Package ‘HDclassif’

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Description Discriminant analysis and data clustering methods for high
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different subspaces with low dimensionality proposing a new parametrization of
the Gaussian mixture model which combines the ideas of dimension reduction and
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R topics documented:

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## Description

Discriminant analysis and data clustering methods for high dimensional data, based on the assumption that high-dimensional data live in different subspaces with low dimensionality, proposing a new parametrization of the Gaussian mixture model which combines the ideas of dimension reduction and constraints on the model.

## Details

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This package is used to make efficient supervised and unsupervised classification with high dimensional data. The supervised method uses the `hdda` function to get the data parameters and the `predict` function to realise the class prediction of a dataset. The unsupervised method is implemented in the `hddc` function, and once the parameters are estimated, the `predict` gives the class prediction of other datasets. The method used in the `hddc` is based on the Expectation - Maximisation algorithm.

## Author(s)

Laurent Berge, Charles Bouveyron and Stephane Girard

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## References


Crabs


Crabs

Morphological Measurements on Leptograpsus Crabs.

Description

The Crabs data frame has 200 rows and 6 columns, describing 5 morphological measurements on 50 crabs each of two colour forms and both sexes, of the species Leptograpsus Variegatus collected at Fremantle, W. Australia.

Usage

data(Crabs)

Format

A data frame with 200 observations on the following 6 variables.

class Type of the crabs: the first character represents the species - "B" or "O" for blue or orange-, the second represents the sex -"M" or "F" for male or female-.

FL Frontal lob size (mm).

RW Rear width (mm).

CL Carapace length (mm).

CW Carapace width (mm).

BD Body depth (mm).

Details

This dataset can also be found in the MASS package, the unique difference is the class vector which is easier to use here.

Source


References

**demo_hddc**  
_Demonstration of the clustering process of HDDC._

**Description**

This demonstration uses a PCA on the first two principal axis of the Crabs dataset -that can be found in the package- to show the clustering process of HDDC. At each step of the clustering, the means and directions are shown by, respectively, points and lines. This function should only be used in demo(hddc).

**Usage**

```r
demo_hddc()
```

**Value**

The plots of the clustering process.

**Note**

The algorithm and the initialization are interactively chosen.

**Author(s)**

Laurent Berge, Charles Bouveyron and Stephane Girard

**See Also**

hddc.

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**hdda**  
_High Dimensional Discriminant Analysis_

**Description**

HDDA is a model-based discriminant analysis method assuming each class of the dataset live in a proper Gaussian subspace which is much smaller than the original one, the hdda.learn function calculates the parameters of each subspace in order to predict the class of new observation of this kind.

**Usage**

```r
hdda(data, cls, model = "AkjBkQkDk", graph = FALSE, d_select = "Cattell",
    threshold = 0.2, com_dim = NULL, show = TRUE, scaling = FALSE,
    cv.dim = 1:10, cv.threshold = c(0.001, 0.005, 0.05, 1:9 * 0.1),
    cv.vfold = 10, LOO = FALSE, noise.ctrl = 1e-08, d)
```
Arguments

data A matrix or a data frame of observations, assuming the rows are the observations and the columns the variables. Note that NAs are not allowed.

cls The vector of the class of each observations, its type can be numeric or string.

model A character string vector, or an integer vector indicating the models to be used. The available models are: "AkjBkQkDk" (default), "AkBkQkDk", "ABkQkDk", "AkjBQkDk", "AkBQkDk", "AbkQkDk", "AkBkQkD", "ABkQkD", "AkjBQkD", "AkBQkD", "ABQkD", "AjBQD", "ABQD". It is not case sensitive and integers can be used instead of names, see details for more information. Several models can be used, if it is, only the results of the one which maximizes the BIC criterion is kept. To run all models, use model="ALL".

graph It is for comparison sake only, when several estimations are run at the same time (either when using several models, or when using cross-validation to select the best dimension/threshold). If graph = TRUE, the plot of the results of all estimations is displayed. Default is FALSE.

d_select Either “Cattell” (default), “BIC” or “CV”. See details for more information. This parameter selects which method to use to select the intrinsic dimensions.

threshold A float strictly within 0 and 1. It is the threshold used in the Cattell’s Scree-Test.

