Package 'npmr'

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Type Package

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Title Nuclear Penalized Multinomial Regression
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Description Fit multinomial logistic regression with a penalty on the nuclear norm of the estimated regression coefficient matrix, using proximal gradient descent.
License GPL-2
NeedsCompilation no
R topics documented:
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Nuclear penalized multinomial regression

Description

As an alternative to an 11- or 12-penalty on multinomial logistic regression, this package fits multinomial regression with a penalty on the nuclear norm of the fitted regression coefficient matrix. The result is often a matrix of reduced rank, leveraging structure among the response classes so that the likelihood of one class informs the likelihood of other classes. Proximal gradient descent is used to solve the NPMR optimization problem.

Details

Package: npmr Type: Package Version: 1.0

Date: 2016-02-05

License: What license is it under?

The primary functions in the package are npmr, which solves nuclear penalized multinomial regression for a sequence of input values for the regularization parameter lambda, and cv.npmr, which chooses the optimal value of the regularization parameter lambda via cross validation. Both npmr and cv.npmr have predict and plot methods.

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

Maintainer: Scott Powers <sspowers@stanford.edu>

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

Examples

```
# Fit NPMR to simulated data

K = 5
n = 1000
m = 10000
p = 10
r = 2

# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)  # low-rank coefficient matrix
X = matrix(rnorm(n*p), n, p)  # covariate matrix with iid Gaussian entries
```

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```
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = npmr(X, Y, lambda = exp(seq(7, -2)))
# Print the NPMR fit:
fit2
# Produce a biplot:
plot(fit2, lambda = 20)
# Compute mean test error using the predict function (for each value of lambda):
getloss = function(pred, Y) {
    -mean(log(rowSums(Y*pred)))
apply(predict(fit2, Xtest), 3, getloss, Ytest)
```

cv.npmr

Cross-validated nuclear penalized multinomial regression

Description

Divide the training data into folds. Hold out each fold and fit NPMR for a range of regularization values on the remaining data, testing the result on the held-out fold. After the optimal value of the regularization parameter is determined, fit NPMR with this tuning parameter to the whole training set.

Usage

```
cv.npmr(X, Y, lambda = exp(seq(7, -2)), s = 0.1/max(X), eps = 1e-06,
    group = NULL, accelerated = TRUE, B.init = NULL, b.init = NULL,
    foldid = NULL, nfolds = 10)
```

Arguments

Χ

Covariate matrix. May be in sparse form from Matrix package

Υ

Multinomial reponse. May be (1) a vector of length equal to nrow(X), which will be interpreted as a factor, with levels representing response classes; or (2) a matrix with nrow(Y) = nrow(X), and each row has exactly one 1 representing the response class for that observation, with the remaining entries of the row being zero.

lambda

Vector of regularization parameter values for penalizing nuclear norm. Default is a wide range of values. We suggest that the user choose this sequence via trial and error. If the model takes too long to fit, try larger values of lambda. If

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> cross validation error is strictly increasing or strictly decreasing over the range of lambda specified, try extending the range in the direction of the smallest cross validation error.

Step size for proximal gradient descent s

Convergence threshold. When relative change in the objective function after an eps

interation drops below this threshold, algorithm halts.

Vector of length equal to number of variables (ncol(X) and nrow(B)). Variables group

> in the same group indexed by a POSITIVE integer will be penalized together (the nuclear norm of the sub-matrix of the regression coefficients will be penalized). Variables without positive integers will NOT be penalized. Default is NULL, which means there are no sub-groups; nuclear norm of entire coefficient

matrix is penalized.

Logical. Should accelerated proximal gradient descent be used? Default is accelerated

TRUE.

B.init Initial value of the regression coefficient matrix for proximal gradient descent b.init Initial value of the regression intercept vector for proximal gradient descent foldid Vector of length equal to nrow(X). Specifies folds for cross validation.

nfolds Number of folds for cross validation. Ignored if foldid is specified. Default is

Value

An object of class "cv.npmr" with values:

call the call that produced this object

A vector of total cross validation error for each value of lambda error

fit An object of class npmr fitted to the entire training data using lambda.min

lambda.min The value of lambda with minimum cross validation error

lambda The input sequence of regularization parameter values for which cross validation

error was calculated

number of rows in the input covariate matrix X n

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

```
npmr, predict.cv.npmr, print.cv.npmr, plot.cv.npmr
```

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Examples

