

Package ‘KODAMA’

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Title Knowledge Discovery by Accuracy Maximization

Description An unsupervised and semi-supervised learning algorithm that performs feature extraction from noisy and high-dimensional data.

Depends R (>= 2.10.0), stats

Imports Rcpp (>= 0.12.4)

LinkingTo Rcpp, RcppArmadillo

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VignetteBuilder knitr

SuggestsNote No suggestions

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core_cpp	<i>Maximization of Cross-Validated Accuracy Methods</i>
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Description

This function performs the maximization of cross-validated accuracy by an iterative process

Usage

```
core_cpp(x,
        xTdata=NULL,
        clbest,
        Tcycle=20,
        FUN=c("KNN", "PLS-DA"),
        fpar=2,
        constrain=NULL,
        fix=NULL,
        shake=FALSE)
```

Arguments

x	a matrix.
xTdata	a matrix for projections. This matrix contains samples that are not used for the maximization of the cross-validated accuracy. Their classification is obtained by predicting samples on the basis of the final classification vector.
clbest	a vector to optimize.
Tcycle	number of iterative cycles that leads to the maximization of cross-validated accuracy.
FUN	classifier to be consider. Choices are "KNN" and "PLS-DA".
fpar	parameters of the classifier. If the classifier is KNN, fpar represents the number of neighbours. If the classifier is PLS-DA, fpar represents the number of components.

constrain	a vector of <code>nrow(data)</code> elements. Supervised constraints can be imposed by linking some samples in such a way that if one of them is changed, all other linked samples change in the same way (<i>i.e.</i> , they are forced to belong to the same class) during the maximization of the cross-validation accuracy procedure. Samples with the same identifying constrain will be forced to stay together.
fix	a vector of <code>nrow(data)</code> elements. The values of this vector must be TRUE or FALSE. By default all elements are FALSE. Samples with the TRUE fix value will not change the class label defined in <code>W</code> during the maximization of the cross-validation accuracy procedure. For more information refer to Cacciatore, <i>et al.</i> (2014).
shake	if <code>shake = FALSE</code> the cross-validated accuracy is computed with the class defined in <code>W</code> , before the maximization of the cross-validation accuracy procedure.

Value

The function returns a list with 3 items:

clbest	a classification vector with a maximized cross-validated accuracy.
accbest	the maximum cross-validated accuracy achieved.
vect_acc	a vector of all cross-validated accuracies obtained.
vect_proj	a prediction of samples in <code>xTdata</code> matrix using the vector <code>clbest</code> . This output is present only if <code>xTdata</code> is not NULL.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[KODAMA](#)

Examples

```
# Here, the famous (Fisher's or Anderson's) iris data set was loaded
data(iris)
u=as.matrix(iris[,-5])
s=sample(1:150,150,TRUE)

# The maximization of the accuracy of the vector s is performed
```

```
results=core_cpp(u, clbest=s, fpar = 10)

print(as.numeric(results$clbest))
```

dinisurface

Ulisse Dini Data Set Generator

Description

This function creates a data set based upon data points distributed on a Ulisse Dini's surface.

Usage

```
dinisurface(N=1000)
```

Arguments

N Number of data points.

Value

The function returns a three dimensional data set.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[helicoid](#), [swissroll](#), [spirals](#)

Examples

```
require("rgl")
x=dinisurface()
open3d()
plot3d(x, col=rainbow(1000),box=FALSE,size=3)
rgl.postscript("dinisurface.pdf", fmt="pdf")
```

`floyd`*Find Shortest Paths Between All Nodes in a Graph*

Description

The `floyd` function finds all shortest paths in a graph using Floyd's algorithm.

Usage

```
floyd(data)
```

Arguments

`data` matrix or distance object

Value

`floyd` returns a matrix with the total lengths of the shortest path between each pair of points.

Author(s)

Stefano Cacciatore

References

Floyd RW. Algorithm 97: Shortest path. Commun ACM 5(6):345.