com_dim It is used only for common dimensions models. The user can give the common dimension he wants. If used, it must be an integer. Its default is set to NULL.

show Use show = FALSE to settle off the informations that may be printed.

scaling Logical: whether to scale the dataset (mean=0 and standard-error=1 for each variable) or not. By default the data is not scaled.

cv.dim A vector of integers. Only when d_select="CV". Gives the dimensions for which the CV is to be done. Note that if some dimensions are greater than what it is possible to have, those are taken off.

cv.threshold A vector of floats strictly within 0 and 1. Only when d_select="CV". Gives the thresholds for which the CV is to be done.

cv.vfold An integer. Only when d_select="CV". It gives the number of different sub-samples in which the dataset is split. If “cv.vfold” is greater than the number of observations, then the program equalize them.

LOO If TRUE, it returns the results (classes and posterior probabilities) for leave-one-out cross-validation.

noise.ctrl This parameter avoids to have a too low value of the ‘noise’ parameter b. It guarantees that the dimension selection process do not select too many dimensions (which leads to a potential too low value of the noise parameter b). When selecting the intrinsic dimensions using Cattell’s scree-test or BIC, the function doesn’t use the eigenvalues inferior to noise.ctrl, so that the intrinsic dimensions selected can’t be higher or equal to the order of these eigenvalues.

d DEPRECATED. This parameter is kept for retro compatibility. Now please use the parameter d_select.
Details

Some information on the signification of the model names:

Akj are the parameters of the classes subspaces:
  • if Akj: each class has its parameters and there is one parameter for each dimension
  • if Ak: the classes have different parameters but there is only one per class
  • if Aj: all the classes have the same parameters for each dimension (it’s a particular case with a common orientation matrix)
  • if A: all classes have the same one parameter

Bk are the noises of the classes subspaces:
  • If Bk: each class has its proper noise
  • if B: all classes have the same noise

Qk is the orientation matrix of each class:
  • if Qk: all classes have its proper orientation matrix
  • if Q: all classes have the same orientation matrix

Dk is the intrinsic dimension of each class:
  • if Dk: the dimensions are free and proper to each class
  • if D: the dimension is common to all classes

The model “all” will compute all the models, give their BIC and keep the model with the highest BIC value. Instead of writing the model names, they can also be specified using an integer. 1 represents the most general model (“AkjBkQkDk”) while 14 is the most constrained (“ABQD”), the others number/name matching are given below. Note also that several models can be run at once, by using a vector of models (e.g. model = c("AKBKQKD","AKJBQKD","AJBQD") is equivalent to model = c(8,4,13); to run the 6 first models, use model=1:6). If all the models are to be run, model="all" is faster than model=1:14.

<table>
<thead>
<tr>
<th>AkjBkQkDk</th>
<th>1</th>
<th>AkjBkQkD</th>
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<td>AkBkQkDk</td>
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<td>10</td>
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</table>

The parameter d, is used to select the intrinsic dimensions of the subclasses. Here are his definitions:

• “Cattell”: The Cattell’s scree-test is used to gather the intrinsic dimension of each class. If the model is of common dimension (models 7 to 14), the scree-test is done on the covariance matrix of the whole dataset.

• “BIC”: The intrinsic dimensions are selected with the BIC criterion. See Bouveyron et al. (2010) for a discussion of this topic. For common dimension models, the procedure is done on the covariance matrix of the whole dataset.

• “CV”: A V-fold cross-validation (CV) can be done in order to select the best threshold (for all models) or the best common dimensions (models 7 to 14). The V-fold cross-validation is done for each dimension (respectively threshold) in the argument “cv.dim” (resp. “cv.threshold”),
then the dimension (resp. threshold) that gives the best good classification rate is kept. The dataset is split in “cv.vfold” (default is 10) random subsamples, then CV is done for each sample: each of them is used as validation data while the remaining data is used as training data. For sure, if “cv.vfold” equals the number of observations, then this CV is equivalent to a leave-one-out.

Value

hdda returns an ‘hdc’ object; it’s a list containing:

- **model**: The name of the model.
- **k**: The number of classes.
- **d**: The dimensions of each class.
- **a**: The parameters of each class subspace.
- **b**: The noise of each class subspace.
- **mu**: The mean of each variable for each class.
- **prop**: The proportion of each class.
- **ev**: The eigen values of the var/covar matrix.
- **Q**: The orthogonal matrix of orientation of each class.
- **kname**: The name of each class.
- **bic**: The BIC value of the model used.
- **scaling**: The centers and the standard deviation of the original dataset.

