Package ‘HTLR’

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Description

Efficient Bayesian multinomial logistic regression based on heavy-tailed priors. This package is suitable for classification and feature selection with high-dimensional features, such as gene expression profiles. Heavy-tailed priors can impose stronger shrinkage (compared to Gaussian and Laplace priors) to the coefficients associated with a large number of useless features, but still allow coefficients of a small number of useful features to stand out without punishment. It can also automatically make selection within a large number of correlated features. The posterior of coefficients and hyper-parameters is sampled with restricted Gibbs sampling for leveraging high-dimensionality and Hamiltonian Monte Carlo for handling high-correlations among coefficients.

References

as.matrix.htlr.fit  

Create a Matrix of Markov Chain Samples

Description

The Markov chain samples (without warmup) included in a htlr.fit object will be coerced to a matrix.

Usage

## S3 method for class 'htlr.fit'
as.matrix(x, k = NULL, include.warmup = FALSE, ...)

Arguments

x  
An object of S3 class htlr.fit.

k  
Coefficients associated with class k will be drawn. Must be a positive integer in 1,2,...,C-1 for C-class traning labels (base class 0 can not be chosen). By default the last class is selected. For binary logistic model this argument can be ignored.

include.warmup  
Whether or not to include warmup samples

...  
Not used.

Value

A matrix with \((p + 1)\) columns and \(i\) rows, where \(p\) is the number of features excluding intercept, and \(i\) is the number of iterations after burnin.

Examples

## No. of features used: 100; No. of iterations after burnin: 15
fit <- htlr(X = colon$X, y = colon$y, fsel = 1:100, iter = 20, warmup = 5)
dim(as.matrix(fit))

colon  
Colon Tissues

Description

In this dataset, expression levels of 40 tumor and 22 normal colon tissues for 6500 human genes are measured using the Affymetrix technology. A selection of 2000 genes with highest minimal intensity across the samples has been made by Alon et al. (1999). The data is preprocessed by carrying out a base 10 logarithmic transformation and standardizing each tissue sample to zero mean and unit variance across the genes.
Usage

data("colon")

Format

A list contains data matrix $X$ and response vector $y$:

- $X$ A matrix with 66 rows (observations) and 2000 columns (features).
- $y$ A binary vector where 0 indicates normal colon tissues and 1 indicates tumor colon tissues.

References


Pima Indians Diabetes

Description

This dataset is originally from the National Institute of Diabetes and Digestive and Kidney Diseases. The objective of the dataset is to diagnostically predict whether or not a patient has diabetes, based on certain diagnostic measurements included in the dataset. Several constraints were placed on the selection of these instances from a larger database. In particular, all patients here are females at least 21 years old of Pima Indian heritage. Different from the UCI original version, the dataset has been preprocessed such that rows with missing values are removed, and features are scaled.

Usage

data("diabetes392")

Format

A list contains data matrix $X$ and response vector $y$:

- $X$ A matrix with 392 rows (observations) and 8 columns (features).
- $y$ A binary vector where 1 indicates diabetes patients and 0 for otherwise.

Source


References

**evaluate_pred**

*Evaluate Prediction Results*

**Description**

This function compares the prediction results returned by a classifier with ground truth, and finally gives a summary of the evaluation.

**Usage**

```r
evaluate_pred(y.pred, y.true, caseid = names(y.true), showplot = TRUE, ...)
```

**Arguments**

- `y.pred`: A matrix of predicted probabilities, as returned by a classifier.
- `y.true`: Ground truth labels vector.
- `caseid`: The names of test cases which we take account of. By default all test cases are used for evaluation.
- `showplot`: Logical; if TRUE, a summary plot will be generated.
- `...`: Not used.

**Value**

A summary of evaluation result.

---

**gendata_FAM**

*Generate Simulated Data with Factor Analysis Model*

**Description**

This function generates inputs \( X \) given by the response variable \( y \) using a multivariate normal model.

**Usage**

```r
gedata_FAM(n, muj, A, sd_g = 0, stdx = FALSE)
```
Arguments

n  
Number of observations.

muj  
C by p matrix, with row c representing y = c, and column j representing x_j. Used to specify y.