Examples

```
# build a graph with 5 nodes
x=matrix(c(0,NA,NA,NA,NA,30,0,NA,NA,NA,10,NA,0,NA,NA,NA,70,50,0,10,NA,40,20,60,0),ncol=5)
print(x)

# compute all path lengths
z=floyd(x)
print(z)
```

helicoid

Helicoid Data Set Generator

Description

This function creates a data set based upon data points distributed on a Helicoid surface.

Usage

```
helicoid(N=1000)
```

Arguments

N Number of data points.

Value

The function returns a three dimensional data set.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[swissroll](#), [dini](#), [surface](#), [spirals](#)

Examples

```
require("rgl")
x=helicoid()
open3d()
plot3d(x, col=rainbow(1000),box=FALSE,size=3)
rgl.postscript("helicoid.pdf", fmt="pdf")
```

`k.test`*K-Test of Statistical Association*

Description

This function performs a permutation test using PLS to assess association between the KODAMA output and any additional related parameters such as clinical metadata.

Usage

```
k.test(data, labels, n = 100)
```

Arguments

<code>data</code>	a matrix.
<code>labels</code>	a classification vector.
<code>n</code>	number of iterations of the permutation test.

Value

The p-value of the test.

Author(s)

Stefano Cacciatore

References

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[KODAMA](#)

Examples

```
# data(iris)
# data=iris[,-5]
# labels=iris[,5]
# kk=KODAMA(data,FUN="PLS-DA")
# kkplot=kk$pp
# k1=k.test(kkplot,labels)
# print(k1)
# k2=k.test(kkplot,sample(labels))
# print(k2)
```

knn.double.cv

*Cross-Validation with k-Nearest Neighbors algorithm.***Description**

This function performs a 10-fold cross validation on a given data set using *k*-Nearest Neighbors (*k*NN) model. To assess the prediction ability of the model, a 10-fold cross-validation is conducted by generating splits with a ratio 1:9 of the data set, that is by removing 10% of samples prior to any step of the statistical analysis, including PLS component selection and scaling. Best number of component for PLS was carried out by means of 10-fold cross-validation on the remaining 90% selecting the best Q_{2y} value. Permutation testing was undertaken to estimate the classification/regression performance of predictors.

Usage

```
knn.double.cv(Xdata,
              Ydata,
              constrain=1:nrow(Xdata),
              compmax=min(c(ncol(Xdata),nrow(Xdata))),
              perm.test=FALSE)
```

Arguments

Xdata	a matrix.
Ydata	the responses. If Ydata is a numeric vector, a regression analysis will be performed. If Ydata is factor, a classification analysis will be performed.
constrain	a vector of nrow(data) elements. Sample with the same identifying constrain will be split in the training set or in the test set of cross-validation together.
compmax	the number of latent components to be used for classification.
perm.test	a classification vector.

Value

A list with the following components:

Ypred	the vector containing the predicted values of the response variables obtained by cross-validation.
Yfit	the vector containing the fitted values of the response variables.
Q2Y	Q _{2y} value.
R2Y	R _{2y} value.
conf	The confusion matrix (only in classification mode).
acc	The cross-validated accuracy (only in classification mode).

Author(s)

Stefano Cacciatore

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

Examples

```
data(iris)
data=iris[,-5]
labels=iris[,5]
pp=knn.double.cv(data,labels)
print(pp$Q2Y)
table(pp$Ypred,labels)

#
# data(MetRef)
# u=MetRef$data;
# u=u[,-which(colSums(u)==0)]
# u=normalization(u)$newXtrain
# u=scaling(u)$newXtrain
# pp=knn.double.cv(u,as.factor(MetRef$donor))
# print(pp$Q2Y)
# table(pp$Ypred,MetRef$donor)
#
```

knn.kodama

k-Nearest Neighbors Classifier.

Description

k-nearest neighbour classification for a test set from a training set.

Usage

```
knn.kodama(Xtrain, Ytrain, Xtest, k)
```

Arguments

Xtrain	a matrix of training set cases.
Ytrain	a classification vector.
Xtest	a matrix of test set cases.
k	the number of nearest neighbors to consider.

