# Package 'TULIP'

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Title A Toolbox for Linear Discriminant Analysis with Penalties

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<b>Description</b> Integrates several popular high-dimensional methods based on Linear Discriminant Analysis (LDA) and provides a comprehensive and user-friendly toolbox for linear, semi-parametric and tensor-variate classification as mentioned in Yuqing Pan, Qing Mai and Xin Zhang (2019) <arxiv:1904.03469>. Functions are included for covariate adjustment, model fitting, cross validation and prediction.</arxiv:1904.03469>
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adjten         adjvec         catch         catch_matrix         csa       1         cv.catch       1         cv.dsda       1         cv.msda       1         cv.SeSDA       1

2 adjten

Index																							37
	SOS		•	 •		•	•	 •		•	•	 •	•	•	 •	•	 •	٠		•	 ٠	•	 35
	sim.tensor.cov																						
	sim.bi.vector																						 33
	SeSDA																						 32
	ROAD																						 31
	predict.SeSDA	٠.																					 30
	predict.msda.																						 29
	predict.dsda .																						 28
	predict.catch .																						 26
	msda																						 22
	getnorm																						 21
	GDS1615																						 20
	dsda.all									•									•				 18

adjten

Adjust tensor for covariates.

### Description

Adjusts tensor with respect to covariates to achieve a more accurate performance. Tensor depends on the covariates through a linear regression model. The function returns the coefficients of covariates in regression and adjusted tensor list for further classifier modeling. It estimates coefficients based on training data, and then adjusts training tensor. When testing data is provided, the function will automatically adjust testing data by learned coefficients as well.

### Usage

```
adjten(x, z, y, testx = NULL, testz = NULL, is.centered = FALSE)
```

### **Arguments**

X	Input tensor or matrix list of length $N$ , where $N$ is the number of observations. Each element of the list is a tensor or matrix. The order of tensor can be any integer not less than 2.
Z	Input covariate matrix of dimension $N \times q$ , where $q < N$ . Each row of z is an observation.
У	Class label vector of dimention $N \times 1$ . For K class problems, y takes values in $\{1, \cdots, {\rm K}\}.$
testx	Input testing tensor or matrix list. Each element of the list is a test case. When testx is not provided, the function will only adjust training data.
testz	Input testing covariate matrix with each row being an observation.
is.centered	Indicates whether the input tensor and covariates have already been centered by their within class mean or not. If is.centered is FALSE, the function adjten will center data by class. If is.centered is TRUE, the function will skip the centering step.

adjten 3

#### **Details**

The model CATCH assumes the linear relationship bewteen covariates and tensor as

$$\mathbf{X} = \boldsymbol{\mu}_k + \boldsymbol{\alpha} \overline{\times}_{M+1} \mathbf{Z} + \mathbf{E},$$

where  $\alpha$  is the matrix of estimated coefficient of covariates. The function removes the effects of covariates on response variable through tensor and obtain  $\mathbf{X} - \alpha \overline{\times}_{M+1} \mathbf{Z}$  as adjusted tensor to fit tensor discriminant analysis model.

In estimating  $\alpha$ , which is the alpha in the package, adjten first centers both tensor and covariates within their individual classes, then performs tensor response regression which regresses X on Z.

#### Value

gamma	The estimated coefficients of covariates to plug in classifier. gamma is the $\gamma_k$ defined function catch of dimension $q \times (K-1)$ , where q is the size of covariates and K is the number of classes.
xres	Adjusted training tensor list $\mathbf{X} - \alpha \overline{\times}_{M+1} \mathbf{Z}$ after adjusting for covariates. The effect of the covariate is removed.
testxres	Adjusted testing tensor list $\mathbf{X} - \boldsymbol{\alpha} \overline{\times}_{M+1} \mathbf{Z}$ after adjusting for covariates. The effect of the covariate is removed.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Pan, Y., Mai, Q., and Zhang, X. (2018), "Covariate-Adjusted Tensor Classification in High-Dimensions." Journal of the American Statistical Association, *accepted*.

### See Also

catch

```
n <- 20
p <- 4
k <- 2
nvars <- p*p*p
x <- array(list(),n)
vec_x <- matrix(rnorm(n*nvars),nrow=n,ncol=nvars)
vec_x[1:10,] <- vec_x[1:10,]+2
z <- matrix(rnorm(n*2),nrow=n,ncol=2)
z[1:10,] <- z[1:10,]+0.5
y <- c(rep(1,10),rep(2,10))
for (i in 1:n){
   x[[i]] <- array(vec_x[i,],dim=c(p,p,p))
}
obj <- adjten(x, z, y)</pre>
```

4 adjvec

adjvec

Adjust vector for covariates.

#### **Description**

Adjusts vector with respect to covariates. Vector depends on the covariates through a linear regression model. The function returns the coefficients of covariates in regression and adjusted predictor matrix for further classifier modeling. It estimates coefficients based on training data, and then adjusts training tensor. When testing data is provided, the function will automatically adjust testing data by learned coefficients as well.

#### Usage

```
adjvec(x, z, y, testx = NULL, testz = NULL, is.centered = FALSE)
```

### **Arguments**

х	Input matrix of dimension $N \times p$ , where $N$ is the number of observations and $p$ is the number of variables. Each row is an observation
Z	Input covariate matrix of dimension $N \times q$ , where $q < N$ . Each row of z is an observation.
У	Class label vector of dimention $N \times 1$ . For K class problems, y takes values in $\{1,\cdots,{\sf K}\}.$
testx	Input testing matrix. Each row is a test case. When testx is not provided, the function will only adjust training data.
testz	Input testing covariate matrix with each row being an observation.
is.centered	Indicates whether the input vector and covariates have already been centered by their within class mean or not. If is.centered is FALSE, the function adjvec will center data by class. If is.centered is TRUE, the function will skip the centering step.

#### **Details**

Similar as CATCH model, assume the linear relationship between vector predictors and covariates as

$$\mathbf{X} = \boldsymbol{\mu}_k + \boldsymbol{\alpha} \times \mathbf{Z} + \mathbf{E},$$

where  $\mathbf{X}$  is a  $N \times p$  matrix and  $\boldsymbol{\alpha}$  is the matrix of estimated coefficient of covariates. The function removes the effects of covariates on response variable through vector and obtain  $\mathbf{X} - \boldsymbol{\alpha} \times \mathbf{Z}$  as adjusted predictors to fit MSDA and DSDA model.

#### Value

gamma

The estimated coefficients of covariates to plug in classifier. gamma is similar as the  $\gamma_k$  defined function catch of dimension  $q \times (K-1)$ , where q is the size of covariates and K is the number of classes.

xres Adjusted training predictor matrix  $\mathbf{X} - \boldsymbol{\alpha} \times \mathbf{Z}$  after adjusting for covariates. The

effect of the covariate is removed.

testxres Adjusted testing predictor matrix  $\mathbf{X} - \boldsymbol{\alpha} \times \mathbf{Z}$  after adjusting for covariates. The

effect of the covariate is removed.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Pan, Y., Mai, Q., and Zhang, X. (2018), "Covariate-Adjusted Tensor Classification in High-Dimensions." Journal of the American Statistical Association, *accepted*.

