp](#)

Examples

```
## generate a BayesMfp object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 method="exhaustive")
```

```
## extract the top ten models
test[1:10]
```

<code>findModel</code>	<i>Find a specific fractional polynomial model in a BayesMfp object</i>
------------------------	---

Description

Returns the index of the wished model if it is present in the model list, and otherwise returns NA.

Usage

```
findModel(model, BayesMfpObject)
```

Arguments

`model` the specific model: a list with entries `powers` and `ucTerms`
`BayesMfpObject` an object of class [BayesMfp](#)

Details

See [BayesMfp](#) for the description of a model.

Value

Index of `model` in `BayesMfpObject` if it is present in the model list, otherwise NA.

Note

The searched model must have exactly the same construction as the models in `BayesMfpObject`. See the example below for the recommended use.

Examples

```
## construct a BayesMfp object
set.seed(92)

x1 <- rnorm (15)
x2 <- rbinom (n=15, size=20, prob=0.6)
x3 <- rexp (15)
y <- rt (15, df=2)

test <- BayesMfp (y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels=2000, method="exhaustive")

## copy one model
myModel <- test[[1]]

## and modify it!
myModel$powers[["x2"]] <- c (1, 2)
stopifnot(identical(findModel (myModel, test),
                     31L))
```

getPosteriorParms	<i>Extract updated posterior parameters for the normal inverse gamma distribution from a model, given a shrinkage factor.</i>
-------------------	---

Description

Conditional on a fixed shrinkage factor $t=g/(g+1)$, the posterior joint distribution of the effects and the regression variance is normal inverse gamma. With this function, you can compute the parameters of this distribution.

Usage

```
getPosteriorParms(x, shrinkage=x[[1]]$postExpectedShrinkage,
                 design = getDesignMatrix(x))
```

Arguments

x	a valid BayesMfp -Object, only first list element will be recognized
shrinkage	shrinkage factor used in the computations (defaults to the posterior expected shrinkage factor in the model <code>x[1]</code>)
design	(centered) design matrix for the model

Value

A list with four parameters:

aStar	the first parameter of the inverse gamma distribution
VStar	the covariance matrix part of the multivariate normal distribution
mStar	the expectation of the multivariate normal distribution
bStar	the second parameter of the inverse gamma distribution

Author(s)

Daniel Saban\`es Bov\`e

Examples

```
## construct a BayesMfp object
set.seed(19)

x1 <- rnorm (n=15)
x2 <- rbinom (n=15, size=20, prob=0.5)
x3 <- rexp (n=15)

y <- rt (n=15, df=2)

test <- BayesMfp (y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 200, method="exhaustive")
```

```
## now get the posterior parameters of the third best model
getPosteriorParms(test[3])
```

inclusionProbs	<i>Compute (model averaged) posterior variable inclusion probabilities</i>
----------------	--

Description

Compute (model averaged) posterior inclusion probabilities for the uncertain variables (including FP variables) based on a [BayesMfp](#) object.

Usage

```
inclusionProbs(BayesMfpObject, postProbs = posteriors(BayesMfpObject, ind = 1))
```

Arguments

`BayesMfpObject` valid [BayesMfp](#) object
`postProbs` posterior probabilities to weight the models (defaults to the normalized probability estimates)

Value

Named numeric vector with the estimated variable inclusion probabilities. Note that these can differ noticeably from the “global” inclusion probabilities computed from all discovered models, from which only the best were saved in the [BayesMfp](#) object.

Author(s)

Daniel Saban\’es Bov\’e

Examples

```
## construct a BayesMfp object
set.seed(19)

x1 <- rnorm (n=15)
x2 <- rbinom (n=15, size=20, prob=0.5)
x3 <- rexp (n=15)

y <- rt (n=15, df=2)

test <- BayesMfp (y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 200, method="exhaustive")

## now get the local inclusion probabilities
local <- inclusionProbs(test)

## they can be compared with the global inclusion probabilities
local - attr(test, "inclusionProbs")
```

ozone

Ozone data from Breiman and Friedman, 1985

Description

This is the Ozone data discussed in Breiman and Friedman (JASA, 1985, p. 580). These data are for 330 days in 1976. All measurements are in the area of Upland, CA, east of Los Angeles.

Usage

`data(ozone)`

Format

A data frame with 366 observations on the following 13 variables.