Details

The function utilizes the Approximate Nearest Neighbor (ANN) C++ library, which can give the exact nearest neighbours or (as the name suggests) approximate nearest neighbours to within a specified error bound. For more information on the ANN library please visit <http://www.cs.umd.edu/~mount/ANN/>.

Value

The function returns a vector of predicted labels.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Bentley JL (1975)
Multidimensional binary search trees used for associative search.
Communication ACM 1975;18:309-517.

Arya S, Mount DM
Approximate nearest neighbor searching
Proc. 4th Ann. ACM-SIAM Symposium on Discrete Algorithms (SODA'93);271-280.

Arya S, Mount DM, Netanyahu NS, Silverman R, Wu AY
An optimal algorithm for approximate nearest neighbor searching
Journal of the ACM 1998;45:891-923.

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[KODAMA](#)

Examples

```
data(iris)
data=iris[,-5]
labels=iris[,5]
```

```

ss=sample(150,15)

z=knn.kodama(data[-ss,], labels[-ss], data[ss,], k=5)
table(z[,5],labels[ss])

```

KODAMA

*Knowledge Discovery by Accuracy Maximization***Description**

KODAMA (KnOwledge Discovery by Accuracy MAximization) is an unsupervised and semi-supervised learning algorithm that performs feature extraction from noisy and high-dimensional data. Unlike other data mining methods, KODAMA is driven by an integrated procedure of cross validation of the results.

Usage

```

KODAMA(data,
        M = 100,
        Tcycle = 20,
        FUN_VAR = function(x) { ceiling(ncol(x)) },
        FUN_SAM = function(x) { ceiling(nrow(x) * 0.75)},
        bagging = FALSE,
        FUN = c("KNN", "PLS-DA"),
        f.par = 5,
        W = NULL,
        constrain = NULL,
        fix=NULL,
        epsilon = 0.05,
        dims=2,
        landmarks=5000)

```

Arguments

data	a matrix.
M	number of iterative processes (step I-III).
Tcycle	number of iterative cycles that leads to the maximization of cross-validated accuracy.
FUN_VAR	function to select the number of variables to select randomly. By default all variable are taken.
FUN_SAM	function to select the number of samples to select randomly. By default the 75 per cent of all samples are taken.
bagging	Should sampling be with replacement, bagging = TRUE. By default bagging = FALSE.
FUN	classifier to be considered. Choices are "KNN" and "PLS-DA".
f.par	parameters of the classifier.

<code>W</code>	a vector of <code>nrow(data)</code> elements. The KODAMA procedure can be started by different initializations of the vector <code>W</code> . Without any <i>a priori</i> information the vector <code>W</code> can be initialized with each element being different from the others (<i>i.e.</i> , each sample categorized in a one-element class). Alternatively, the vector <code>W</code> can be initialized by a clustering procedure, such as <code>kmeans</code> .
<code>constrain</code>	a vector of <code>nrow(data)</code> elements. Supervised constraints can be imposed by linking some samples in such a way that if one of them is changed the remaining linked samples must change in the same way (<i>i.e.</i> , they are forced to belong to the same class) during the maximization of the cross-validation accuracy procedure. Samples with the same identifying <code>constrain</code> will be forced to stay together.
<code>fix</code>	a vector of <code>nrow(data)</code> elements. The values of this vector must to be <code>TRUE</code> or <code>FALSE</code> . By default all elements are <code>FALSE</code> . Samples with the <code>TRUE</code> <code>fix</code> value will not change the class label defined in <code>W</code> during the maximization of the cross-validation accuracy procedure.
<code>epsilon</code>	cut-off value for low proximity. High proximity are typical of intracluster relationships, whereas low proximities are expected for intercluster relationships. Very low proximities between samples are ignored by (default) setting <code>epsilon = 0.05</code> .
<code>dims</code>	dimensions of the configurations of Sammon's non-linear mapping based on the KODAMA dissimilarity matrix.
<code>landmarks</code>	number of landmarks to use.