Author(s)

Laurent Berge, Charles Bouveyron and Stephane Girard

References


See Also

hddc, predict.hdc, plot.hdc
Examples

# Example 1:
data <- simuldata(1000, 1000, 50, K=5)
X <- data$X
clx <- data$clx
Y <- data$Y
cly <- data$cly
# we get the HDDA parameters:
prms1 <- hdda(X, clx)

c1l <- predict(prms1, Y, cly)
# the class vector of Y estimated with HDDA:
c1l$class

# another model is used:
prms1 <- hdda(X, clx, model=12)
# model=12 is equivalent to model="ABQKD"
c1l <- predict(prms1, Y, cly)

# Example 2:
data(wine)
a <- wine[,1]
z <- wine[,2]
prms2 <- hdda(a, z, model='all', scaling=TRUE, d_select="bic", graph=TRUE)
c12 <- predict(prms2, a, z)

# getting the best dimension
# using a common dimension model
# we do LOO-CV using cv.vfold=nrow(a)
prms3 <- hdda(a, z, model="akjbqkd", d_select="CV", cv.vfold=nrow(a), scaling=TRUE, graph=TRUE)
c13 <- predict(prms3, a, z)

# Example 3:
# Validation with LOO
prms4 <- hdda(a, z, LOO=TRUE, scaling=TRUE)
sum(prms4$class==z) / length(z)

---

hddc  
High Dimensional Data Clustering

Description

HDDC is a model-based clustering method. It is based on the Gaussian Mixture Model and on the idea that the data lives in subspaces with a lower dimension than the dimension of the original space. It uses the Expectation - Maximisation algorithm to estimate the parameters of the model.
Usage

hddc(data, K = 1:10, model = c("AkjBkQkDk"), threshold = 0.2,
criterion = "bic", com_dim = NULL, itermax = 200, eps = 0.001,
algo = "EM", d_select = "Cattell", init = "kmeans", init.vector,
show = TRUE, mini.nb = c(5, 10), scaling = FALSE, min.individuals = 2,
noise.ctrl = 1e-08, mc.cores = 1, nb.rep = 1, keepAllRes = TRUE,
kmeans.control = list(), d_max = 100, d)

Arguments

data A matrix or a data frame of observations, assuming the rows are the observations and the columns the variables. Note that NAs are not allowed.

K A vector of integers specifying the number of clusters for which the BIC and the parameters are to be calculated; the function keeps the parameters which maximises the criterion. Default is 1:10.

model A character string vector, or an integer vector indicating the models to be used. The available models are: "AkjBkQkDk" (default), "AkBkQkDk", "AkjBQkDk", "AkBQkDk", "AkjBkQkD", "AkBkQkD", "AkjBQkD", "AkBQkD", "AkjBQkD", "AkBQkD", "AkjBQD", "AkBQD", "AkjBQD", "AkBQD", "AkjQBD", "AkBQD", "AkjQBD", "AkBQD", "AkjQBD", "AkBQD", "AkjQBD", "AkBQD", "AkjQBD", "AkBQD". It is not case sensitive and integers can be used instead of names, see details for more information. Several models can be used, if it is, only the results of the one which maximizes the BIC criterion is kept. To run all models, use model="ALL".

threshold A float strictly within 0 and 1. It is the threshold used in the Cattell’s Scree-Test.

criterion Either “BIC” or “ICL”. If several models are run, the best model is selected using the criterion defined by criterion.

com_dim It is used only for common dimensions models. The user can give the common dimension he wants. If used, it must be an integer. Its default is set to NULL.

itermax The maximum number of iterations allowed. The default is 200.

esp A positive double. It is the stopping criterion: the algorithm stops when the difference between two successive Log Likelihoods is lower than 'esp'.

algo A character string indicating the algorithm to be used. The available algorithms are the Expectation-Maximisation ("EM"), the Classification E-M ("CEM") and the Stochastic E-M ("SEM"). The default algorithm is the "EM".