`month` month of the year

`day` day of the month

`weekday` day of the week: a factor with levels Monday, Tuesday, Wednesday, Thursday, Friday, Saturday, Sunday

`hourAverageMax` maximum 1-hour average ozone level [ppm]

`pressure500Height` 500 millibar pressure height [meters]

`windSpeed` wind speed [mph]

`humidity` relative humidity [%]

`tempSandburg` temperature at Sandberg, CA [degrees F]

`tempElMonte` temperature at El Monte, CA [degrees F]

`inversionBaseHeight` inversion base height [feet]

`pressureGradientDaggett` pressure gradient from LAX to Daggett, CA [mm Hg]

`inversionBaseTemp` inversion base temperature [degrees F]

`visibility` visibility [miles]

Source

Breiman, L and Friedman, J. (1985), "Estimating Optimal Transformations for Multiple Regression and Correlation", *Journal of the American Statistical Association*, 80, 580-598.

plotCurveEstimate *Generic function for plotting a fractional polynomial curve estimate*

Description

Plot a fractional polynomial curve estimate for either a single model or a Bayesian model average over [BayesMfp](#) objects. Optionally, credible intervals and / or bands can be added to the plot.

Usage

```
plotCurveEstimate(model, termName, plevel = 0.95, slevel = plevel,
  plot = TRUE, legendPos = "topleft", rug = FALSE, partialResids=TRUE,
  hpd=TRUE,..., main = NULL)
```

Arguments

model	an object of class BayesMfp or BmaSamples
termName	string denoting an FP term, as written by the summary method
plevel	credible level for pointwise intervals, and NULL means no pointwise intervals (default: 0.95)
slevel	credible level for simultaneous credible band (SCB), NULL means no SCB (defaults to plevel)
plot	if FALSE, only return values needed to produce the plot, but do not plot (default is TRUE, so a plot is made)
legendPos	position of coefficient estimates (for BayesMfp) or sample size (for BmaSamples) in the plot, NULL suppresses the printing (default is "topleft")
rug	add a rug to the plot? (default: FALSE)
partialResids	add partial residuals to the plot? (default: TRUE)
hpd	use HPD intervals (TRUE, default) or quantile-based (FALSE) intervals?
...	further arguments in case of a BayesMfp object (see details) and arguments for plotting with matplot
main	optional main argument for the plot

Details

Further arguments for application on a [BayesMfp](#) object:

grid vector of unscaled abscissae, default is a length `gridSize` grid over the observed range specified by providing the argument NULL.

post list with posterior parameters of the model, which may be provided manually to accelerate plotting in a loop

gridSize default number of grid points used when no `grid` is supplied (default is 201)

numSim number of simulations for estimation of the SCB (default is 500)

Value

a list of various plotting information:

original	grid on the original covariate scale
grid	grid on the transformed scale
mode	mode curve values, only for BayesMfp object
mean	pointwise mean curve values, only for BmaSamples object
median	pointwise median curve values, only for BmaSamples object
plower	lower boundaries for pointwise intervals
pupper	upper boundaries for pointwise intervals
slower	lower boundaries for SCB
supper	upper boundaries for SCB
obsVals	observed values of the covariate on the original scale
sampleSize	sample size underlying the curve estimate, only for BmaSamples object
partialResids	partial residuals
transform	vector of shift and scale parameter

See Also

[BayesMfp](#), [BmaSamples](#)

Examples

```
## construct a BayesMfp object
set.seed(19)
x1 <- rnorm (n=15)
x2 <- rbinom (n=15, size=20, prob=0.5)
x3 <- rexp (n=15)
y <- rt (n=15, df=2)

test <- BayesMfp (y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
method="exhaustive")

## plot the x2 curve estimate for the 20-th best model
p1 <- plotCurveEstimate (test[20], "x2")

## look at the returned list
str(p1)

## plot the BMA curve estimate for the same covariate
testBma <- BmaSamples (test)
p2 <- plotCurveEstimate (testBma, "x2")

## look at the returned list
str(p2)

## try the new options:
plotCurveEstimate (testBma, "x2", partialResids=FALSE, hpd=FALSE)
```

 posteriors

Extract posterior model probability estimates from BayesMfp objects

Description

Extract posterior model probability estimates (either normalized estimates or sampling frequencies) from [BayesMfp](#) objects.

Usage

```
posteriors(BayesMfpObject, ind = 1)
```

Arguments

`BayesMfpObject` a valid [BayesMfp](#) object, containing the models the probabilities of which one wants to estimate

`ind` `ind = 1` means normalized posteriors, `ind = 2` means sampling frequencies

Value

The vector of probability estimates.

Author(s)

Daniel Saban\`es Bov\`e

Examples

```
## construct a BayesMfp object
set.seed(19)
x1 <- rnorm (n=15)
x2 <- rbinom (n=15, size=20, prob=0.5)
x3 <- rexp (n=15)
y <- rt (n=15, df=2)

test <- BayesMfp (y ~ bfp (x1, max = 2) + bfp (x2, max = 2) + uc (x3), nModels = 100,
  method="exhaustive")

## this works:
posteriors(test)

## this must not work:
## SoDA::muststop(posteriors(test, ind=2))

## only if we do model sampling there are model frequencies:
test2 <- BayesMfp (y ~ bfp (x1, max = 2) + bfp (x2, max = 2) + uc (x3), nModels = 100,
  method="sampling")
posteriors(test2, ind=2)
```

predict.BayesMfp *Predict method for BayesMfp objects*

Description

Predict new responses from a single multiple FP model.