#### See Also

adjten

#### **Examples**

```
n <- 50
p <- 200
k <- 2
q <- 2
x <- matrix(rnorm(n*p), n, p)
z <- matrix(rnorm(n*q), n, q)
x[1:20, ] <- x[1:20, ] + 2
z[1:20, ] <- z[1:20, ] + 0.5
y <- c(rep(1, 20), rep(2, 30))
obj <- adjvec(x, z, y)</pre>
```

catch

Fit a CATCH model and predict categorical response.

### Description

The catch function solves classification problems and selects variables by fitting a covariate-adjusted tensor classification in high-dimensions (CATCH) model. The input training predictors include two parts: tensor data and low dimensional covariates. The tensor data could be matrix as a special case of tensor. In catch, tensor data should be stored in a list form. If the dataset contains no covariate, catch can also fit a classifier only based on the tensor predictors. If covariates are provided, the method will adjust tensor for covariates and then fit a classifier based on the adjusted tensor along with the covariates. If users specify testing data at the same time, predicted response will be obtained as well.

#### Usage

```
catch(x, z = NULL, y, testx = NULL, testz = NULL, nlambda = 100,
lambda.factor = ifelse((nobs - nclass) <= nvars, 0.2, 1E-03),
lambda = NULL,dfmax = nobs, pmax = min(dfmax * 2 + 20, nvars),
pf = rep(1, nvars), eps = 1e-04, maxit = 1e+05, sml = 1e-06,
verbose = FALSE, perturb = NULL)</pre>
```

#### **Arguments**

lambda

dfmax

pmax

pf

eps

х	Input tensor (or matrix) list of length $N$ , where $N$ is the number of observations. Each element of the list is a tensor or matrix. The order of tensor can be any positive integer not less than 2.
Z	Input covariate matrix of dimension $N \times q$ , where $q < N$ . z can be omitted if covariate is absent.
у	Class label. For K class problems, y takes values in $\{1, \dots, K\}$ .
testx	Input testing tensor or matrix list. Each element of the list is a test case. When testx is not provided, the function will only fit the model and return the classifier. When testx is provided, the function will predict response on testx as well.
testz	Input testing covariate matrix. Can be omitted if covariate is absent. However, training covariates z and testing covariates testz must be provided or not at the same time.
nlambda	The number of tuning values in sequence lambda. If users do not specify lambda values, the package will generate a solution path containing nlambda many tuning values of lambda. Default is 100.
lambda.factor	When lambda is not supplied, catch first finds the largest value in lambda which yields $\beta=0$ . Then the minimum value in lambda is obtained by (largest

value\*lambda.factor). The sequence of lambda is generated by evenly sampling nlambda numbers within the range. Default value of lambda.factor is 0.2 if N < p and 0.0001 if N > p.

A sequence of user-specified lambda values. lambda is the weight of L1 penalty and a smaller lambda allows more variables to be nonzero. If NULL, then the algorithm will generate a sequence of nlambda many potential lambdas according to lambda.factor.

The maximum number of selected variables in the model. Default is the number of observations N.

The maximum number of potential selected variables during iteration. In middle step, the algorithm can select at most pmax variables and then shrink part of them such that the nubmer of final selected variables is less than dfmax. Default is

 $\min(df max \times 2 + 20, N).$ 

Weight of lasso penalty. Default is a vector of value 1 and length p, representing

L1 penalty of length p. Can be mofidied to use adaptive lasso penalty.

Convergence threshold for coordinate descent difference between iterations. De-

fault value is 1e-04.

maxit Maximum iteration times for all lambda. Default value is 1e+05.

sml Threshold for ratio of loss function change after each iteration to old loss func-

tion value. Default value is 1e-06.

verbose Indicates whether print out lambda during iteration or not. Default value is

FALSE.

perturb Perturbation scaler. If it is specified, the value will be added to diagonal of

estimated covariance matrix. A small value can be used to accelerate iteration.

Default value is NULL.

#### **Details**

The catch function fits a linear discriminant analysis model as follows:

$$\mathbf{Z}|(Y=k) \sim N(\boldsymbol{\phi_k}, \boldsymbol{\psi}),$$

$$\mathbf{X}|(\mathbf{Z}=\mathbf{z},Y=k) \sim TN(\boldsymbol{\mu}_k + \boldsymbol{\alpha} \bar{\times}_{M+1} \mathbf{z}, \boldsymbol{\Sigma}_1, \cdots, \boldsymbol{\Sigma}_M).$$

The categorical response is predicted from the estimated Bayes rule:

$$\widehat{Y} = \arg \max_{k=1,\dots,K} a_k + \gamma_k^T \mathbf{Z} + \langle \boldsymbol{\beta}_k, \mathbf{X} - \boldsymbol{\alpha} \times \mathbf{X} \rangle,$$

where **X** is the tensor, **Z** is the covariates,  $a_k$ ,  $\gamma_k$  and  $\alpha$  are parameters estimated by CATCH. A detailed explanation can be found in reference. When Z is not NULL, the function will first adjust tensor on covariates by modeling

$$\mathbf{X} = \boldsymbol{\mu}_k + \boldsymbol{\alpha} \overline{\times}_{M+1} \mathbf{Z} + \mathbf{E},$$

where **E** is an unobservable tensor normal error independent of **Z**. Then catch fits model on the adjusted training tensor  $\mathbf{X} - \alpha \overline{\times}_{M+1} \mathbf{Z}$  and makes predictions on testing data by using the adjusted tensor list. If Z is NULL, it reduces to a simple tensor discriminant analysis model.

The coefficient of tensor  $\beta$ , represented by beta in package, is estimated by

$$\min_{\boldsymbol{\beta}_2,...,\boldsymbol{\beta}_K} \left[ \sum_{k=2}^K \left( \langle \boldsymbol{\beta}_k, [\![ \boldsymbol{\beta}_k; \widehat{\boldsymbol{\Sigma}}_1, \ldots, \widehat{\boldsymbol{\Sigma}}_M ]\!] \rangle - 2 \langle \boldsymbol{\beta}_k, \widehat{\boldsymbol{\mu}}_k - \widehat{\boldsymbol{\mu}}_1 \rangle \right) + \lambda \sum_{j_1...j_M} \sqrt{\sum_{k=2}^K \beta_{k,j_1...j_M}^2} \right].$$

When response is multi-class, the group lasso penalty over categories is added to objective function through parameter lambda, and it reduces to a lasso penalty in binary problems.

The function catch will predict categorical response when testing data is provided. If testing data is not provided or if one wishes to perform prediction separately, catch can be used to only fit model with a catch object outcome. The object outcome can be combined with the adjusted tensor list from adjten to perform prediction by predict.catch.

#### Value

beta	Output variable coefficients for each lambda, which is the estimation of $oldsymbol{eta}$ in the
	Bayes rule. beta is a list of length being the number of lambdas. Each element
	of beta is a matrix of dimension $nvars \times (nclass - 1)$ .

df The number of nonzero variables for each value in sequence lambda.

dim Dimension of coefficient array.

lambda	The actual lambda sequence used. The user specified sequence or automatically generated sequence could be truncated by constraints on dfmax and pmax.
obj	Objective function value for each value in sequence lambda.
X	The tensor list after adjustment in training data. If covariate is absent, this is the original input tensor list.
У	Class label in training data.
npasses	Total number of iterations.
jerr	Error flag.
sigma	Estimated covariance matrix on each mode. sigma is a list with the ith element being covariance matrix on ith mode.
delta	Estimated delta matrix $(vec(\widehat{\boldsymbol{\mu}}_2 - \widehat{\boldsymbol{\mu}}_1), \cdots, vec(\widehat{\boldsymbol{\mu}}_K - \widehat{\boldsymbol{\mu}}_1))$ .
mu	Estimated mean array of $X$ .
prior	Proportion of samples in each class.
call	The call that produces this object.
pred	Predicted categorical response for each value in sequence lambda when testx is provided.

### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Pan, Y., Mai, Q., and Zhang, X. (2018), "Covariate-Adjusted Tensor Classification in High-Dimensions." Journal of the American Statistical Association, *accepted*.

### See Also

```
cv.catch, predict.catch, adjten
```

```
#without prediction
n <- 20
p <- 4
k <- 2
nvars <- p*p*p
x <- array(list(),n)
vec_x <- matrix(rnorm(n*nvars), nrow=n, ncol=nvars)
vec_x[1:10,] <- vec_x[1:10,]+2
z <- matrix(rnorm(n*2), nrow=n, ncol=2)
z[1:10,] <- z[1:10,]+0.5
y <- c(rep(1,10),rep(2,10))
for (i in 1:n){
    x[[i]] <- array(vec_x[i,],dim=c(p,p,p))
}
obj <- catch(x,z,y=y)</pre>
```

catch\_matrix 9

catch_matrix Fit a CATCH model for matrix and predict categorical response.
---

### **Description**

Fits a classifier for matrix data. catch\_matrix is a special case of catch when each observation  $X_i$  is a matrix. Different from catch takes list as input, data need to be formed in an array to call the function (see arguments). The function will perform prediction as well.

### Usage

```
catch_matrix(x, z = NULL, y, testx = NULL, testz = NULL, ...)
```

### **Arguments**

Х	Input matrix array. The array should be organized with dimension $p_1 \times p_2 \times N$ .
Z	Input covariate matrix of dimension $N \times q$ , where $q < N$ . z can be omitted if covariate is absent.
у	Class label. For K class problems, y takes values in $\{1, \dots, K\}$ .
testx	Input testing matrix array. When testx is not provided, the function will only fit model. When testx is provided, the function will predict response on testx as well.
testz	Input testing covariate matrix. Can be omitted if there is no covariate.
	Other arguments that can be passed to catch.

#### **Details**

The function fits a matrix classifier as a special case of catch. The fitted model and predictions should be identical to catch when matrix data is provided. Input matrix should be organized as three-way array where sample size is the last dimension. If the matrix is organized in a list, users can either reorganize it or use catch directly to fit model, which takes a matrix or tensor list as input and has the same output as catch\_matrix.

#### Value

beta	Output variable coefficients for each lambda. beta is a list of length being the number of lambdas. Each element of beta is a matrix of dimension $(p_1 \times p_2) \times (nclass - 1)$ .
df	The number of nonzero variables for each value in sequence lambda.
dim	Dimension of coefficient array.
lambda	The actual lambda sequence used. The user specified sequence or automatically generated sequence could be truncated by constraints on dfmax and pmax.
obj	Objective function value for each value in sequence lambda.
X	The matrix list after adjustment in training data. If covariate is absent, this is the original input matrix.

10 catch\_matrix

у	Class label in training data.
npasses	Total number of iterations.
jerr	Error flag.
sigma	Estimated covariance matrix on each mode. sigma is a list with the ith element being covariance matrix on ith mode.
delta	Estimated delta matrix $(vec(\widehat{\boldsymbol{\mu}}_2-\widehat{\boldsymbol{\mu}}_1),\cdots,vec(\widehat{\boldsymbol{\mu}}_K-\widehat{\boldsymbol{\mu}}_1)).$
mu	Estimated mean array.
prior	Prior proportion of observations in each class.
call	The call that produces this object.
pred	Predicted categorical response for each value in sequence lambda when testx is provided.

### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

### References

Pan, Y., Mai, Q., and Zhang, X. (2018), "Covariate-Adjusted Tensor Classification in High-Dimensions." Journal of the American Statistical Association, *accepted*.

### See Also

catch

```
#without prediction n <-20 p <-4 k <-2 nvars <-p*p x=array(rnorm(n*nvars),dim=c(p,p,n)) x[,,11:20]=x[,,11:20]+0.3 z <-matrix(rnorm(n*2), nrow=n, ncol=2) z[1:10,] <-z[1:10,]+0.5 y <-c(rep(1,10),rep(2,10)) obj <- catch_matrix(x,z,y=y)
```

csa 11

csa

Colorimetric sensor array (CSA) data

#### Description

A dataset collected from a series of CSA experiments to identify volatile chemical toxicants (VCT). Chemical dyes were exposed to VCT under different concentration conditions and colors of dyes were recorded to identify the class of VCT. There are two concentration conditions PEL (permissible exposure level) and IDLH (immediately dangerous to life of health).

#### Usage

data(csa)

#### **Format**

Two lists, *PEL* and *IDLH*, and a numeric vector y. Each list contains 147 matrices of dimension  $36 \times 3$ .

PEL A list of matrices containing the observations after exposure at PEL.

IDLH A list of matrices containing the observations after exposure at IDLH level.

y Class label ranging from 1 to 21.

#### **Details**

This dataset is provided in the Supplementary matrial of Zhong (2015). In each concentration case, there are 147 observations and 21 classes. We reorganize the data into a list to be directly called by catch. For matrices in the list, each row represents a dye and the three columns correspond to red, green and blue.

### Source

Wenxuan Zhong and Kenneth S. Suslick (2015). "Matrix discriminant analysis with application to colorimetric sensor array data" *Technometrics* **57**(4), 524–534.

cv.catch

Cross-validation for CATCH

### Description

Performs k-fold cross validation for CATCH and returns the best tuning parameter  $\lambda$  in the user-specified or automatically generated choices.

#### Usage

```
cv.catch(x, z = NULL, y, nfolds = 5, lambda = NULL,
lambda.opt = "min",...)
```

12 cv.catch

#### **Arguments**

)	X	Input tensor or matrix list of length $N$ , where $N$ is the number of observations. Each element of the list is a tensor or matrix. The order of tensor can be any number and not limited to three.
2	Z	Input covariate matrix of dimension $N \times q$ , where $q < N$ . z can be omitted if covariate is absent.
,	y	Class label. For K class problems, y takes values in $\{1, \dots, K\}$ .
1	nfolds	Number of folds. Default value is 5.
	lambda	User-specified lambda sequence for cross validation. If not specified, the algorithm will generate a sequence of lambdas based on all data and cross validate on the sequence.
-	lambda.opt	The optimal criteria when multiple elements in lambda return the same minimum classification error. "min" will return the smallest lambda with minimum cross validation error. "max" will return the largest lambda with the minimum cross validation error.
		Other arguments that can be passed to catch.

#### **Details**

The function cv.catch runs function catch nfolds+1 times. The first one fits model on all data. If lambda is specified, it will check if all lambda satisfies the constraints of dfmax and pmax in catch. If not, a lambda sequence will be generated according to lambda.factor in catch. Then the rest nfolds many replicates will fit model on nfolds-1 many folds data and predict on the omitted fold, repectively. Return the lambda with minimum average cross validation error and the largest lambda within one standard error of the minimum.