Details

KODAMA consists of five steps. These can be in turn divided into two parts: (i) the maximization of cross-validated accuracy by an iterative process (step I and II), resulting in the construction of a proximity matrix (step III), and (ii) the definition of a dissimilarity matrix (step IV and V). The first part entails the core idea of KODAMA, that is, the partitioning of data guided by the maximization of the cross-validated accuracy. At the beginning of this part, a fraction of the total samples (defined by `FUN_SAM`) are randomly selected from the original data. The whole iterative process (step I-III) is repeated `M` times to average the effects owing to the randomness of the iterative procedure. Each time that this part is repeated, a different fraction of samples is selected. The second part aims at collecting and processing these results by constructing a dissimilarity matrix to provide a holistic view of the data while maintaining their intrinsic structure (steps IV and V). Then, Sammon's non-linear mapping is used to visualise the results of KODAMA dissimilarity matrix.

Value

The function returns a list with 4 items:

<code>dissimilarity</code>	a dissimilarity matrix.
<code>acc</code>	a vector with the <code>M</code> cross-validated accuracies.
<code>proximity</code>	a proximity matrix.
<code>v</code>	a matrix containing the all classification obtained maximizing the cross-validation accuracy.
<code>pp</code>	a matrix containing the score of the Sammon's non-linear mapping.

res a matrix containing all classification vectors obtained through maximizing the cross-validation accuracy.

f.par parameters of the classifier.

entropy Shannon's entropy of the KODAMA proximity matrix.

landpoints indexes of the landmarks used.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

Examples

```
data(iris)
data=iris[,-5]
labels=iris[,5]
kk=KODAMA(data)
plot(kk$pp,col=as.numeric(labels), xlab="First component", ylab="Second component",cex=2)
```

loads *Variable Ranking*

Description

This function can be used to extract the variable ranking when KODAMA is performed with the PLS-DA classifier.

Usage

```
loads(model,method=c("loadings","kruskal.test"))
```

Arguments

model output of KODAMA.

method method to be used. Choices are "loadings" and "kruskal.test".

Value

The function returns a vector of values indicating the "importance" of each variable. If "method="loadings" the average of the loading of the first component of PLS models based on the cross-validated accuracy maximized vector is computed. If "method="kruskal.test" the average of minus logarithm of p-value of Kruskal-Wallis Rank Sum test is computed.

Author(s)

Stefano Cacciatore

References

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[KODAMA](#)

Examples

```
# data(iris)
# data=iris[,-5]
# labels=iris[,5]
# kk=KODAMA(data,FUN="PLS-DA")
# loads(kk)
```

lymphoma

Lymphoma Gene Expression Dataset

Description

This dataset consists of gene expression profiles of the three most prevalent adult lymphoid malignancies: diffuse large B-cell lymphoma (DLBCL), follicular lymphoma (FL), and B-cell chronic lymphocytic leukemia (B-CLL). The dataset consists of 4,682 mRNA genes for 62 samples (42 samples of DLBCL, 9 samples of FL, and 11 samples of B-CLL). Missing value are imputed and data are standardized as described in Dudoit, *et al.* (2002).

Usage

```
data(lymphoma)
```

Value

A list with the following elements:

data	Gene expression data. A matrix with 62 rows and 4,682 columns.
class	Class index. A vector with 62 elements.

References

- Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.
- Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.
- Alizadeh AA, Eisen MB, Davis RE, *et al.*
Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling.
Nature 2000;403(6769):503-511.
- Dudoit S, Fridlyand J, Speed TP
Comparison of discrimination methods for the classification of tumors using gene expression data.
J Am Stat Assoc 2002;97(417):77-87.

Examples

```
data(lymphoma)
class=1+as.numeric(lymphoma$class)
cc=prcomp(lymphoma$data)$x
plot(cc,pch=21,bg=class,xlab="First Component",ylab="Second Component")

#
# kk=KODAMA(lymphoma$data)
# plot(kk$pp,pch=21,bg=class,xlab="First Component",ylab="Second Component")
#
```

mcplot

Evaluation of the Monte Carlo accuracy results

Description

This function can be used to plot the accuracy values obtained during KODAMA procedure.

Usage

```
mcplot(model)
```

Arguments

model output of KODAMA.