show Use show = FALSE to settle off the informations that may be printed.
mini.nb  A vector of integers of length two. This parameter is used in the “mini-em” initialization. The first integer sets how many times the algorithm is repeated; the second sets the maximum number of iterations the algorithm will do each time. For example, if init=“mini-em” and mini.nb=\(c(5,10)\), the algorithm will be launched 5 times, doing each time 10 iterations; finally the algorithm will begin with the initialization that maximizes the log-likelihood.

scaling  Logical: whether to scale the dataset (mean=0 and standard-error=1 for each variable) or not. By default the data is not scaled.

min.individuals  This parameter is used to control for the minimum population of a class. If the population of a class becomes strictly inferior to 'min.individuals’ then the algorithm stops and gives the message: 'pop<min.indiv.’. Here the meaning of "population of a class" is the sum of its posterior probabilities. The value of 'min.individuals’ cannot be lower than 2.

noise.ctrl  This parameter avoids to have a too low value of the ‘noise’ parameter b. It guarantees that the dimension selection process do not select too many dimensions (which leads to a potential too low value of the noise parameter b). When selecting the intrinsic dimensions using Cattell’s scree-test or BIC, the function doesn’t use the eigenvalues inferior to noise.ctrl, so that the intrinsic dimensions selected can’t be higher or equal to the order of these eigenvalues.

mc.cores  Positive integer, default is 1. If mc.cores>1, then parallel computing is used, using mc.cores cores. Warning for Windows users only: the parallel computing can sometimes be slower than using one single core (due to how parLapply works).

nb.rep  A positive integer (default is 1). Each estimation (i.e. combination of (model, K, threshold)) is repeated nb.rep times and only the estimation with the highest log-likelihood is kept.

keepAllRes  Logical. Should the results of all runs be kept? If so, an argument all_results is created in the results. Default is TRUE.

kmeans.control  A list. The elements of this list should match the parameters of the kmeans initialization (see kmeans help for details). The parameters are “iter.max”, “nstart” and “algorithm”.

d_max  A positive integer. The maximum number of dimensions to be computed. Default is 100. It means that the intrinsic dimension of any cluster cannot be larger than d_max. It quickens a lot the algorithm for datasets with a large number of variables (e.g. thousands).

d  DEPRECATED. This parameter is kept for retro compatibility. Now please use the parameter d_select.

details
Some information on the signification of the model names:

**Akj are the parameters of the classes subspaces:**  
- if Akj: each class has its parameters and there is one parameter for each dimension
- if Ak: the classes have different parameters but there is only one per class
• if $A_j$: all the classes have the same parameters for each dimension (it’s a particular case with a common orientation matrix)
• if $A$: all classes have the same one parameter

**Bk are the noises of the classes subspaces:**
• if $B_k$: each class has its proper noise
• if $B$: all classes have the same noise

**Qk is the orientation matrix of each class:**
• if $Q_k$: all classes have its proper orientation matrix
• if $Q$: all classes have the same orientation matrix

**Dk is the intrinsic dimension of each class:**
• if $D_k$: the dimensions are free and proper to each class
• if $D$: the dimension is common to all classes

The model “ALL” will compute all the models, give their BIC and keep the model with the highest BIC value. Instead of writing the model names, they can also be specified using an integer. 1 represents the most general model (“$A_k B_k Q_k D_k$”) while 14 is the most constrained (“$A B Q D$”), the others number/name matching are given below. Note also that several models can be run at once, by using a vector of models (e.g. model = c(“AKBQKD”,“AKJBQKD”,”ABQD”) is equivalent to model = c(8,4,13); to run the 6 first models, use model=1:6). If all the models are to be run, model=“all” is faster than model=1:14.

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<thead>
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</table>

The parameter $d$, is used to select the intrinsic dimensions of the subclasses. Here are his definitions:

• “Cattell”: The Cattell’s scree-test is used to gather the intrinsic dimension of each class. If the model is of common dimension (models 7 to 14), the scree-test is done on the covariance matrix of the whole dataset.

• “BIC”: The intrinsic dimensions are selected with the BIC criterion. See Bouveyron et al. (2010) for a discussion of this topic. For common dimension models, the procedure is done on the covariance matrix of the whole dataset.