#### Value

lambda	The actual lambda sequence used. The user specified sequence or automatically generated sequence could be truncated by constraints on dfmax and pmax.
CVM	The mean of cross validation errors for each lambda.
cvsd	The standard error of cross validaiton errors for each lambda.
lambda.min	The lambda with minimum cross validation error. If lambda.opt is min, then returns the smallest lambda with minimum cross validation error. If lambda.opt is max, then returns the largest lambda with minimum cross validation error.
lambda.1se	The largest lambda with cross validation error within one standard error of the minimum.
catch.fit	The fitted catchobj object.

### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Pan, Y., Mai, Q., and Zhang, X. (2018), "Covariate-Adjusted Tensor Classification in High-Dimensions." Journal of the American Statistical Association, *accepted*.

cv.dsda 13

### See Also

catch

#### **Examples**

```
n <- 20
p <- 4
k <- 2
nvars <- p*p*p
x <- array(list(),n)
vec_x <- matrix(rnorm(n*nvars), nrow=n, ncol=nvars)
vec_x[1:10,] <- vec_x[1:10,]+2
z <- matrix(rnorm(n*2),nrow=n,ncol=2)
z[1:10,] <- z[1:10,]+0.5
y <- c(rep(1,10),rep(2,10))
for (i in 1:n){
    x[[i]] <- array(vec_x[i,], dim=c(p,p,p))
}
object <- cv.catch(x, z, y=y)</pre>
```

cv.dsda

Cross validation for direct sparse discriminant analysis

### **Description**

Choose the optimal lambda for direct sparse discriminant analysis by cross validation.

#### Usage

```
cv.dsda(x, y, nfolds = 5, lambda=lambda, lambda.opt="min",
    standardize=FALSE, alpha=1, eps=1e-7)
```

#### **Arguments**

x	An n by p matrix containing the predictors.
У	An n-dimensional vector containing the class labels.
nfolds	The number of folds to be used in cross validation. Default is 5.
lambda	A sequence of lambda's.
lambda.opt	Should be either "min" or "max", specifying whether the smallest or the largest lambda with the smallest cross validation error should be used for the final classification rule.
standardize	A logic object indicating whether x.matrix should be standardized before performing DSDA. Default is FALSE.
alpha	The elasticnet mixing parameter, the same as in glmnet. Default is alpha=1 so that the lasso penalty is used.
eps	Convergence threshold for coordinate descent, the same as in glmnet. Default is 1e-7.

14 cv.msda

#### Value

lambda The sequence of lambda's used in cross validation.

cvm Cross validation errors.

cvsd The standard error of the cross validation errors.

lambda.min The optimal lambda chosen by cross validation.

model.fit The fitted model.

#### References

Mai, Q., Zou, H. and Yuan, M. (2013). A direct approach to sparse discriminant analysis in ultrahigh dimensions. Biometrika, 99, 29-42.

#### See Also

```
cv.dsda predict.dsda dsda
```

cv.msda

Cross-validation for DSDA/MSDA through function msda

#### **Description**

Performs K-fold cross validation for msda and returns the best tuning parameter  $\lambda$  in the user-specified or automatically generated choices.

### Usage

```
cv.msda(x, y, model = NULL, nfolds = 5, lambda = NULL,
  lambda.opt = "min", ...)
```

#### **Arguments**

У

X	Input matrix of predictors. x is of dimension $N \times p$ ; each row is an observation
	vector

Class label. For K class problems, y takes values in  $\{1, \dots, K\}$ .

model Method type. The model argument can be one of 'binary', 'multi.original',

'multi.modified' and the default is NULL. The function supports fitting DSDA and MSDA models by specifying method type. Without specification, the function will automatically choose one of the methods. If the response variable is binary, the function will fit a DSDA model. If the response variable is multiclass, the function will fit an original MSDA model for dimension p <= 2000

and a modified MSDA model for dimension p > 2000.

nfolds Number of folds. Default value is 5. Although nfolds can be as large as the

sample size (leave-one-out CV), it is not recommended for large datasets. Smallest value allowable is nfolds=3 for multi.original and multi.modified.

cv.msda 15

lambda User-specified lambda sequence for cross validation. If not specified, the algo-

rithm will generate a sequence of lambdas based on all data and cross validate

on the sequence.

lambda.opt The optimal criteria when multiple elements in lambda return the same mini-

mum classification error. "min" will return the smallest lambda with minimum cross validation error. "max" will return the largest lambda with the minimum

cross validation error.

... other arguments that can be passed to msda.

#### **Details**

The function cv.msda runs function msda nfolds+1 times. The first one fits model on all data. If lambda is specified, it will check if all lambda satisfies the constraints of dfmax and pmax in msda. If not, a lambda sequence will be generated according to lambda factor in msda. Then the rest nfolds many replicates will fit model on nfolds-1 many folds data and predict on the omitted fold, repectively. Return the lambda with minimum average cross validation error and the largest lambda within one standard error of the minimum.

Similar as msda, user can specify which method to use by inputing argument model. Without specification, the function can automatically decide the method by number of classes and variables.

#### Value

An object of class cv.dsda or cv.msda.original or cv.msda.modified is returned, which is a list with the ingredients of the cross-validation fit.

lambda The actual lambda sequence used. The user specified sequence or automatically

generated sequence could be truncated by constraints on dfmax and pmax.

cvm The mean of cross validation errors for each lambda.

cvsd The standard error of cross validation errors for each lambda.

lambda.min The lambda with minimum cross validation error. If lambda.opt is min, then

 $returns\ the\ smallest\ lambda\ with\ minimum\ cross\ validation\ error.\ If\ lambda\ .opt$ 

is max, then returns the largest lambda with minimum cross validation error.

lambda.1se The largest value of lambda such that error is within one standard error of the

minimum. This arguement is only available for object cv.msda.original and

cv.msda.modified.

model.fit A fitted cv.dsda or cv.msda.original or cv.msda.modified object for the

full data.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Mai, Q., Zou, H. and Yuan, M. (2012), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrica, 99, 29-42.

16 cv.SeSDA

Mai, Q., Yang, Y., and Zou, H. (2017), "Multiclass sparse discriminant analysis." Statistica Sinica, in press.

```
URL: https://github.com/emeryyi/msda
```

#### See Also

msda

### **Examples**

```
data(GDS1615)
x <- GDS1615$x
y <- GDS1615$y
obj.cv <- cv.msda(x=x, y=y, nfolds=5, lambda.opt="max")
lambda.min <- obj.cv$lambda.min
obj <- msda(x=x, y=y, lambda=lambda.min)
pred <- predict(obj,x)</pre>
```

cv.SeSDA

Cross validation for semiparametric sparse discriminant analysis

### Description

Choose the optimal lambda for semiparametric sparse discriminant analysis by cross validation.

### Usage

```
cv.SeSDA(x, y, nfolds = 5, lambda=NULL, lambda.opt="min",
    standardize=FALSE, alpha=1, eps=1e-7)
```

### Arguments

X	An n by p matrix containing the predictors.
У	An n-dimensional vector containing the class labels.
nfolds	The number of folds to be used in cross validation. Default is 5.
lambda	A sequence of lambda's.
lambda.opt	Should be either "min" or "max", specifying whether the smallest or the largest lambda with the smallest cross validation error should be used for the final classification rule.
standardize	A logic object indicating whether x.matrix should be standardized before performing DSDA. Default is FALSE.
alpha	The elasticnet mixing parameter, the same as in glmnet. Default is alpha=1 so that the lasso penalty is used.
eps	Convergence threshold for coordinate descent, the same as in glmnet. Default is 1e-7.

dsda 17

### Value

transform The transformation functions.

objdsda The output of cross validation from cv.dsda on transformed data.