Usage

```
## S3 method for class 'BayesMfp'  
predict(object, newdata, ...)
```

Arguments

object	valid BayesMfp object, from which only the first model will be used.
newdata	new covariate data with exactly the names (and preferably ranges) as for the original BayesMfp call
...	unused

Author(s)

Daniel Saban\`es Bov\`e

See Also

[bmaPredict](#)

Examples

```
## generate a BayesMfp object  
set.seed(19)  
  
x1 <- rnorm(n=15)  
x2 <- rbinom(n=15, size=20, prob=0.5)  
x3 <- rexp(n=15)  
  
y <- rt(n=15, df=2)  
  
test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,  
                  method="exhaustive")  
  
## predict new responses at (again random) covariates  
predict(test,  
      newdata = list(x1 = rnorm (15),  
                      x2 = rbinom (n=15, size=5, prob=0.2) + 1,  
                      x3 = rexp (15)))
```

predict.BmaSamples *Predict method to extract point and interval predictions from BmaSamples objects*

Description

Predict new responses from a Bayesian model average over FP models, from which predictive samples have already been produced.

Usage

```
## S3 method for class 'BmaSamples'
predict(object, level=0.95, hpd=TRUE, ...)
## S3 method for class 'predict.BmaSamples'
print(x, ...)
```

Arguments

object	valid BmaSamples object
level	credible level for the credible intervals (default: 95%)
hpd	should empirical hpd intervals be used (default) or simple quantile-based?
...	unused
x	object of S3 class predict.BmaSamples

Details

This function summarizes the predictive samples saved in the [BmaSamples](#) object. Using these functions, one can obtain predictive credible intervals, as opposed to just using the function [bmaPredict](#), which only gives the means of the predictive distributions.

Value

A list of class `predict.BmaSamples`, which has then a separate print method. The elements of the list are:

intervalType	which credible intervals have been computed (either “HPD” or “equitailed”)
level	the credible level
newdata	the covariate data for the predicted data points (just copied from object)
sampleSize	the sample size (just copied from object)
nModels	the number of models (just copied from object)
summaryMat	the summary matrix for the predictions, with median, mean, lower and upper credible interval borders.

Author(s)

Daniel Saban\`es Bov\`e

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

```
## generate a BmaSamples object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 method="exhaustive")

## predict new responses at (again random) covariates with BMA:
testBma <- BmaSamples(test,
                      newdata=data.frame(x1 = rnorm (15),
                                          x2 = rbinom (n=15, size=5, prob=0.2) + 1,
                                          x3 = rexp (15)))

predict(testBma)
```

scrBesag

*Simultaneous credible band computation (Besag algorithm)***Description**

Simultaneous credible band computation

Usage

scrBesag(samples, level=0.95)

Arguments

samples	m by n matrix where m is the number of parameters, n is the number of samples and hence each (multivariate) sample is a column in the matrix samples
level	the credible level (default: 0.95)

Details

Calculates a series of simultaneous credible bounds for one parameter type.

Value

matrix with 'lower' and 'upper' rows

Author(s)

Thomas Kneib

References

J. Besag, P. Green, D. Higdon, K. Mengersen (1995): Bayesian Methods for Nonlinear Classification and Regression, *Statistical Science* **10**/1, 3–66, doi: [10.1214/ss/1177010123](https://doi.org/10.1214/ss/1177010123)

scrHpd

*Calculate an SCB from a samples matrix***Description**

Calculate an SCB from a samples matrix, which minimizes the absolute distances of the contained samples to a mode vector, at each gridpoint. Therefore the SCB might be considered an “HPD SCB”.

Usage

```
scrHpd(samples, mode = apply(samples, 2, median), level = 0.95)
```

Arguments

samples	m by n matrix where m is the number of samples and n the number of parameters, hence each (multivariate) sample is a row in the matrix samples
mode	mode vector of length n (defaults to the vector of medians)
level	credible level for the SCB (default: 0.95)

Details

This function first computes the matrix of absolute distances of the samples to the mode vector. Then based on this distance matrix, a one-sided SCB as described in Besag et al. (1995) is computed, which is then mapped back to the samples.

Value

A matrix with rows “lower” and “upper”, with the lower and upper SCB bounds.

Author(s)

Daniel Saban\`es Bov\`e

References

Besag, J.; Green, P.; Higdon, D. \& Mengersen, K. (1995): “Bayesian computation and stochastic systems (with discussion)”, *Statistical Science*, 10, 3-66.