A  
Factor loading matrix of size p by p, see details.

sd_g  
Numeric value indicating noise level δ, see details.

stdx  
Logical; if TRUE, data X is standardized to have mean = 0 and sd = 1.

Details

The means of each covariate x_j depend on y specified by the matrix muj; the covariate matrix Σ of the multivariate normal is equal to AA^t δ^2 I, where A is the factor loading matrix and δ is the noise level.

Value

A list contains input matrix X, response variables y, covariate matrix SGM and muj (standardized if stdx = TRUE).

See Also

gendata_MLR

Examples

## feature #1: marginally related feature
## feature #2: marginally unrelated feature, but feature #2 is correlated with feature #1
## feature #3-5: marginally related features and also internally correlated
## feature #6-10: noise features without relationship with the y

set.seed(12345)
n <- 100
p <- 10

means <- rbind(  
c(0, 1, 0),  
c(0, 0, 0),  
c(0, 0, 1),  
c(0, 0, 1),  
c(0, 0, 1))  
  * 2
means <- rbind(means, matrix(0, p - 5, 3))

A <- diag(1, p)
A[1:5, 1:3] <- rbind(  
c(1, 0, 0),  
c(2, 1, 0),  
c(0, 0, 1),  
c(0, 0, 1),  
c(0, 0, 1),  
c(0, 0, 1),  
c(0, 0, 1),  
c(0, 0, 1),  
c(0, 0, 1))
**gendata_MLR**

Generate Simulated Data with Multinomial Logistic Regression Model

**Description**

This function generates the response variables $y$ given optional supplied $X$ using a multinomial logistic regression model.

**Usage**

```r
gedata_MLR(n, p, NC = 3, nu = 2, w = 1, X = NULL, betas = NULL)
```

**Arguments**

- `n` Number of observations.
- `p` Number of features.
- `NC` Number of classes for response variables.
- `nu, w` If `betas` is not supplied (default), the regression coefficients are generated with t prior with df = `nu`, scale = $\sqrt{w}$; will be ignored if `betas` is supplied.
- `X` The design matrix; will be generated from standard normal distribution if not supplied.
- `betas` User supplied regression coefficients.

**Value**

A list contains input matrix $X$, response variables $y$, and regression coefficients $\delta$.

**See Also**

- `gendata_FAM`

**Examples**

```r
set.seed(12345)
dat <- gendata_MLR(n = 100, p = 10)
ggplot2::qplot(dat$y, bins = 6)
corrplot::corrplot(cor(dat$X))
```
htlr

Fit a HTLR Model

Description

This function trains linear logistic regression models with HMC in restricted Gibbs sampling.

Usage

htlr(
  X,
  y,
  fsel = 1:ncol(X),
  stdx = TRUE,
  prior = "t",
  df = 1,
  iter = 2000,
  warmup = floor(iter/2),
  thin = 1,
  init = "lasso",
  leap = 50,
  leap.warm = floor(leap/10),
  leap.stepsize = 0.3,
  cut = 0.05,
  verbose = FALSE,
  rep.legacy = FALSE,
  keep.warmup.hist = FALSE,
  ...
)

Arguments

X
Input matrix, of dimension nobs by nvars; each row is an observation vector.

y
Vector of response variables. Must be coded as non-negative integers, e.g.,
1,2,...,C for C classes, label 0 is also allowed.

fsel
Subsets of features selected before fitting, such as by univariate screening.

stdx
Logical; if TRUE, the original feature values are standardized to have
mean = 0 and sd = 1.

prior
The prior to be applied to the model. Either a list of hyperparameter settings
returned by htlr_prior or a character string from "t" (student-t), "ghs" (horse-
shoe), and "neg" (normal-exponential-gamma).

df
The degree freedom of t/ghs/neg prior for coefficients. Will be ignored if the
configuration list from htlr_prior is passed to prior.

iter
A positive integer specifying the number of iterations (including warmup).
**warmup**
A positive integer specifying the number of warmup (aka burnin). The number of warmup iterations should not be larger than iter and the default is \( \text{iter} / 2 \).