#### References

Mai, Q., Zou, H. and Yuan, M. (2013). A direct approach to sparse discriminant analysis in ultrahigh dimensions. Biometrika, 99, 29-42.

### See Also

cv.dsda SeSDA

dsda

Solution path for direct sparse discriminant analysis

### **Description**

Compute the solution path for direct sparse discriminant analysis (DSDA).

### Usage

```
dsda(x, z=NULL, y, testx=NULL, testz=NULL, standardize=FALSE,
lambda=lambda, alpha=1, eps=1e-7)
```

### Arguments

X	Input matrix of predictors. x is of dimension $N \times p$ ; each row is an observation vector.
Z	Input covariate matrix of dimension $N \times q$ , where $q < N$ . z can be omitted if covariate is absent.
У	An n-dimensional vector containing the class labels. The classes have to be labeled as 1 and 2.
testx	Input testing matrix. Each row is a test case. When testx is not provided, the function will only fit the model and return the classifier. When testx is provided, the function will predict response on testx as well.
testz	Input testing covariate matrix. Can be omitted if covariate is absent. However, training covariates z and testing covariates testz must be provided or not at the same time.
standardize	A logic object indicating whether x should be standardized before performing DSDA. Default is FALSE.
lambda	A sequence of lambda's. If lambda is missed, the function will automatically generates a sequence of lambda's to fit model.
alpha	The elasticnet mixing parameter, the same as in glmnet. Default is alpha=1 so that the lasso penalty is used.
eps	Convergence threshold for coordinate descent, the same as in glmnet. Default is 1e-7.

18 dsda.all

### Value

beta	Output variable coefficients for each lambda. The first element of each solution is the intercept.
lambda	The sequence of lambda's used in computing the solution path.
Х	The predictor matrix in training data.
у	The class label in training data.
pred	Predicted categorical response for each value in sequence lambda when testx is provided.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Mai, Q., Zou, H. and Yuan, M. (2013). A direct approach to sparse discriminant analysis in ultrahigh dimensions. Biometrika, 99, 29-42.

### **Examples**

```
data(GDS1615) ##load the prostate data
x<-GDS1615$x
y<-GDS1615$y
x=x[which(y<3),]
y=y[which(y<3)]
obj.path <- dsda(x, y=y)</pre>
```

dsda.all

Direct sparse discriminant analysis

### **Description**

Performs direct sparse discriminant analysis, with the optimal lambda chosen by cross validation. The function can perform prediction on test data as well.

### Usage

```
dsda.all(x, y, x.test.matrix=NULL, y.test=NULL, standardize=FALSE, lambda.opt="min", nfolds=10, lambda=lambda, alpha=1, eps=1e-7)
```

dsda.all

#### **Arguments**

x	An n by p matrix containing the predictors.
у	An n-dimensional vector containing the class labels 1 and 2.

x.test.matrix The predictors of a testing set. (Optional.)

y.test The class labels of the testing set. (Required if x.test.matrix is supplied, but

otherwise optional.)

standardize A logic object indicating whether x.matrix should be standardized before per-

forming DSDA. Default is FALSE.

lambda.opt Should be either "min" or "max", specifying whether the smallest or the largest

lambda with the smallest cross validation error should be used for the final clas-

sification rule.

nfolds The number of folds to be used in cross validation. Default is 10.

lambda A sequence of lambda's.

alpha The elasticnet mixing parameter, the same as in glmnet. Default is alpha=1 so

that the lasso penalty is used.

eps Convergence threshold for coordinate descent, the same as in glmnet. Default is

1e-7.

#### Value

error Testing error if x.test.matrix is supplied.

beta The coefficients of the classification rule corresponding to the optimal lambda

chosen by cross validation.

s The optimal lambda chosen by cross validation.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Mai, Q., Zou, H. and Yuan, M., (2012), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrika, 99, 29-42.

### See Also

dsda

```
data(GDS1615) ##load the prostate data
x<-GDS1615$x
y<-GDS1615$y

x=x[which(y<3),]
y=y[which(y<3)]</pre>
```

20 GDS1615

```
n<-length(y)  ##split the original dataset to a training set and a testing set
n.test<-round(n/3)
set.seed(20120822)
id<-sample(n,n.test,replace=FALSE)
x.train<-x[-id,]
x.test<-x[id,]
y.train<-y[-id]
y.test<-y[id]

set.seed(123)
##perform direct sparse discriminant analysis
obj<-dsda.all(x.train,y.train,x.test,y.test)
obj$error</pre>
```

GDS1615

GDS1615 data introduced in Burczynski et al. (2012).

### **Description**

The dataset is a subset of the dataset available on Gene Expression Omnibus with the accession number GDS1615. The original dataset contains 22283 gene expression levels and the disease states of the observed subjects. In Mai, Yang and Zou, the dimension of the original dataset was first reduced to 127 by F-test screening.

#### Usage

```
data(GDS1615)
```

#### Value

This data frame contains the following:

- x Gene expression levels.
- y Disease state that is coded as 1,2,3. 1: normal; 2: ulcerative colitis; 3: Crohn's disease.

#### References

M. E. Burczynski, R. L Peterson, N. C. Twine, K. A. Zuberek, B. J. Brodeur, L. Casciotti, V. Maganti, P. S. Reddy, A. Strahs, F. Immermann, W. Spinelli, U. Schwertschlag, A. M. Slager, M. M. Cotreau, and A. J. Dorner. (2012), "Molecular classification of crohn's disease and ulcerative colitis patients using transcriptional profiles in peripheral blood mononuclear cells". *Journal of Molecular Diagnostics*, 8:51–61.

Mai, Q., Zou, H. and Yuan, M. (2012), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrica, 99, 29-42.

```
data(GDS1615)
```

getnorm 21

getnorm Direct sparse discriminant analysis	getnorm	Direct sparse discriminant analysis
---	---------	-------------------------------------

#### **Description**

Transform the predictors to achieve normality.

#### Usage

```
getnorm(x, y, type="pooled")
```

### Arguments

X	an n dimensional	vector containing n o	bservations for one predictor.
---	------------------	-----------------------	--------------------------------

y an n-dimensional vector containing the class labels.

type The type of estimator. Two estimators were proposed in Mai & Zou (2015), the

naive estimator and the pooled estimator. The function getnorm() uses the naive estimator if type="naive", and it uses the pooled estimator if type="pooled". The default is "pooled". When the naive estimator is used, it is recommended to label

the class with more samples as Class 0.

#### Value

x.norm	Transformed x.
f0	The transformation computed based on observations from Class 0. Not applicable if type="naive".
f1	The transformation computed based on observations from Class 1. Not applicable if type="naive".
mu.hat	The sample mean for transformed x from Class 1.
transform	The transformation that was actually used to transform x.

### References

Mai, Q., Zou, H. and Yuan, M. (2013). A direct approach to sparse discriminant analysis in ultrahigh dimensions. Biometrika, 99, 29-42.