• Note that ”Cattell” (resp. ”BIC”) can be abreviated to ”C” (resp. ”B”) and that this argument is not case sensitive.

The different initializations are:

**“param”**: it is initialized with the parameters, the means being generated by a multivariate normal distribution and the covariance matrix being common to the whole sample

**“mini-em”**: it is an initialization strategy, the classes are randomly initialized and the E-M algorithm makes several iterations, this action is repeted a few times (the default is 5 iterations and 10 times), at the end, the initialization choosen is the one which maximise the log-likelihood (see mini.nb for more information about its parametrization)
“random”: the classes are randomly given using a multinomial distribution

“kmeans”: the classes are initialized using the kmeans function (with: algorithm="Hartigan-Wong"; nstart=4; iter.max=50); note that the user can use his own arguments for kmeans using the dot-dot-dot argument

A prior class vector: It can also be directly initialized with a vector containing the prior classes of the observations. To do so use init="vector" and provide the vector in the argument init.vector.

The BIC criterion used in this function is to be maximized and is defined as $2^{\text{LL}} - k \cdot \log(n)$ where LL is the log-likelihood, $k$ is the number of parameters and $n$ is the number of observations.

Value

hddc returns an 'hdc' object; it’s a list containing:

- model: The name of the model.
- K: The number of classes.
- d: The dimensions of each class.
- a: The parameters of each class subspace.
- b: The noise of each class subspace.
- mu: The mean of each variable for each class.
- prop: The proportion of each class.
- ev: The eigen values of the var/covar matrix.
- Q: The orthogonal matrix of orientation of each class.
- loglik: The log-likelihood.
- loglik_all: The log-likelihood of all iterations.
- posterior: The matrix of the probabilities to belong to a class for each observation and each class.
- class: The class vector obtained by the clustering.
- com_ev: Only if this is a common dimension model. The eigenvalues of the var/covar matrix of the whole dataset.
- N: The number of observations.
- complexity: The number of parameters of the model.
- threshold: The threshold used for the Cattell scree-test.
- d_select: The way the dimensions were selected.
- BIC: The BIC of the model.
- ICL: The ICL of the model.
- criterion: The criterion used to select the model.
- call: The call.
- complexity_allModels: The number of parameters for all the models that were run (used to compute the slope heuristic).
The data.frame with the combination (model, K, threshold) and the associated values of the likelihood (LL), BIC and ICL, as well as the rank of each of the models with respect to the selection criterion.

Only if keepAllRes=TRUE. The parameters of all estimations that were run.

Only if scaling=TRUE. The centers and the standard deviation of the original dataset.

Laurent Berge, Charles Bouveyron and Stephane Girard


hdda, predict.hdc, plot.hdc.

# Example 1:
data <- simuldata(1000, 1000, 50)
X <- data$X
clx <- data$clx
Y <- data$Y
cly <- data$cly

#clustering of the simulated dataset:
prms1 <- hddc(X, K=3, algo="CEM", init='param')

#class vector obtained by the clustering:
prms1$class

#We can look at the adjusted rand index to assess the goodness of fit
res1 <- predict(prms1, X, clx)
res2 <- predict(prms1, Y)
#the class predicted using hddc parameters on the test dataset:
res2$class

# Example 2:
data(Crabs)

# clustering of the Crabs dataset:
prms3 <- hddc(Crabs[,-1], K=4, algo="EM", init='mini-em')
res3 <- predict(prms3, Crabs[,-1], Crabs[,1])
# another example using the Crabs dataset
prms4 <- hddc(Crabs[-1], K=1:8, model=c(1,2,7,9))

# model=c(1,2,7,9) is equivalent to:
# model=c("AKJBQKDk","AkBkQKDk","AkjbQKDk","ABQkDk")
res4 <- predict(prms4, Crabs[-1], Crabs[,-1])

# PARALLEL COMPUTING
## Not run:
## Same example but with Parallel Computing => platform specific
## (slower for Windows users)
## To enable it, just use the argument 'mc.cores'
prms5 <- hddc(Crabs[-1], K=1:8, model=c(1,2,7,9), mc.cores=2)

## End(Not run)

---

**hdmda**  
*Mixture Discriminant Analysis with HD Gaussians*

**Description**

HD-MDA implements mixture discriminant analysis (MDA, Hastie & Tibshirani, 1996) with HD Gaussians instead of full Gaussians. Each class is assumed to be made of several class-specific groups in which the data live in low-dimensional subspaces. From a technical point of view, a clustering is done using *hddc* in each class.