**thin**
A positive integer specifying the period for saving samples.

**init**
The initial state of Markov Chain; it accepts three forms:

- a previously fitted fithtlr object,
- a user supplied initial coefficient matrix of \((p+1)\times K\), where \(p\) is the number of features, \(K\) is the number of classes in \(y\) minus 1,
- a character string matches the following:
  - "lasso" - (Default) Use Lasso initial state with \(\lambda\) chosen by cross-validation. Users may specify their own candidate \(\lambda\) values via optional argument \(\text{lasso.lambda}\). Further customized Lasso initial states can be generated by \(\text{lasso.deltas}\).
  - "bcbc" - Use initial state generated by package BCBSF (Bias-corrected Bayesian classification). Further customized BCBSF initial states can be generated by \(\text{bcbsf.deltas}\). WARNING: This type of initial states can be used for continuous features such as gene expression profiles, but it should not be used for categorical features such as SNP profiles.
  - "random" - Use random initial values sampled from \(N(0, 1)\).

**leap**
The length of leapfrog trajectory in sampling phase.

**leap.warm**
The length of leapfrog trajectory in burnin phase.

**leap.stepsize**
The integrator step size used in the Hamiltonian simulation.

**cut**
The coefficients smaller than this criteria will be fixed in each HMC updating step.

**verbose**
Logical; setting it to \(\text{TRUE}\) for tracking MCMC sampling iterations.

**rep.legacy**
Logical; if \(\text{TRUE}\), the output produced in HTLR versions up to legacy-3.1-1 is reproduced. The speed will be typically slower than non-legacy mode on multi-core machine. Default is \(\text{FALSE}\).

**keep.warmup.hist**
Warmup iterations are not recorded by default, set \(\text{TRUE}\) to enable it.

**Value**
An object with S3 class \(\text{htlr.fit}\).

**References**
Examples

```r
set.seed(12345)
data("colon")

## fit HTLR models with selected features, note that the chain length setting is for demo only

## using t prior with 1 df and log-scale fixed to -10
fit.t <- htlr(X = colon$X, y = colon$y, fsel = 1:100,
prior = htlr_prior("t", df = 1, logw = -10),
init = "bcbc", iter = 20, thin = 1)

## using NEG prior with 1 df and log-scale fixed to -10
fit.neg <- htlr(X = colon$X, y = colon$y, fsel = 1:100,
prior = htlr_prior("neg", df = 1, logw = -10),
init = "bcbc", iter = 20, thin = 1)

## using horseshoe prior with 1 df and auto-selected log-scale
fit.ghs <- htlr(X = colon$X, y = colon$y, fsel = 1:100,
prior = "ghs", df = 1, init = "bcbc",
iter = 20, thin = 1)
```

htlr_prior

Generate Prior Configuration

Description

Configure prior hyper-parameters for HTLR model fitting

Usage

```r
htlr_prior(
    ptype = c("t", "ghs", "neg"),
    df = 1,
    logw = -(1/df) * 10,
    eta = ifelse(df > 1, 3, 0),
    sigmab0 = 2000
)
```

Arguments

- **ptype**
  The prior to be applied to the model. Either "t" (student-t, default), "ghs" (horseshoe), or "neg" (normal-exponential-gamma).

- **df**
  The degree freedom (aka alpha) of t/ghs/neg prior for coefficients.

- **logw**
  The log scale of priors for coefficients.

- **eta**
  The sd of the normal prior for logw. When it is set to 0, logw is fixed. Otherwise, logw is assigned with a normal prior and it will be updated during sampling.

- **sigmab0**
  The sd of the normal prior for the intercept.
Details
The output is a configuration list which is to be passed to prior argument of htlr. For naive users, you only need to specify the prior type and degree freedom, then the other hyper-parameters will be chosen automatically. For advanced users, you can supply each prior hyper-parameters by yourself. For suggestion of picking hyper-parameters, see references.

Value
A configuration list containing ptype, alpha, logw, eta, and sigmab0.

References

nzero_idx
Get Indices of Non-Zero Coefficients

Description
Get the indices of non-zero coefficients from fitted HTLR model objects.

Usage
nzero_idx(fit, cut = 0.1)

Arguments
fit An object of S3 class htlr.fit.
cut Threshold on relative SDB to distinguish zero coefficients.