Mai, Q. and Zou, H. (2015). Sparse semiparametric discriminant analysis. Journal of Multivariate Analysis, 135, 175-188.

```
data(GDS1615) ##load the prostate data
x<-GDS1615$x
y<-GDS1615$y
x<-exp(x[which(y<3),])
y<-y[which(y<3)]</pre>
```

```
n<-length(y)
n1 < -sum(y==1)
n2<-n-n1
n1.test < -round(n1/2)
n2.test < -round(n2/2)
n.test < -n1.test + n2.test
n.train<-n-n.test
id.test<-c(sample(which(y==1),n1.test),sample(which(y==2),n2.test))</pre>
p < -ncol(x)
x.train<-x[-id.test,]
y.train<-y[-id.test]
x.test<-x[id.test,]</pre>
y.test<-y[id.test]
 x.norm<-matrix(0,n.train,p)</pre>
  x.test.norm<-matrix(0,n.test,p)</pre>
  for(i in 1:p){
    obj.norm<-getnorm(x.train[,i],y.train)
    x.norm[,i]<-obj.norm$x.norm</pre>
    x.test.norm[,i]<-obj.norm$transform(x.test[,i])</pre>
  }
  obj<-dsda.all(x.norm,y.train,x.test.norm,y.test)</pre>
```

msda

Fits a regularization path of Sparse Discriminant Analysis and predicts

#### **Description**

Fits a regularization path of Sparse Discriminant Analysis at a sequence of regularization parameters lambda. Performs prediction when testing data is provided. The msda function solves classification problem by fitting a sparse discriminant analysis model. When covariates are provided, the function will first make adjustment on the training data. It provides three models: binary for fitting DSDA model to solve binary classification problems, multi.original and multi.modified for fitting MSDA model to solve multi-class classification problems. multi.original runs faster for small dimension case but the computation ability is limited to a relatively large dimension. multi.modified has no such limitation and works in ultra-high dimensions. User can specify method by argument or use the default settings.

#### Usage

```
msda(x, z=NULL, y, testx=NULL, testz=NULL, model = NULL, lambda = NULL,
  standardize=FALSE, alpha=1, nlambda = 100,
```

```
lambda.factor = ifelse((nobs - nclass)<= nvars, 0.2, 1e-03), dfmax = nobs,
pmax = min(dfmax * 2 + 20, nvars), pf = rep(1, nvars), eps = 1e-04,
maxit = 1e+06, sml = 1e-06, verbose = FALSE, perturb = NULL)</pre>
```

#### **Arguments**

У

X	Input matrix of predictors. x is of dimension $N \times p$ ; each row is an observation
	vector

z Input covariate matrix of dimension  $N \times q$ , where q < N. z can be omitted if covariate is absent.

Class labl. This argument should be a factor for classification. For model='binary', y should be a binary variable with values 1 and 2. For model='multi.original' or 'multi.modified', y should be a multi-class variable starting from 1.

Input testing matrix. Each row is a test case. When testx is not provided, the function will only fit the model and return the classifier. When testx is provided, the function will predict response on testx as well.

Input testing covariate matrix. Can be omitted if covariate is absent. However, training covariates z and testing covariates testz must be provided or not at the same time.

Method type. The model argument can be one of 'binary', 'multi.original', 'multi.modified' and the default is NULL. The function supports fitting DSDA and MSDA models by specifying method type. Without specification, the function will automatically choose one of the methods. If the response variable is binary, the function will fit a DSDA model. If the response variable is multiclass, the function will fit an original MSDA model for dimension p <= 2000 and a modified MSDA model for dimension p > 2000.

A user supplied lambda sequence. Typically, by leaving this option unspecified users can have the program compute its own lambda sequence based on nlambda and lambda.factor. Supplying a value of lambda overrides this. It is better to supply a decreasing sequence of lambda values than a single (small) value, if not, the program will sort user-defined lambda sequence in decreasing order automatically.

A logic object indicating whether x should be standardized before performing DSDA. Default is FALSE. This argument is only valid for model = 'binary'.

The elasticnet mixing parameter, the same as in glmnet. Default is alpha=1 so that the lasso penalty is used in DSDA. This argument is only valid for model = 'binary'.

The number of tuning values in sequence lambda. If users do not specify lambda values, the package will generate a solution path containing nlambda many tuning values of lambda. Default is 100 for model = 'multi.original' and 50 for model = 'multi.modified'.

The factor for getting the minimal lambda in lambda sequence, where min(lambda) = lambda. factor \* max(lambda). max(lambda) is the smallest value of lambda for which all coefficients are zero. The default depends on p (the number of predictors) and its relationship with N (the number of rows in the matrix of predictors). For Original MSDA, if N > p, the default is 0.0001, close to zero.

testx

testz

model

lambda

standardize

alpha

nlambda

lambda.factor

If N < p, the default is 0.2. For Modified MSDA, if  $p \le 5000$ , the default is 0.2. If 5000 , the default is 0.4. If <math>p > 30000, the default is 0.5. A very small value of lambda factor will lead to a saturated fit. It takes no effect if there is user-defined lambda sequence. This argument is only valid for multi.original and multi.modified.

dfmax

The maximum number of selected variables in the model. Default is the number of observations N. This argument is only valid for multi.original and multi.modified.

pmax

The maximum number of potential selected variables during iteration. In middle step, the algorithm can select at most pmax variables and then shrink part of them such that the nubmer of final selected variables is less than dfmax. Default is  $\min(dfmax \times 2 + 20, N)$ .

pf

L1 penalty factor of length p. Separate L1 penalty weights can be applied to each coefficient of  $\theta$  to allow differential L1 shrinkage. Can be 0 for some variables, which implies no L1 shrinkage, and results in that variable always being included in the model. Default is 1 for all variables (and implicitly infinity for variables listed in exclude). This argument is only valid for multi.original and multi.modified.

eps

Convergence threshold for coordinate descent. Each inner coordinate descent loop continues until the relative change in any coefficient. Defaults value is 1e-4.

maxit

Maximum number of outer-loop iterations allowed at fixed lambda value. Default is 1e6. If models do not converge, consider increasing maxit. This argument is only valid for multi.original and multi.modified.

sml

Threshold for ratio of loss function change after each iteration to old loss function value. Default is 1e-06. This argument is only valid for multi.original and multi.modified.

verbose

Whether to print out computation progress. The default is FALSE. This argument is only valid for multi.original and multi.modified.

perturb

A scalar number. If it is specified, the number will be added to each diagonal element of the covariance matrix as perturbation. The default is NULL. This argument is only valid for multi.original and multi.modified.

#### **Details**

The msda function fits a linear discriminant analysis model for vector X as follows:

$$\mathbf{X}|Y = k \sim N(\boldsymbol{\mu}_k, \boldsymbol{\Sigma}).$$

The categorical response is predicted from the Bayes rule:

$$\widehat{Y} = \arg\max_{k=1,\dots,K} (\mathbf{X} - \frac{\boldsymbol{\mu}_k}{2})^T \boldsymbol{\beta}_k + \log \pi_k.$$

The parameter model specifies which method to use in estimating  $\beta$ . Users can use binary for binary problems and binary and multi.modified for multi-class problems. In multi.original, the algorithm first computes and stores  $\Sigma$ , while it doesn't compute or store the entire covariance matrix in multi.modified. Since the algorithm is element-wise based, multi.modified computes

each element of covariance matrix when needed. Therefore, multi.original is faster for low dimension but multi.modified can fit model for a much higher dimension case.