Author(s)

Stefano Cacciatore

References

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[KODAMA](#)

Examples

```
data=as.matrix(iris[,-5])  
kk=KODAMA(data)  
mcplot(kk)
```

MetRef

Nuclear Magnetic Resonance Spectra of Urine Samples

Description

The data belong to a cohort of 22 healthy donors (11 male and 11 female) where each provided about 40 urine samples over the time course of approximately 2 months, for a total of 873 samples. Each sample was analysed by Nuclear Magnetic Resonance Spectroscopy. Each spectrum was divided in 450 spectral bins.

Usage

```
data(MetRef)
```

Value

A list with the following elements:

data	Metabolomic data. A matrix with 873 rows and 450 columns.
gender	Gender index. A vector with 873 elements.
donor	Donor index. A vector with 873 elements.

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

Examples

```

data(MetRef)
u=MetRef$data;
u=u[,-which(colSums(u)==0)]
u=normalization(u)$newXtrain
u=scaling(u)$newXtrain
class=as.numeric(as.factor(MetRef$gender))
cc= prcomp(u)$x
plot(cc,pch=21,bg=class,xlab="First Component",ylab="Second Component")

class=as.numeric(as.factor(MetRef$donor))
plot(cc,pch=21,bg=rainbow(22)[class],xlab="First Component",ylab="Second Component")

#
# kk=KODAMA(u)
# plot(kk$pp,pch=21,bg=rainbow(22)[class],xlab="First Component",ylab="Second Component")
#

```

normalization

Normalization methods

Description

Collection of Different Normalization Methods.

Usage

```
normalization(Xtrain,Xtest=NULL, method = "pqn",ref=NULL)
```

Arguments

Xtrain	a matrix of data (training data set).
Xtest	a matrix of data (test data set).(by default = NULL).
method	the normalization method to be used. Choices are "none", "pqn", "sum", "median", "sqrt" (by default = "pqn"). A partial string sufficient to uniquely identify the choice is permitted.
ref	Reference sample for Probabilistic Quotient Normalization. (by default = NULL).

Details

A number of different normalization methods are provided:

- "none": no normalization method is applied.
- "pqn": the Probabilistic Quotient Normalization is computed as described in *Dieterle, et al.* (2006).

- "sum": samples are normalized to the sum of the absolute value of all variables for a given sample.
- "median": samples are normalized to the median value of all variables for a given sample.
- "sqrt": samples are normalized to the root of the sum of the squared value of all variables for a given sample.

Value

The function returns a list with 2 items or 4 items (if a test data set is present):

<code>newXtrain</code>	a normalized matrix (training data set).
<code>coeXtrain</code>	a vector of normalization coefficient of the training data set.
<code>newXtest</code>	a normalized matrix (test data set).
<code>coeXtest</code>	a vector of normalization coefficient of the test data set.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Dieterle F, Ross A, Schlotterbeck G, Senn H.
 Probabilistic Quotient Normalization as Robust Method to Account for Dilution of Complex Biological Mixtures. Application in ¹H NMR Metabolomics.
Anal Chem 2006;78:4281-90.

Cacciatore S, Luchinat C, Tenori L
 Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
 KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[scaling](#)

Examples

```
data(MetRef)
u=MetRef$data;
u=u[,-which(colSums(u)==0)]
u=normalization(u)$newXtrain
u=scaling(u)$newXtrain
class=as.numeric(as.factor(MetRef$gender))
cc= prcomp(u)$x
plot(cc,pch=21,bg=class,xlab="First Component",ylab="Second Component")
```

Description

This function performs a 10-fold cross validation on a given data set using Partial Least Squares (PLS) model. To assess the prediction ability of the model, a 10-fold cross-validation is conducted by generating splits with a ratio 1:9 of the data set, that is by removing 10% of samples prior to any step of the statistical analysis, including PLS component selection and scaling. Best number of component for PLS was carried out by means of 10-fold cross-validation on the remaining 90% selecting the best Q_{2y} value. Permutation testing was undertaken to estimate the classification/regression performance of predictors.