```
Fit NPMR to simulated data
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
X = matrix(rnorm(n*p), n, p)
                                # covariate matrix with iid Gaussian entries
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
fold = sample(rep(1:10, length = nrow(X)))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = cv.npmr(X, Y, lambda = exp(seq(7, -2)), foldid = fold)
# Print the NPMR fit:
fit2
# Produce a biplot:
plot(fit2)
# Compute mean test error using the predict function:
-mean(log(rowSums(Ytest*predict(fit2, Xtest))))
```

logL

Log-likelihood for multinomial regression model

Description

Computes the log-likelihood of the fitted regression parameters given the data observed. Intended for internal use only.

Usage

```
logL(B, b, X, Y)
```

Arguments

В

Regression coefficient matrix

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b	Regression intercept vector
Χ	Covariate matrix

Y Multinomial response matrix

Value

The log-likelihood of B and b given X and Y

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

See Also

nuclear, objective

npmr

Nuclear penalized multinomial regression

Description

Fit a multinomial logistic regression model for a sequence of regularization parameters penalizing the nuclear norm of the regression coefficient matrix.

Usage

```
npmr(X, Y, lambda = exp(seq(7, -2)), s = 0.1/max(X), eps = 1e-06, group = NULL, accelerated = TRUE, B.init = NULL, b.init = NULL, quiet = TRUE)
```

Arguments

Χ	Covariate matrix. May be in sparse form from Matrix package
Y	Multinomial reponse. May be (1) a vector of length equal to $nrow(X)$, which will be interpreted as a factor, with levels representing response classes; or (2) a matrix with $nrow(Y) = nrow(X)$, and each row has exactly one 1 representing the response class for that observation, with the remaining entries of the row being zero.
lambda	Vector of regularization parameter values for penalizing nuclear norm. Default is a wide range of values. We suggest that the user choose this sequence via trial and error. If the model takes too long to fit, try larger values of lambda.
S	Step size for proximal gradient descent
eps	Convergence threshold. When relative change in the objective function after an interation drops below this threshold, algorithm halts.
group	Vector of length equal to number of variables (ncol(X) and nrow(B)). Variables in the same group indexed by a POSITIVE integer will be penalized together (the nuclear norm of the sub-matrix of the regression coefficients will be penalized). Variables without positive integers will NOT be penalized. Default is NULL, which means there are no sub-groups; nuclear norm of entire coefficient matrix is penalized.

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accelerated	Logical. Should accelerated proximal gradient descent be used? Default is TRUE.
B.init	Initial value of the regression coefficient matrix for proximal gradient descent
b.init	Initial value of the regression intercept vector for proximal gradient descent
quiet	Logical. Should output be silenced? If not, print the value of the objective function after each step of proximal gradient descent. Perhaps useful for debugging. Default is TRUE.

Details

In multinomial regression (in contrast with Gaussian regression or logistic regression) there is a matrix of regression coefficients, not just a vector. NPMR fits a logistic multinomial regression with a penalty on the nuclear norm of this regression coefficient matrix B. Specifically, the objective is

```
-loglik(B,b|X,Y) + lambda*||B|| *
```

where \|\.\|_* denotes the nuclear norm. This implementation solves the problem using proximal gradient descent, which iteratively steps in the direction of the negative gradient of the loss function and soft-thresholds the singular values of the result.

This function makes available the option, through the groups argument, of dividing the regression coefficient matrix into sub-matrices (by row) and penalizing the sum of the nuclear norms of these submatrices. Rows (correspond to variables) can be given no penalty in this way.

Value

An object of class "npmr" with values:

call	the call that produced this object
В	A 3-dimensional array, with dimensions (ncol(X), ncol(Y), length(lambda)). For each lambda, this array stores the regression coefficient matrix which solves the NPMR optimization problem for that value of lambda.
b	A matrix with ncol(Y) rows and length(lambda) columns. Each column stores the regression intercept vector which solves the NPMR optimization problem for that value of lambda.
objective	A vector of length equal to the length of lambda, giving the value value of the objective for the solution corresponding to each value of lambda.
lambda	The input sequence of values for the regularization parameter, for each of which NPMR has been solved.

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

```
cv.npmr, predict.npmr, print.npmr, plot.npmr
```

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Examples

```
Fit NPMR to simulated data
K = 5
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
X = matrix(rnorm(n*p), n, p)
                                # covariate matrix with iid Gaussian entries
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = npmr(X, Y, lambda = exp(seq(7, -2)))
# Print the NPMR fit:
fit2
# Produce a biplot:
plot(fit2, lambda = 20)
# Compute mean test error using the predict function (for each value of lambda):
getloss = function(pred, Y) {
    -mean(log(rowSums(Y*pred)))
apply(predict(fit2, Xtest), 3, getloss, Ytest)
```

nuclear

Nuclear norm of a matrix

Description

Returns the nuclear norm of a matrix, which is the sum of its singular values, obtained through a singular value decomposition. Intended for internal use only.

Usage

