# Package 'pclogit'

April 17, 2021

Type Package

| based p   | zed conditional (unconditional) logistic regression using a network-<br>peanlty for matched (unmatched) case-control data with grouped or graph-<br>nined variables. |  |
|---|--|--|
| Version 0.3   |  |  |
| <b>Date</b> 2021-0  | 14-17  |  |
| Author Hok  | eun Sun  |  |
| Maintainer  | Hokeun Sun <hsun@pusan.ac.kr></hsun@pusan.ac.kr>   |  |
| Depends Ma  | atrix  |  |
| <b>Description</b> An efficient algorithm for fitting the regularization path and providing selection probabilities of individual variables for analysis of high-dimensional matched (unmatched) case-control data. The algorithm uses cyclical coordinate descent in a pathwise fashion. |  |  |
| License GPI   | L-2  |  |
| R topics  | documented:  |  |
| -   | pclogit  |  |
| Index   | 7  |  |
| pclogit   | penalized conditional (unconditional) logistic regression for grouped or graph-constrained variables   |  |
| Description   |  |  |

Fit a regularization path of conditional (unconditional) logistic regression model for a matched (un-

When predictors are correlated witin either a group or a network graph, Laplacian matrix can be

matched) case-control response at a grid of values for regularization parameter lambda.

imposed into the regularization procedure to efficiently select relevant variables.

2 pclogit

#### Usage

```
pclogit(x,y,stra=NULL,alpha=1.0,nlam=100,lambda=NULL,eps=NULL,
    maxit=100000,maxp=p,thre=1e-6,group=NULL,type=c("ring","fcon"),
    sgnc=NULL)
```

# Arguments

| X      | The design matrix $(n \times p)$ without an intercept. pclogit standardizes the data by default, but the coefficients are always returned on the original scale.  |
|--------|---|
| у      | The response variable coded as 1 for cases and 0 for the matched controls.  |
| stra   | A vector of consecutive integers indicating the stratum of each observation. Each stratum must have exact one case and at least one control. If not specified, pclogit fits an ordinary logistic regression.  |
| alpha  | The penalty mixing parameter with $0 \le \alpha \le 1$ and default is 1. See details.   |
| nlam   | The number of lambda values and default is 100.   |
| lambda | A user supplied sequence of lambda values. Typically, this is left unspecified, and the program automatically computes its own lambda sequence based on nlam and eps.   |
| eps    | The smallest value for lambda as a fraction of lambda.max. The value should be greater than 1E-5. The default is .0001 if $n>p$ and .05 if $n\leq p$ .  |
| maxit  | Maximum number of passes over the data for all lambda values, and default is 10^5.  |
| maxp   | Limit the maximum number of variables ever to be nonzero.   |
| thre   | Convergence threshold for coordinate descent algorithm. The default value is 1E-6.  |
| group  | Either an integer vector of group sizes or a symmetric adjacency matrix. group describes either grouped or graph structure of predictors x. If no information between predictors is available, i.e., not specified, the pclogit performs an elastic-net regularization procedure. See details.                        |
| type   | A type of grouping network when group is defined as a vector of group sizes. "ring" and "fcon" represent a ring and fully connected network, respectively. Default is "ring". See details.  |
| sgnc   | Signs of regression coefficients. This can be provided only if group is specified as either a list of a group size or an adjacency matrix. The estimated signs of ridge regression for $n \leq p$ or ordinary regression for $n > p$ can be used for an adaptive network-based regularization procedure. See details. |

# **Details**

The penalty function of pclogit is defined as

$$\alpha||\beta||_1 + (1 - \alpha)(\beta^T S^T L S \beta)/2,$$

where S is a p dimensional diagonal matrix with estimated signs of regression coefficients on its diagonal entries, and L is a Laplacian matrix describing a graph structure of covariates. This penalty is equivalent to the Lasso penalty if alpha=1. When group and sgnc are not defined, L and S in the penalty function are replaced by an identity matrix, respectively. In this case, pclogit performs an elastic-net regularization procedure since the second term of the penalty simply reduces to the  $l_2$ 

pclogit 3

norm of  $\beta$ .

If group sizes of predictors are listed in group, it is assumed that all variables of the same groups are linked with each other like a ring or a fully connected network. In this case, the Laplacian matrix forms a block-wise diagonal matrix. The signs of regression coefficients sgnc can provide more accuarate estimates in case some variables either in the same group or linked with each other have different signs of thier regression coefficients, where the coefficients are not expected to be loccally smooth.

#### Value

b0 Intercept sequence of length of lambda. This is present only if an ordinary

logistic regression is fit, i.e., stra was not defined.

strata The strata of obsevations if stra was defined.

beta The coefficient matrix with a dimension (p x nlam), stored in sparse column

format("CsparseMatrix")

lambda The actual sequence of lambda values used

df The number of nonzero coefficients for each value of lambda

nobs The number of observations, n

alpha The value of alpha used

iterations Total passes over the data summed over all lambda values

jerr The error flag, for warnings and errors (largely for internal debugging)

# Author(s)

Hokeun Sun <hsun@pusan.ac.kr>

#### References

- H. Sun and S. Wang (2012) *Penalized Logistic Regression for High-dimensional DNA Methylation Data with Case-Control Studies*, Bioinformatics 28(10), 1368–1375
- H. Sun and S. Wang (2012) Network-based Regularization for Matched Case-Control Analysis of High-dimensional DNA Methylation Data, Statistics in Medicine 32(12), 2127–2139