Usage

```
pls.double.cv(Xdata,
              Ydata,
              constrain=1:nrow(Xdata),
              compmax=min(c(ncol(Xdata),nrow(Xdata))),
              perm.test=FALSE)
```

Arguments

Xdata	a matrix.
Ydata	the responses. If Ydata is a numeric vector, a regression analysis will be performed. If Ydata is factor, a classification analysis will be performed.
constrain	a vector of nrow(data) elements. Sample with the same identifying constrain will be split in the training set or in the test set of cross-validation together.
compmax	the number of latent components to be used for classification.
perm.test	a classification vector.

Value

A list with the following components:

B	the (p x m x length(ncomp)) array containing the regression coefficients. Each row corresponds to a predictor variable and each column to a response variable. The third dimension of the matrix B corresponds to the number of PLS components used to compute the regression coefficients. If ncomp has length 1, B is just a (p x m) matrix.
Ypred	the vector containing the predicted values of the response variables obtained by cross-validation.
Yfit	the vector containing the fitted values of the response variables.
P	the (p x max(ncomp)) matrix containing the X-loadings.
Q	the (m x max(ncomp)) matrix containing the Y-loadings.

T	the (ntrain x max(ncomp)) matrix containing the X-scores (latent components)
R	the (p x max(ncomp)) matrix containing the weights used to construct the latent components.
Q2Y	Q2y value.
R2Y	R2y value.
R2X	vector containing the explained variance of X by each PLS component.

Author(s)

Stefano Cacciatore

References

Cacciatore S, Luchinat C, Tenori L
 Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
 KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

Examples

```
data(iris)
data=iris[,-5]
labels=iris[,5]
pp=pls.double.cv(data,labels)
print(pp$Q2Y)
table(pp$Ypred,labels)

#
# data(MetRef)
# u=MetRef$data;
# u=u[,-which(colSums(u)==0)]
# u=normalization(u)$newXtrain
# u=scaling(u)$newXtrain
# pp=pls.double.cv(u,as.factor(MetRef$donor))
# print(pp$Q2Y)
# table(pp$Ypred, MetRef$donor)
#
```

pls.kodama

Partial Least Squares regression.

Description

Partial Least Squares (PLS) regression for test set from training set.

Usage

```
pls.kodama(Xtrain, Ytrain, Xtest, ncomp)
```

Arguments

Xtrain	a matrix of training set cases.
Ytrain	a classification vector.
Xtest	a matrix of test set cases.
ncomp	the number of components to consider.

Value

A list with the following components:

B	the (p x m x length(ncomp)) matrix containing the regression coefficients. Each row corresponds to a predictor variable and each column to a response variable. The third dimension of the matrix B corresponds to the number of PLS components used to compute the regression coefficients. If ncomp has length 1, B is just a (p x m) matrix.
Ypred	the (ntest x m x length(ncomp)) containing the predicted values of the response variables for the observations from Xtest. The third dimension of the matrix Ypred corresponds to the number of PLS components used to compute the regression coefficients.
P	the (p x max(ncomp)) matrix containing the X-loadings.
Q	the (m x max(ncomp)) matrix containing the Y-loadings.
T	the (ntrain x max(ncomp)) matrix containing the X-scores (latent components)
R	the (p x max(ncomp)) matrix containing the weights used to construct the latent components.
meanX	the p-vector containing the means of the columns of Xtrain.

Author(s)

Stefano Cacciatore

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennet PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also[KODAMA](#)**Examples**

```

data(iris)
data=iris[,-5]
labels=iris[,5]
ss=sample(150,15)
ncomponent=3

z=pls.kodama(data[-ss,], labels[-ss], data[ss,], ncomp=ncomponent)
table(z$Ypred[,ncomponent],labels[ss])

```

scaling

*Scaling methods***Description**

Collection of Different Scaling Methods.

Usage

```
scaling(Xtrain,Xtest=NULL, method = "autoscaling")
```

Arguments

Xtrain	a matrix of data (training data set).
Xtest	a matrix of data (test data set).(by default = NULL).
method	the scaling method to be used. Choices are "none", "centering", "autoscaling", "rangescaling", "paretoscaling" (by default = "autoscaling"). A partial string sufficient to uniquely identify the choice is permitted.