```
nuclear(B)
```

Arguments

В

a matrix

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Value

the nuclear norm of the matrix

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

See Also

logL, objective, objectiveFast

objective

NPMR objective function

Description

Return the objective function of the data and the fitted parameters for nuclear penalized multinomial regression. The objective is the sum of the negative log-likelihood and the product of the regularization parameter and nuclear norm of the fitted regression coefficient matrix. Intended for internal use only.

Usage

```
objective(B, b, X, Y, lambda)
```

Arguments

В	fitted regression coefficient matrix
b	fitted regression intercept vector
Χ	covariate matrix

Y multinomial response matrix

lambda regularization parameter (maybe be a vector of values)

Value

a vector of objective values for the NPMR optimization problem, one for each value of lambda

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

```
logL, nuclear, objectiveFast
```

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objectiveFast	Shortcut computation for NPMR objective function	

Description

Computes the objective function for NPMR more quickly than objective by leveraging precomputed fitted values, which is the bottleneck in computing the objective. Intended for internal use only.

Usage

```
objectiveFast(B, P, W, lambda)
```

Arguments

n	C44 - J		CC -: 4	
D	muear	egression	coefficient	maurx

P matrix of fitted multinomial class probabilities

wector containing indices of P which correspond to observed data

lambda regularization parameter (maybe be a vector of values)

Value

a vector of objective values for the NPMR optimization problem, one for each value of lambda

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

See Also

objective, nuclear, PGDnpmr

PGDnpmr	Proximal gradient descent for nuclear penalized multinomial regression

Description

Iterates steps of proximal gradient descent until convergence, by repeatedly taking steps in the direction of the negative of the gradient and soft-thresholding the singular values of the result. Intended for internal use only.

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Usage

```
PGDnpmr(B, b, X, Y, lambda, s, group = NULL, accelerated = TRUE, eps = 1e-07, maxit = 1e+05, quiet = TRUE)
```

Arguments

В	Initial regression coefficient matrix
b	Initial intercept vector
Χ	Covariate matrix. May be in sparse form from Matrix package
Υ	Response matrix. Each row has exactly one 1 indicating response category for that observation. All other entries are zero.
lambda	Vector of regularization parameter values for penalizing nuclear norm
S	Step size for proximal gradient descent
group	Vector of length equal to number of variables (ncol(X) and nrow(B)). Variables in the same group indexed by a POSITIVE integer will be penalized together (the nuclear norm of the sub-matrix of the regression coefficients will be penalized). Variables without positive integers will NOT be penalized. Default is NULL, which means there are no sub-groups; nuclear norm of entire coefficient matrix is penalized.
accelerated	Logical. Should accelerated proximal gradient descent be used? Default is TRUE.
eps	Convergence threshold. When relative change in the objective function after an interation drops below this threshold, algorithm halts.
maxit	Maximum number of iterations for proximal gradient descent.
quiet	Logical. Should output be silenced? If not, print the value of the objective function after each step of proximal gradient descent. Perhaps useful for debugging. Default is TRUE.

Value

В	Optimal value of the regression coefficent matrix at convergence
b	Optimal value of the regression intercept vector at convergence
objectivePath	Vector showing the value of the objective function at each step in proximal gradient descent
time	Time taken until convergence

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

See Also

npmr, prox, objective, objectiveFast

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plot.cv.npmr

Visualize the regression coefficient matrix fit by cross-validated NPMR

Description

Plots features (in orange) by their weights on the first two latent variables in the singular value decomposition of the regression coefficient matrix. Plots response classes (as blue arrows) by their loadings on the first two latent variables. Does this for the regression coefficient matrix fit with the value of lambda that led to the minimum cross validation error among all those tried.

Usage

```
## S3 method for class 'cv.npmr'
plot(x, feature.names = TRUE, ...)
```

Arguments

```
    an object of class cv.npmr
    logical. Should the names of the covariates be used in the plot? If FALSE, use standard plotting symbol (pch=1) instead.
    additional arguments to be passed to plot
```

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

See Also