**Usage**

```r
hdmda(X, cls, K=1:10, model='AkjBkQkDk', show=FALSE, ...)
```

**Arguments**

- **X**  
  A matrix or a data frame of observations, assuming the rows are the observations and the columns the variables. Note that NAs are not allowed.

- **cls**  
  The vector of the class of each observations, its type can be numeric or string.

- **K**  
  A vector of integers specifying the number of clusters for which the BIC and the parameters are to be calculated; the function keeps the parameters which maximises the BIC. Note that the length of the vector K can't be larger than 20. Default is 1:10.

- **model**  
  A character string vector, or an integer vector indicating the models to be used. The available models are: "AkjBkQkDk" (default), "AkBkQkDk", "ABQkDk", "AkjBQkDk", "AkBQkDk", "AkjBkQkD", "AkBkQkD", "AkjBQkD", "AkBQkD", "ABQkD", "AjbQkD", "ABQD", "ABQD". It is not case sensitive and integers can be used instead of names, see details for more information. Several models can be used, if it is, only the results of the one which maximizes the BIC criterion is kept. To run all models, use model="ALL".
show Use show = TRUE to display some information related to the clustering.

Any argument that can be used by the function hddc.

Details

Some information on the signification of the model names:

**Akj are the parameters of the classes subspaces:**
- if Akj: each class has its parameters and there is one parameter for each dimension
- if Ak: the classes have different parameters but there is only one per class
- if Aj: all the classes have the same parameters for each dimension (it’s a particular case with a common orientation matrix)
- if A: all classes have the same one parameter

**Bk are the noises of the classes subspaces:**
- if Bk: each class has its proper noise
- if B: all classes have the same noise

**Qk is the orientation matrix of each class:**
- if Qk: all classes have their proper orientation matrix
- if Q: all classes have the same orientation matrix

**Dk is the intrinsic dimension of each class:**
- if Dk: the dimensions are free and proper to each class
- if D: the dimension is common to all classes

The model “all” will compute all the models, give their BIC and keep the model with the highest BIC value. Instead of writing the model names, they can also be specified using an integer. 1 represents the most general model (“AkjBkQkDk”) while 14 is the most constrained (“ABQD”), the others number/name matching are given below:

<table>
<thead>
<tr>
<th>Model Name</th>
<th>Integer Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>AkjBkQkDk</td>
<td>1</td>
</tr>
<tr>
<td>AkBkQkDk</td>
<td>2</td>
</tr>
<tr>
<td>ABkQkDk</td>
<td>3</td>
</tr>
<tr>
<td>AkjBQkDk</td>
<td>4</td>
</tr>
<tr>
<td>AkBQkDk</td>
<td>5</td>
</tr>
<tr>
<td>ABQkDk</td>
<td>6</td>
</tr>
<tr>
<td>AjBQD</td>
<td>13</td>
</tr>
</tbody>
</table>

**Value**

hdmda returns an 'hdmda' object which is a list containing:

- alpha Estimated prior probabilities for the classes.
- prms Estimated mixture parameters for each class.
- kname The name (level) of each class.

**Author(s)**

Laurent Berge, Charles Bouveyron and Stephane Girard
References


See Also

hdda, hddc

Examples

```r
# Load the Wine data set
data(wine)
cls = wine[,1]; X = scale(wine[,,-1])

# A simple use...
out = hdmda(X[1:100,],cls[1:100])
res = predict(out,X[101:nrow(X),])

# Comparison between hdmda and hdda in a CV setup
set.seed(123); nb = 10; Err = matrix(NA,2,nb)
for (i in 1:nb){
  cat('
')
  test = sample(nrow(X),50)
  out0 = lda(X[-test,],cls[-test])
  res0 = predict(out0,X[test,])
  Err[1,i] = sum(res0$class != cls[test]) / length(test)
  out = hdmda(X[-test,],cls[-test],K=3,model="AKJBQK0K")
  res = predict(out,X[test,])
  Err[2,i] = sum(res$class != cls[test]) / length(test)
}

boxplot(t(Err),names=c('LDA','HD-MDA'),col=2:3,ylab="CV classification error",
       main='CV classification error on Wine data')
```
Description

This function plots Cattell’s scree-test or the BIC selection, using parameters coming from \texttt{hdda} or \texttt{hddc} functions.