# **Examples**

```
n<-200
p<-1000
x<-matrix(rnorm(n*p),n,p)

# one-to-one matched set
y<-c(rep(0,n/2),rep(1,n/2))
st<-rep(seq(n/2),2)

# one-to-four matched set
y<-c(rep(0,4*n/5),rep(1,n/5))
st<-c(rep(0,4*n/5),rep(4,n/5)),rep(seq(n/5),1))

# a total of 100 groups each of which consists of 5, 10, or 20 members
gr<-c(rep(5,40),rep(10,40),rep(20,20))</pre>
```

4 sel.pclogit

```
# an example of adjacency matrix
adjm < -cov(x)
diag(adjm)<-0</pre>
adjm[abs(adjm) \le 0.3] \le 0
adjm[abs(adjm)>0.3]<-1
# an example of signs of coefficients
sg<-sign(rnorm(p))</pre>
# Lasso
g1<-pclogit(x,y,st)</pre>
# Elastic-net
g2<-pclogit(x,y,st,alpha=0.1)
# Ring network of grouped covariates
g3<-pclogit(x,y,st,alpha=0.1,group=gr)</pre>
# Fully connected network of grouped covariates
g4<-pclogit(x,y,st,alpha=0.1,group=gr,type="fcon")
# Graph-constrained covariates
g5<-pclogit(x,y,st,alpha=0.1,group=adjm)</pre>
# Adaptive graph-constrained covariates
g6<-pclogit(x,y,st,alpha=0.1,group=adjm,sgnc=sg)</pre>
```

sel.pclogit

selection probabilities of regression coefficients

# **Description**

The selection probability of each regression coefficient is computed based on resamplings.

# Usage

```
sel.pclogit(x,y,stra=NULL,...,psub=0.5,N.lam=5,K=100)
```

# Arguments

| Х     | The design matrix $(n \times p)$ without an intercept. pclogit standardizes the data by default, but the coefficients are always returned on the original scale.   |
|-------|--|
| У     | The response variable coded as 1 for cases and 0 for the matched controls.   |
| stra  | A vector of consecutive integers indicating the stratum of each observation. Each stratum must have exact one case and at least one control. If not specified, sel.pclogit fits an ordinary logistic regression. |
|       | Other arguments that can be passed to pclogit.   |
| psub  | The proportion of subsamples used for resamplings, and $psub \in [0.5, 1)$ . The defalut is $0.5$ .  |
| N.lam | The number of lambda values used for resamplings, and default is 5.  |
| K     | The number of resamplings, and default is 100.   |

sel.pclogit 5

#### **Details**

The half of the strata stra are randomly selected without replacement K times. For each replication, the paired (x,y) in the selected strata are only used for pclogit to find non-zero coefficients along with N.lam lambda values. The selection probability of each coefficient is then computed based on the proportion of non-zeros out of K replciations. In an ordinary logistic model, the half of cases and controls are selected each time.

#### Value

beta The selection prbabilities ( $p \times N.1am$ )

maxsel The maximum selection probability of each coefficient are listed in descending

order along with the corresponding variable.

1ambda The actual sequence of 1ambda values usedK The actual number of resamplings used

#### Author(s)

Hokeun Sun <hsun@pusan.ac.kr>

#### References

H. Sun and S. Wang (2012) Penalized Logistic Regression for High-dimensional DNA Methylation Data with Case-Control Studies, Bioinformatics 28(10), 1368–1375

H. Sun and S. Wang (2012) Network-based Regularization for Matched Case-Control Analysis of High-dimensional DNA Methylation Data, Statistics in Medicine 32(12), 2127–2139

#### **Examples**

```
n<-200
p<-1000
x<-matrix(rnorm(n*p),n,p)</pre>
# one-to-one matched set
y < -c(rep(0,n/2),rep(1,n/2))
st < -rep(seq(n/2), 2)
# one-to-four matched set
y < -c(rep(0, 4*n/5), rep(1, n/5))
st<-c(rep(seq(n/5), rep(4, n/5)), rep(seq(n/5), 1))
# a total of 100 groups each of which consists of 5, 10, or 20 members
gr<-c(rep(5,40),rep(10,40),rep(20,20))
# an example of adjacency matrix
adjm < -cov(x)
diag(adjm)<-0
adjm[abs(adjm) \le 0.3] \le 0
adjm[abs(adjm)>0.3]<-1
# an example of signs of coefficients
sg<-sign(rnorm(p))</pre>
# Lasso
```

sel.pclogit

```
g1<-sel.pclogit(x,y,st)
# Elastic-net
g2<-sel.pclogit(x,y,st,alpha=0.1)
# Ring network of grouped covariates
g3<-sel.pclogit(x,y,st,alpha=0.1,group=gr)
# Fully connected network of grouped covariates
g4<-sel.pclogit(x,y,st,alpha=0.1,group=gr,type="fcon")
# Graph-constrained covariates
g5<-sel.pclogit(x,y,st,alpha=0.1,group=adjm)
# Adaptive graph-constrained covariates
g6<-sel.pclogit(x,y,st,alpha=0.1,group=adjm,sgnc=sg)</pre>
```

# Index

```
pclogit, 1
sel.pclogit, 4
```