Value
Indices vector of non-zero coefficients in the model.

Examples
set.seed(12345)
data("colon")

fit <- htlr(X = colon$X, y = colon$y, fsel = 1:100, iter = 20)
nzero_idx(fit)
predict.htlr.fit  Make Prediction on New Data

Description

Similar to other predict methods, this function returns predictions from a fitted htlrfit object.

Usage

```r
## S3 method for class 'htlr.fit'
predict(object, newx, type = c("response", "class"), ...)
```

Arguments

- `object`: A fitted model object with S3 class htlrfit.
- `newx`: A Matrix of values at which predictions are to be made.
- `type`: Type of prediction required. Type "response" gives the fitted probabilities. Type "class" produces the class label corresponding to the maximum probability.
- `...`: Advanced options to specify the Markov chain iterations used for inference. See htlr_predict.

Value

The object returned depends on type.

split_data  Split Data into Train and Test Partitions

Description

This function splits the input data and response variables into training and testing parts.

Usage

```r
split_data(X, y, p.train = 0.7, n.train = round(nrow(X) * p.train))
```

Arguments

- `X`: Input matrix, of dimension nobs by nvars; each row is an observation vector.
- `y`: Vector of response variables.
- `p.train`: Percentage of training set.
- `n.train`: Number of cases for training; will override p.train if specified.
Value

List of training data \( x.\text{tr}, y.\text{tr} \) and testing data \( x.\text{te}, y.\text{te} \).

Examples

dat <- gendata_MLR(n = 100, p = 10)
dat <- split_data(dat$X, dat$y, p.train = 0.7)
dim(dat$x.tr)
dim(dat$x.te)

std

Standardizes a Design Matrix

Description

This function accepts a design matrix and returns a standardized version of that matrix, the statistics of each column such as median and sd are also provided.

Usage

\[
\text{std}(X, \text{tol} = 1e-06)
\]

Arguments

\( X \)  
Design matrix, of dimension \( nobs \) by \( nvars \); each row is an observation vector; can also be an object that can be coerced to a matrix, e.g. a data.frame.

\( \text{tol} \)  
The tolerance value; a column of \( X \) is considered as singular if the sd of its entries (observations) is less than \( \text{tol} \). Singular columns will be dropped by the end.

Details

For each column of \( X \), the standardization is done by first subtracting its median, then dividing by its sample standard deviation, while the original version in ncvreg uses mean and population standard deviation. Its speed is slower than ncvreg because of the complexity of median finding, but still substantially faster than \text{scale()} provided by R base.

Value

The standardized design matrix with the following attributes:

- **nonsingular**  Indices of non-singular columns.
- **center**  Median of each non-singular column which is used for standardization.
- **scale**  Standard deviation of each non-singular column which is used for standardization.
Author(s)

Patrick Breheny (original)
Steven Liu (modification)

See Also


Examples

```r
set.seed(123)
mat <- matrix(rnorm(n = 80 * 90, mean = 100, sd = 50), 80, 90)
mat %>% as.numeric() %>% ggplot2::qplot(bins = 30, xlab = "Var")
mat %>% std() %>% as.numeric() %>% ggplot2::qplot(bins = 30, xlab = "Var")
```

Description

This function gives a summary of posterior of parameters.

Usage

```r
## S3 method for class 'htlr.fit'
summary(
  object,
  features = 1L:object$p,
  method = median,
  usedmc = get_sample_indice(dim(object$mcdeltas)[3], object$mc.param$iter.rmc),
  ...
)
```

Arguments

- `object`: An object of S3 class htlr.fit.
- `features`: A vector of indices (int) or names (char) that specify the parameters we will look at. By default all parameters are selected.
- `method`: A function that is used to aggregate the MCMC samples. The default is median, other built-in/customized statistical functions such as mean, sd, and mad can also be used.
- `usedmc`: Indices of Markov chain iterations used for inference. By default all iterations are used.
- `...`: Not used.
Value

A point summary of MCMC samples.

Examples

```r
set.seed(12345)
data("colon")

fit <- htlr(X = colon$X, y = colon$y, fsel = 1:100, iter = 20)
summary(fit, features = 1:16)
```
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