```
cv.npmr, plot.npmr
```

Examples

```
# Fit NPMR to simulated data

K = 5
n = 1000
m = 10000
p = 10
r = 2

# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)  # low-rank coefficient matrix
X = matrix(rnorm(n*p), n, p)  # covariate matrix with iid Gaussian entries
eta = X
```

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```
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
fold = sample(rep(1:10, length = nrow(X)))

# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))

# Fit NPMR for a sequence of lambda values without CV:
fit2 = cv.npmr(X, Y, lambda = exp(seq(7, -2)), foldid = fold)
# Produce a biplot:
plot(fit2)
```

plot.npmr

Visualize the regression coefficient matrix fit by cross-validated NPMR

Description

Plots features (in orange) by their weights on the first two latent variables in the singular value decomposition of the regression coefficient matrix. Plots response classes (as blue arrows) by their loadings on the first two latent variables. Does this for the regression coefficient matrix fit with the value of lambda closest among all those tried to the value of lambda specified.

Usage

```
## S3 method for class 'npmr'
plot(x, lambda, feature.names = TRUE, ...)
```

Arguments

x an object of class npmr

lambda a single regularization parameter value

feature.names logical. Should the names of the covariates be used in the plot? If FALSE, use

standard plotting symbol (pch=1) instead.

... additional arguments to be passed to plot

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

```
npmr, plot.cv.npmr
```

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Examples

```
Fit NPMR to simulated data
K = 5
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
                                # covariate matrix with iid Gaussian entries
X = matrix(rnorm(n*p), n, p)
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = npmr(X, Y, lambda = exp(seq(7, -2)))
# Produce a biplot:
plot(fit2, lambda = 20)
```

predict.cv.npmr

Make predictions from a "cv.npmr" object

Description

Return predicted reponse class probabilities from a cross-validated NPMR model, using the value of the regularization parameter that led to the minimum cross validation error

Usage

```
## S3 method for class 'cv.npmr'
predict(object, newx, ...)
```

Arguments

object an object of class cv.npmr

newx covariate matrix on which for which to make response class probability predic-

tions. Must have same number of columns as X used original to fit object.

... ignored

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Value

a matrix giving the predicted probability that each row of news belongs to each class, corresponding the value of the regularization parameter that led to minimum cross validation error.

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

See Also

```
cv.npmr, predict.npmr
```

Examples

```
Fit NPMR to simulated data
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
X = matrix(rnorm(n*p), n, p)
                               # covariate matrix with iid Gaussian entries
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
fold = sample(rep(1:10, length = nrow(X)))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = cv.npmr(X, Y, lambda = exp(seq(7, -2)), foldid = fold)
# Compute mean test error using the predict function:
-mean(log(rowSums(Ytest*predict(fit2, Xtest))))
```

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predict.npmr

Make predictions from a "npmr" object

Description

Return predicted reponse class probabilities from a fitted NPMR model, for each value of lambda on which the NPMR model was originally fit.

Usage

```
## S3 method for class 'npmr'
predict(object, newx, ...)
```

Arguments

object an object of class npmr

newx covariate matrix on which for which to make response class probability predic-

tions. Must have same number of columns as X used original to fit object.

... ignored

Value

a 3-dimensional array, with dimensions (nrow(newx), ncol(Y), length(lambda)). For each lambda, this array stores for that value of lambda the predicted response class probabilites for each observation.

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

See Also

```
npmr, predict.cv.npmr
```

Examples

```
# Fit NPMR to simulated data
K = 5
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
```

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```
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
                               # covariate matrix with iid Gaussian entries
X = matrix(rnorm(n*p), n, p)
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = npmr(X, Y, lambda = exp(seq(7, -2)))
# Compute mean test error using the predict function (for each value of lambda):
getloss = function(pred, Y) {
    -mean(log(rowSums(Y*pred)))
apply(predict(fit2, Xtest), 3, getloss, Ytest)
```

print.cv.npmr

summarize a "cv.npmr" object

Description

Print (1) the call that produced the cv.npmr object; (2) the value of the regularization parameter lambda that led to the minimum cross validation error; (3) the rank of the fitted regression coefficient matrix; and (4) the per-observation cross validation error using the optimal lambda.

Usage

```
## S3 method for class 'cv.npmr'
print(x, ...)
```

Arguments

```
x an object of class cv.npmr
... ignored
```

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

```
cv.npmr, print.npmr
```

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Examples