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

```
## create some samples
time <- 1:10
nSamples <- 50
samples <- t(replicate(nSamples,
                      time * rnorm(1) + rexp(1))) +
           rnorm(length(time) * nSamples)
matplot(time, t(samples), type="l", lty=1, col=1,
        xlab="time", ylab="response")

## now test the function: 50% credible band
scb <- scrHpd(samples, level=0.5)
matlines(time, t(scb), col=2, lwd=2, lty=1)
```

 Summary of BayesMfp object

Calculate and print the summary of a BayesMfp object

Description

Calculate and print the summary of a [BayesMfp](#) object, using S3 methods for the class.

Usage

```
## S3 method for class 'BayesMfp'
summary(object, level=0.95, table=TRUE,
        shrinkage=NULL, ...)
## S3 method for class 'summary.BayesMfp'
print(x, ...)
```

Arguments

object	a valid BayesMfp object
x	a return value of summary.BayesMfp
level	credible level for coefficients HPD intervals (default: 0.95)
table	should a data.frame of the models be included? (default)
shrinkage	shrinkage factor used, where NULL defaults to the posterior expected shrinkage factor
...	only used by summary.BayesMfp to pass arguments to as.data.frame.BayesMfp

Value

`summary.BayesMfp` returns a list with S3 class `summary.BayesMfp`, where the arguments “call”, “numVisited”, “termNames”, “shiftScaleMax”, “inclusionProbs”, “chainlength” (only for model sampling results) are copied from the attributes of the `BayesMfp` object, please see its help page for details.

The other elements are:

<code>dataframe</code>	the model overview as <code>data.frame</code> (only if <code>table=TRUE</code> was specified)
<code>localInclusionProbs</code>	local variable inclusion probability estimates
<code>nModels</code>	number of models contained in object

If there are multiple models in object, the list element `postProbs` contains the exact (for exhaustively explored model spaces) or estimated (if model sampling has been done) posterior model probabilities.

If object contains only one FP model, then this one is summarized in more detail:

<code>level</code>	used credible level for coefficients HPD intervals
<code>shrinkage</code>	used shrinkage factor
<code>summaryMat</code>	matrix with posterior summaries of the single coefficients: “mode” gives the posterior mode, “HPDlower” and “HPDupper” give the boundaries of the HPD intervals with specified credible level
<code>sigma2Sum</code>	posterior summary for the regression variance: again mode, and lower and upper HPD bounds are given in a rowvector.

Note

Note that if you extract the summary of a single model with these functions, you ignore the uncertainty about the shrinkage factor $t=g/(g+1)$ by plugging in the number `shrinkage`. If you want to incorporate this uncertainty, you must run `BmaSamples` on this model and call the corresponding method `summary.BmaSamples`.

Author(s)

Daniel Saban\`es Bov\`e

See Also

[summary.BmaSamples](#)

Examples

```
## generate a BayesMfp object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)
```

```

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                method="exhaustive")

## summary of multiple models:
summary(test)

## summary of just one model (no. 10):
summary(test[10])

## internal structure is usually not interesting:
str(summary(test[10]))

```

Summary of *BmaSamples* object

*Calculate and print the summary of a *BmaSamples* object*

Description

Calculate and print the summary of a [BmaSamples](#) object, using S3 methods for the class.

Usage

```

## S3 method for class 'BmaSamples'
summary(object, level = 0.95, hpd = TRUE, ...)
## S3 method for class 'summary.BmaSamples'
print(x, table = TRUE, ...)

```

Arguments

<code>object</code>	a valid BmaSamples object
<code>level</code>	credible level for coefficients credible intervals
<code>hpd</code>	should empirical hpd intervals be used (default) or simple quantile-based?
<code>x</code>	a return value of summary.BmaSamples
<code>table</code>	should the model table been shown? (default)
<code>...</code>	unused

Value

The summary method returns an S3 object, where “sampleSize”, “modelData” and “modelFreqs” are copied from the [BmaSamples](#) object, please see its help page for the details. “intervalType” and “level” copy the function’s parameters.

“summaryMat” contains the posterior summaries for the intercept and uncertain fixed form covariates. “sigma2Sum” and “shrinkageSum” contain the posterior summaries for the regression variance and the shrinkage factor, respectively. The summaries are always the median, mean, lower and upper credible bounds for the coefficients.

Author(s)

Daniel Saban\`es Bov\`e

See Also

[summary.BayesMfp](#)

Examples

```
## generate a BmaSamples object
set.seed(19)

x1 <- rnorm(n=15)
x2 <- rbinom(n=15, size=20, prob=0.5)
x3 <- rexp(n=15)

y <- rt(n=15, df=2)

test <- BayesMfp(y ~ bfp (x2, max = 4) + uc (x1 + x3), nModels = 100,
                 method="exhaustive")

testBma <- BmaSamples(test)

## look at the summary
summary(testBma)

## and its structure
str(summary(testBma))
```

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