Note that for computing speed reason, if models are not converging or running slow, consider increasing eps and sml, or decreasing nlambda, or increasing lambda.factor before increasing maxit. Users can also reduce dfmax to limit the maximum number of variables in the model.

The arguments list out all parameters in the three models, but not all of them are necessary in applying one of the methods. See the specific explaination of each argument for more detail. Meanwhile, the output of DSDA model only includes beta and lambda.

#### Value

An object with S3 class dsda or msda.original and msda.modified.

beta	Output variable coefficients for each lambda, which is the estimation of $\beta$ in the Bayes rule. beta is a list of length being the number of lambdas. Each element of beta is a matrix of dimension $nvars \times (nclass-1)$ . For model = 'dsda', beta is a vector of length $nvars+1$ , where the first element is intercept.
df	The number of nonzero coefficients for each value of lambda.
obj	The fitted value of the objective function for each value of lambda.
dim	Dimension of each coefficient matrix.
lambda	The actual lambda sequence used. The user specified sequence or automatically generated sequence could be truncated by constraints on dfmax and pmax.
х	The input matrix of predictors for training.
у	Class label in training data.
npasses	Total number of iterations (the most inner loop) summed over all lambda values
jerr	Error flag, for warnings and errors, 0 if no error.
sigma	Estimated sigma matrix. This argument is only available in object msda.original.
delta	Estimated delta matrix. delta[k] = mu[k]-mu[1].
mu	Estimated mu vector.
prior	Prior probability that y belong to class k, estimated by mean(y that belong to k).
call	The call that produced this object
pred	Predicted categorical response for each value in sequence lambda when testx is provided.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Mai, Q., Zou, H. and Yuan, M. (2012), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrica, 99, 29-42.

Mai, Q., Yang, Y., and Zou, H. (2017), "Multiclass sparse discriminant analysis." Statistica Sinica, in press.

URL: https://github.com/emeryyi/msda

26 predict.catch

#### See Also

```
cv.msda, predict.msda
```

### **Examples**

```
data(GDS1615)
x<-GDS1615$x
y<-GDS1615$y
obj <- msda(x = x, y = y)</pre>
```

predict.catch

Predict categorical responses for matrix/tensor data.

### **Description**

Predict categorical responses on new matrix/tensor data given the fitted CATCH model input.

#### Usage

```
## S3 method for class 'catch'
predict(object, newx, z = NULL, ztest = NULL, gamma = NULL,...)
```

### Arguments

object	Input catchobj class object as fitted model.
newx	Input adjusted testing tensor or matrix list. Each element of the list is a tensor. The tensor should of the same dimension as training data.
Z	Input training covariates matrix. z can be omitted if there is no covariate.
ztest	Input testing covariates matrix. ztest can be omitted if there is no covariate.
gamma	Coefficients of covariates obtained from adjten. gamma is NULL if there is no covariate.
	Other arguments that can be passed to predict.

#### Details

The function fits LDA model on selected discriminant vectors. Call predict or predict. catch to perform predictions.

There are two ways to make predictions. One way is to directly predict at the same time as fitting model by catch since predict.catch has already been embedded in catch and it will predicts response when testing data is provided. The other way is to first use adjten to adjuste tensor and catch to fit model. predict.catch will take the input adjusted tensor list newx, covariate coefficient gamma from adjten and the fitted model from catch to perform prediction. The prediction is identical to providing catch testing data.

predict.catch 27

### Value

Predicted response of news for each lambda in model object.

#### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Pan, Y., Mai, Q., and Zhang, X. (2018) Covariate-Adjusted Tensor Classification in High-Dimensions, arXiv:1805.04421.

#### See Also

```
catch, adjten
```

```
#generate training data
n <- 20
p <- 4
k <- 2
nvars <- p*p*p
x <- array(list(),n)</pre>
vec_x <- matrix(rnorm(n*nvars),nrow=n,ncol=nvars)</pre>
vec_x[1:10,] \leftarrow vec_x[1:10,]+2
z <- matrix(rnorm(n*2),nrow=n,ncol=2)</pre>
z[1:10,] \leftarrow z[1:10,]+0.5
y <- c(rep(1,10), rep(2,10))
for (i in 1:n){
  x[[i]] \leftarrow array(vec_x[i,],dim=c(p,p,p))
#generate testing data
newx <- array(list(),n)</pre>
vec_newx <- matrix(rnorm(n*nvars),nrow=n,ncol=nvars)</pre>
vec_newx[1:10,] <- vec_newx[1:10,]+2</pre>
newz <- matrix(rnorm(n*2),nrow=n,ncol=2)</pre>
newz[1:10,] \leftarrow newz[1:10,]+0.5
for (i in 1:n){
  newx[[i]] <- array(vec_newx[i,],dim=c(p,p,p))</pre>
}
#Make adjustment and fit model
obj <- adjten(x, z, y, newx, newz)
fit <- catch(x, z, y)
#Predict
pred <- predict(fit, obj$testxres, z, newz, obj$gamma)</pre>
#The adjusting, fitting model and predicting step can also be completed
#by one command.
pred <- catch(x, z, y, newx, newz)$pred</pre>
```

28 predict.dsda

predict.dsda Prediction for direct sparse discriminant analysis	
---	--

### Description

Predict the class labels by direct sparse discriminant analysis.

### Usage

```
## S3 method for class 'dsda'
predict(object, newx, z=NULL, ztest=NULL, gamma=NULL,...)
```

### Arguments

object	An object returned by dsda or msda with binary setting.
newx	An n by p matrix containing the predictors.
z	Input training covariates matrix. z can be omitted if there is no covariate.
ztest	Input testing covariates matrix. ztest can be omitted if there is no covariate.
gamma	Coefficients of covariates obtained from adjvec. gamma is NULL if there is no covariate.
	Other arguments that can be passed to predict.

### Value

pred The the predicted class labels.

#### References

Mai, Q., Zou, H. and Yuan, M. (2013), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrika, 99, 29-42.

### See Also

```
dsda, dsda.all, predict.msda
```

predict.msda 29

predict.msda	Predict categorical responses for vector data.	

### **Description**

Predict categorical responses on new vector data given the fitted DSDA/MSDA model input.

### Usage

```
## S3 method for class 'msda'
predict(object, newx, z = NULL, ztest = NULL, gamma = NULL,...)
```

### Arguments

object	Fitted model object from msda. The model object can be anyone of binary, multi.original and multi.modified.
newx	The matrix of new values for x at which predictions are to be made. If covariates exist, then news should be adjusted matrix.
Z	Input training covariates matrix. z can be omitted if there is no covariate.
ztest	Input testing covariates matrix. ztest can be omitted if there is no covariate.
gamma	Coefficients of covariates obtained from adjvec. gamma is NULL if there is no covariate.
	Other arguments that can be passed to predict.

#### **Details**

The function fits LDA model on selected discriminant vectors. Call predict or predict.msda to perform prediction. When covariates exist, users could first call adjvec to make adjustment and obtain obtain gamma. The fitted model from msda should also takes adjusted vector as input. The newx in predict.msda should be adjusted vector as well.

#### Value

Predicted class label(s) at the entire sequence of the penalty parameter lambda used to create the model.

### Author(s)

30 predict.SeSDA

#### References

Mai, Q., Zou, H. and Yuan, M. (2012), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrica, 99, 29-42.

Mai, Q., Yang, Y., and Zou, H. (2017), "Multiclass sparse discriminant analysis." Statistica Sinica, in press.