```
Fit NPMR to simulated data
K = 5
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
X = matrix(rnorm(n*p), n, p)
                                # covariate matrix with iid Gaussian entries
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
fold = sample(rep(1:10, length = nrow(X)))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = cv.npmr(X, Y, lambda = exp(seq(7, -2)), foldid = fold)
# Print the NPMR fit:
fit2
```

print.npmr

Summarize a "npmr" object

Description

Print the call that produced the npmr object and a dataframe showing, for each value of the regularization parameter on which the NPMR object was fit, the rank of the resulting regression coefficient matrix and the corresponding value of the NPMR objective function.

Usage

```
## S3 method for class 'npmr'
print(x, ...)
```

Arguments

```
x an object of class npmr
... ignored
```

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

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References

Scott Powers, Trevor Hastie and Rob Tibshirani (2016). "Nuclear penalized multinomial regression with an application to predicting at bat outcomes in baseball." In prep.

See Also

```
npmr, print.cv.npmr
```

Examples

```
# Fit NPMR to simulated data
K = 5
n = 1000
m = 10000
p = 10
r = 2
# Simulated training data
set.seed(8369)
A = matrix(rnorm(p*r), p, r)
C = matrix(rnorm(K*r), K, r)
B = tcrossprod(A, C)
                                # low-rank coefficient matrix
                                # covariate matrix with iid Gaussian entries
X = matrix(rnorm(n*p), n, p)
eta = X
P = exp(eta)/rowSums(exp(eta))
Y = t(apply(P, 1, rmultinom, n = 1, size = 1))
# Simulate test data
Xtest = matrix(rnorm(m*p), m, p)
etatest = Xtest
Ptest = exp(etatest)/rowSums(exp(etatest))
Ytest = t(apply(Ptest, 1, rmultinom, n = 1, size = 1))
# Fit NPMR for a sequence of lambda values without CV:
fit2 = npmr(X, Y, lambda = exp(seq(7, -2)))
# Print the NPMR fit:
fit2
```

prox

Proximal operator for nuclear norm

Description

Return the value of the proximal operator of the nuclear norm (scaled by threshold) applied to a matrix

Usage

```
prox(B, threshold, group)
```

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Arguments

B matrix

threshold scaling factor applied to the nuclear norm. In proximal gradient descent for

NPMR, this is the product of the stepsize and the regularization parameter lambda

group Vector of length equal to number of variables, i.e. nrow(B). Variables in the

same group indexed by a POSITIVE integer will be penalized together (the nuclear norm of the sub-matrix of the regression coefficients will be penalized). Variables without positive integers will NOT be penalized. Default is NULL, which means there are no sub-groups; nuclear norm of entire coefficient matrix

is penalized.

Value

the value of the proximal operator of the nuclear norm (scaled by threshold) applied to B

Author(s)

Scott Powers, Trevor Hastie, Rob Tibshirani

References

Neal Parikh and Stephen Boyd (2013) "Proximal algorithms." Foundations and Trends in Optimization 1, 3:123-231.

See Also

nuclear, PGDnpmr

Description

Vowel

Speaker independent recognition of the eleven steady state vowels of British English using a specified training set of lpc derived log area ratios.

Format

A data frame with 990 observations on the following 12 variables.

Vowel Recognition

y Class label indicating vowel spoken

subset a factor with levels test train

- x.1 a numeric vector
- x.2 a numeric vector
- x.3 a numeric vector
- x.4 a numeric vector
- x.5 a numeric vector
- x.6 a numeric vector
- x.7 a numeric vector

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- x.8 a numeric vector
- x.9 a numeric vector
- x.10 a numeric vector

Details

The speech signals were low pass filtered at 4.7kHz and then digitised to 12 bits with a 10kHz sampling rate. Twelfth order linear predictive analysis was carried out on six 512 sample Hamming windowed segments from the steady part of the vowel. The reflection coefficients were used to calculate 10 log area parameters, giving a 10 dimensional input space. For a general introduction to speech processing and an explanation of this technique see Rabiner and Schafer [RabinerSchafer78].

Each speaker thus yielded six frames of speech from eleven vowels. This gave 528 frames from the eight speakers used to train the networks and 462 frames from the seven speakers used to test the networks.

The eleven vowels, along with words demonstrating their sound, are: i (heed) I (hid) E (head) A (had) a: (hard) Y (hud) O (hod) C: (hoard) U (hood) u: (who'd) 3: (heard)

Source

https://archive.ics.uci.edu/ml/machine-learning-databases/undocumented/connectionist-bench/vowel/

References

D. H. Deterding, 1989, University of Cambridge, "Speaker Normalisation for Automatic Speech Recognition", submitted for PhD.

Examples

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summary(Vowel)

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