Usage

```r
## S3 method for class 'hdc'
plot(x, method = NULL, threshold = NULL, noise.ctrl=1e-8, ...)
```

Arguments

- `x`: A `hdc` class object obtained using \texttt{hdda} or \texttt{hddc} methods.
- `method`: The method used to select the intrinsic dimension. It can be "BIC" or "Cattell". By default it takes the method used when obtaining the parameters using \texttt{hdda} or \texttt{hddc}. Note that "Cattell" (resp. "BIC") can be abbreviated to "C" (resp. "B") and that this argument is not case sensitive.
- `threshold`: The threshold used in Cattell’s Scree-Test. By default it takes the threshold in the argument \(x\), if none, the default value of the threshold is 0.2.
- `noise.ctrl`: This parameter avoids to have a too low value of the 'noise' parameter \(b\). It garantees that the dimension selection process do not select too many dimensions (which leads to a potential too low value of the noise parameter \(b\)). When selecting the intrinsic dimensions using Cattell’s scree-test or BIC, the function doesn’t use the eigenvalues inferior to noise.ctrl, so that the intrinsic dimensions selected can’t be higher or equal to the order of these eigenvalues.

Value

- **If** `method = "Cattell"` The plot of the eigen values and of the sequential differences of the eigen values. The dimension to retain is the one before the last fall of the eigenvalues’ differences below the threshold.
- **If** `method = "BIC"` The BIC related to the dimension for each class. It stops after the first fall of the BIC.

Author(s)

Laurent Berge, Charles Bouveyron and Stephane Girard
References


See Also

hdda, hddc, predict.hdc.

Examples

```r
# Example 1 :
data(wine)
a <- wine[, -1]
z <- wine[, 1]
prms1 <- hdda(a, z, model = "AkBkQkDk", scaling = TRUE, d_select = "bic")

# the plot related to the selection that has been done: BIC
plot(prms1)

# it shows the plot of Cattell's scree-test, with a threshold of .3
plot(prms1, "Cattell", 0.3)

prms2 <- hdda(a, z, model = "AkBkQkDk", scaling = TRUE, d_select = "cattell")
# the plot related to the selection that has been done: Cattell's scree-test
plot(prms2)
# the plot of the BIC
plot(prms2, "b")
```

predict.hdc  
**Prediction method for 'hdc' class objects.**

Description

This function computes the class prediction of a dataset with respect to the model-based supervised and unsupervised classification methods hdda and hddc.

Usage

```r
## S3 method for class 'hdc'
predict(object, data, cls = NULL, ...)
```
**Arguments**

- **object**: An 'hdc' class object obtained by using hdda or hddc function.
- **data**: A matrix or a data frame of observations, assuming the rows are the observations and the columns the variables. Note that NAs are not allowed.
- **cls**: It is optional and used to be compared to the predicted classes, default is NULL.
- **...**: Arguments based from or to other methods. Not currently used.

**Value**

- **class**: vector of the predicted class.
- **prob**: The matrix of the probabilities to belong to a class for each observation and each class.

If the initial class vector is given to the argument ‘cls’ then the adjusted rand index (ARI) is also returned. Also the following object is returned:

- **ARI**: The confusion matrix of the classification.

**Author(s)**

Laurent Berge, Charles Bouveyron and Stephane Girard

**References**


**See Also**

- hdda, hddc.

**Examples**

```
# Example 1:
data <- simuldata(1000, 1000, 50)
X <- data$X
clx <- data$clx
Y <- data$Y
cly <- data$cly

#clustering of the gaussian dataset:
prms1 <- hddc(X, K=3, algo="CEM", init='param')

#class vector obtained by the clustering:
```
prms1$class

# only to see the good classification rate and
# the Adjusted Rand Index:
res1 <- predict(prms1, X, clx)
res2 <- predict(prms1, Y)

# the class predicted using hddc parameters on the test dataset:
res2$class

# Example 2:
data(Crabs)
# clustering of the Crabs dataset:
prms3 <- hddc(Crabs[, 1], K=4, algo="EM", init='kmeans')
res3 <- predict(prms3, Crabs[, 1], Crabs[, 1])

predict.hdmda

Prediction method for ‘hdmda’ class objects.