Pan, Y., Mai, Q., and Zhang, X. (2018), "Covariate-Adjusted Tensor Classification in High-Dimensions." Journal of the American Statistical Association, *accepted*.

#### See Also

msda

### **Examples**

```
data(GDS1615)
x<-GDS1615$x
y<-GDS1615$y
obj <- msda(x = x, y = y)
pred<-predict(obj,x)</pre>
```

predict.SeSDA

Prediction for semiparametric sparse discriminant analysis

### **Description**

Predict the class labels by semiparametric sparse discriminant analysis.

#### **Usage**

```
## S3 method for class 'SeSDA'
predict(object, x.test,...)
```

### Arguments

object An object returned by SeSDA.x.test An n by p matrix containing the predictors.... Other arguments that can be passed to predict.

### Value

pred The the predicted class labels.

### References

Mai, Q., Zou, H. and Yuan, M. (2013), "A direct approach to sparse discriminant analysis in ultrahigh dimensions." Biometrika, 99, 29-42.

ROAD 31

### See Also

dsda, SeSDA

ROAD	Solution path for regularized optimal affine discriminant	

### Description

Compute the solution path for regularized optimal affine discriminant (ROAD).

### Usage

```
ROAD(x,y,standardize=FALSE,lambda=NULL,eps=1e-7)
```

### **Arguments**

X	Input matrix of predictors. ${\bf x}$ is of dimension $N\times p$ ; each row is an observation vector.
У	An n-dimensional vector containing the class labels. The classes have to be labeled as 1 and 2.
standardize	A logic object indicating whether x should be standardized before performing ROAD. Default is FALSE.
lambda	A sequence of lambda's. If lambda is missed, the function will automatically generates a sequence of lambda's to fit model.
eps	Convergence threshold for coordinate descent, the same as in glmnet. Default is 1e-7.

### **Details**

The function obtains the solution path of ROAD through dsda.

### Value

beta Output variable coefficients for each lambda.

lambda The sequence of lambda's used in computing the solution path.

### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

### References

Mai, Q. and Zou, H. (2013), "A note on the connection and equivalence of three sparse linear discriminant analysis methods." Technometrics, 55, 243-246.

32 SeSDA

### **Examples**

```
data(GDS1615) ##load the prostate data
x<-GDS1615$x
y<-GDS1615$y
x=x[which(y<3),]
y=y[which(y<3)]
obj.path <- ROAD(x, y)</pre>
```

SeSDA

Solution path for semiparametric sparse discriminant analysis

### Description

Compute the solution path for semiparametric sparse discriminant analysis.

### Usage

```
SeSDA(x,y,standardize=FALSE,lambda=NULL,alpha=1,eps=1e-7)
```

### Arguments

X	Input matrix of predictors. x is of dimension $N \times p$ ; each row is an observation vector.
У	An n-dimensional vector containing the class labels. The classes have to be labeled as 1 and 2.
standardize	A logic object indicating whether x should be standardized after transformation but before fitting classifier. Default is FALSE.
lambda	A sequence of lambda's. If lambda is missed or NULL, the function will automatically generates a sequence of lambda's to fit model.
alpha	The elasticnet mixing parameter, the same as in glmnet. Default is alpha=1 so that the lasso penalty is used.
eps	Convergence threshold for coordinate descent, the same as in glmnet. Default is 1e-7.

### Value

transform The tranformation functions.

objdsda A DSDA object fitted on transformed data.

### Author(s)

sim.bi.vector 33

#### References

Mai, Q., Zou, H. and Yuan, M. (2013). A direct approach to sparse discriminant analysis in ultrahigh dimensions. Biometrika, 99, 29-42.

### **Examples**

```
data(GDS1615) ##load the prostate data
x<-GDS1615$x
y<-GDS1615$y
x=x[which(y<3),]
y=y[which(y<3)]
obj.path <- SeSDA(x,y)</pre>
```

sim.bi.vector

Simulate data

### Description

Simulate a binary data set with vector predictor.

### Usage

```
sim.bi.vector(tesize = 100)
```

### Arguments

tesize

Number of observations in testing data.

### **Details**

The function simulates a data set with p = 500. Response are binary.

### Value

X	Simulated vector predictor.
testx	Simulated testing vector predictor.
у	Response corresponding to x.
testy	Response corresponding to testx.

### Author(s)

34 sim.tensor.cov

sim.tensor.cov

Simulate data

### Description

Simulate a data set with tensor predictor and covariates.

### Usage

```
sim.tensor.cov(tesize = 100)
```

### Arguments

tesize

Number of observations in testing data.

#### **Details**

The function simulates a data set with  $10 \times 10 \times 10$  tensor and covariate being a two-dimensional vector. Response are binary.

#### Value

x Simulated tensor predictor.

z Simulated covariate.

testx Simulated testing tensor predictor.

testz Simualted testing covariate.

vec\_x Vectorization of x.

vec\_testx Vectorization of testx.

y Response corresponding to x and z.

testy Response corresponding to testx and testz.

### Author(s)

SOS 35

SOS	Solution path for sparse discriminant analysis	

### Description

Compute the solution path for sparse optimal scoring (SOS).

### Usage

```
SOS(x,y,standardize=FALSE,lambda=NULL,eps=1e-7)
```

### **Arguments**

X	Input matrix of predictors. ${\bf x}$ is of dimension $N\times p;$ each row is an observation vector.
У	An n-dimensional vector containing the class labels. The classes have to be labeled as 1 and 2.
standardize	A logic object indicating whether x should be standardized before performing SOS. Default is FALSE.
lambda	A sequence of lambda's. If lambda is missed, the function will automatically generates a sequence of lambda's to fit model.
eps	Convergence threshold for coordinate descent, the same as in glmnet. Default is 1e-7.

#### **Details**

The function obtains the solution path of sparse optimal scoring model through dsda.

#### Value

beta Output variable coefficients for each lambda.

1ambda The sequence of lambda's used in computing the solution path.

### Author(s)

Yuqing Pan, Qing Mai, Xin Zhang

#### References

Mai, Q. and Zou, H. (2013), "A note on the connection and equivalence of three sparse linear discriminant analysis methods." Technometrics, 55, 243-246.

36 SOS

```
data(GDS1615) ##load the prostate data
x<-GDS1615$x
y<-GDS1615$y
x=x[which(y<3),]
y=y[which(y<3)]
obj.path <- SOS(x, y)</pre>
```

## **Index**

```
* datasets
    csa, 11
    GDS1615, 20
adjten, 2, 3, 5, 7, 8, 26, 27
adjvec, 4, 28, 29
catch, 3, 4, 5, 7, 9, 10, 12, 13, 26, 27
catch_matrix, 9, 9
csa, 11
cv.catch, 8, 11, 12
cv.dsda, 13
cv.msda, 14, 15, 26
cv. SeSDA, 16
dsda, 17, 19, 28, 31, 35
dsda.all, 18, 28
GDS1615, 20
getnorm, 21
msda, 15, 16, 22, 29, 30
predict, 29
predict.catch, 7, 8, 26, 26
predict.dsda, 28
predict.msda, 26, 28, 29, 29
predict.SeSDA, 30
ROAD, 31
SeSDA, 31, 32
sim.bi.vector, 33
sim.tensor.cov, 34
SOS, 35
x (GDS1615), 20
y (GDS1615), 20
```