Description
This function computes the class prediction of a dataset with respect to the model-based supervised classification method \texttt{hdmda}.

Usage

\texttt{## S3 method for class ‘hdmda’}
\texttt{predict(object, X, \ldots)}

Arguments

\begin{itemize}
  \item \texttt{object} \hspace{1cm} An object of class ‘hdmda’.
  \item \texttt{X} \hspace{1cm} A matrix or a data frame of observations, assuming the rows are the observations and the columns the variables. Note that NAs are not allowed.
  \item \ldots \hspace{1cm} Arguments based from or to other methods. Not currently used.
\end{itemize}

Value

\begin{itemize}
  \item \texttt{class} \hspace{1cm} vector of the predicted class.
  \item \texttt{posterior} \hspace{1cm} The matrix of the probabilities to belong to a class for each observation and each class.
\end{itemize}

Author(s)
Laurent Berge, Charles Bouveyron and Stephane Girard
References


See Also

hdmda

Examples

# Load the Wine data set
data(wine)
cls = wine[,1]; X = scale(wine[,1:3])

# A simple use...
out = hmda(X[1:100,],cls[1:100])
res = predict(out,X[101:nrow(X),])

simuldata

Gaussian Data Generation

Description

This function generates two datasets according to the model \([A_kB_kQ_kD_k]\) of the HDDA gaussian mixture model paramatrisation (see ref.).

Usage

simuldata(nlearn, ntest, p, K = 3, prop = NULL, d = NULL, a = NULL, b = NULL)

Arguments

nlearn
  The size of the learning dataset to be generated.
ntest
  The size of the testing dataset to be generated.
p
  The number of variables.
K
  The number of classes.
prop  The proportion of each class.
d  The dimension of the intrinsic subspace of each class.
a  The value of the main parameter of each class.
b  The noise of each class.

Value

X  The learning dataset.
clx  The class vector of the learning dataset.
Y  The test dataset.
cly  The class vector of the test dataset.
prms  The principal parameters used to generate the datasets.

Author(s)

Laurent Berge, Charles Bouveyron and Stephane Girard

References


See Also

hddc, hdda.

Examples

data <- simuldata(500, 1000, 50, K=5, prop=c(0.2,0.25,0.25,0.15,0.15))
X <- data$X
clx <- data$clx
f <- hdda(X, clx)
Y <- data$Y
cly <- data$clly
e <- predict(f, Y, cly)

slopHeuristic  

Slope Heuristic for HDDC objects

Description

This function computes the slope heuristic for an

Usage

slopHeuristic(x)
Arguments

x

An hdc object, obtained from the function `hddc`.

Value

A list of two elements:

- **best_model**: The index of the best model, among all estimated models.
- **crit**: The slope heuristic criterion for each of the estimated models.

Examples

```r
# Clustering of the Crabs data set
data(Crabs)
prms = hddc(Crabs[,1], K=1:10)
slope = slopeHeuristic(prms)
plot(slope$crit) # The best model is indeed for 4 clusters
prms$all_results[[slope$best_model]] # we extract the best model
```

---

**wine**  

*Wine dataset*

Description

These data are the results of a chemical analysis of wines grown in the same region in Italy but derived from three different cultivars. The analysis determined the quantities of 13 constituents found in each of the three types of wines.

Usage

```r
data(wine)
```

Format

A data frame with 178 observations on the following 14 variables:

- **class**: The class vector, the three different cultivars of wine are represented by the three integers: 1 to 3.
- **V1**: Alcohol
- **V2**: Malic acid
- **V3**: Ash
- **V4**: Alcalinity of ash
- **V5**: Magnesium
- **V6**: Total phenols
V7 Flavanoids
V8 Nonflavanoid phenols
V9 Proanthocyanins
V10 Color intensity
V11 Hue
V12 OD280/OD315 of diluted wines
V13 Proline

Source
This dataset is from the UCI machine learning repository, provided here: http://archive.ics.uci.edu/ml/datasets/Wine.

Examples
data(wine)
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