Details

A number of different scaling methods are provided:

- "none": no scaling method is applied.
- "centering": centers the mean to zero.
- "autoscaling": centers the mean to zero and scales data by dividing each variable by the variance.
- "rangescaling": centers the mean to zero and scales data by dividing each variable by the difference between the minimum and the maximum value.
- "paretoscaling": centers the mean to zero and scales data by dividing each variable by the square root of the standard deviation. Unit scaling divides each variable by the standard deviation so that each variance equal to 1.

Value

The function returns a list with 1 item or 2 items (if a test data set is present):

`newXtrain` a scaled matrix (training data set).
`newXtest` a scale matrix (test data set).

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

van den Berg RA, Hoefsloot HCJ, Westerhuis JA, *et al.*
Centering, scaling, and transformations: improving the biological information content of metabolomics data.
BMC Genomics 2006;7(1):142.

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[normalization](#)

Examples

```
data(MetRef)
u=MetRef$data;
u=u[,-which(colSums(u)==0)]
u=normalization(u)$newXtrain
u=scaling(u)$newXtrain
class=as.numeric(as.factor(MetRef$gender))
cc= prcomp(u)$x
plot(cc,pch=21,bg=class,xlab="First Component",ylab="Second Component")
```

spirals

Spirals Data Set Generator

Description

Produces a data set of spiral clusters.

Usage

```
spirals(n=c(100,100,100),sd=c(0,0,0))
```

Arguments

n a vector of integer. The length of the vector is the number of clusters and each number corresponds to the number of data points in each cluster.

sd amount of noise for each spiral.

Value

The function returns a two dimensional data set.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[helicoid](#),[dinisurface](#),[swissroll](#)

Examples

```
pdf(file="spirals.pdf")
par(mfrow=c(2,2),mai=c(0,0,0,0))
v1=spirals(c(100,100,100),c(0.1,0.1,0.1))
plot(v1,col=rep(2:4,each=100))
v2=spirals(c(100,100,100),c(0.1,0.2,0.3))
plot(v2,col=rep(2:4,each=100))
v3=spirals(c(100,100,100,100,100),c(0,0,0.2,0,0))
plot(v3,col=rep(2:6,each=100))
v4=spirals(c(20,40,60,80,100),c(0.1,0.1,0.1,0.1,0.1))
plot(v4,col=rep(2:6,c(20,40,60,80,100)))
dev.off()
```

`swissroll`*Swiss Roll Data Set Generator*

Description

Computes the Swiss Roll data set of a given number of data points.

Usage

```
swissroll(N=1000)
```

Arguments

N Number of data points.

Value

The function returns a three dimensional matrix.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Balasubramanian M, Schwartz EL
The isomap algorithm and topological stability.
Science 2002;295(5552):7.

Roweis ST, Saul LK
Nonlinear dimensionality reduction by locally linear embedding.
Science 2000;290(5500):2323-6.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

See Also

[helicoid](#),[dini](#),[surface](#),[spirals](#)

Examples

```
require("rgl")
x=swissroll()
open3d()
plot3d(x, col=rainbow(1000),box=FALSE,size=3)
rgl.postscript("swissroll.pdf", fmt="pdf")
```

transformy

Conversion Classification Vector to Matrix

Description

This function converts a classification vector into a classification matrix.

Usage

```
transformy(y)
```

Arguments

y a vector or factor.

Details

This function converts a classification vector into a classification matrix.

Value

A matrix.

Author(s)

Stefano Cacciatore and Leonardo Tenori

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

Examples

```
y=rep(1:10,3)
print(y)
z=transformy(y)
print(z)
```

Description

This dataset consists of the spoken, not written, addresses from 1900 until the sixth address by Barack Obama in 2014. Punctuation characters, numbers, words shorter than three characters, and stop-words (e.g., "that", "and", and "which") were removed from the dataset. This resulted in a dataset of 86 speeches containing 834 different meaningful words each. Term frequency-inverse document frequency (TF-IDF) was used to obtain feature vectors. It is often used as a weighting factor in information retrieval and text mining. The TF-IDF value increases proportionally to the number of times a word appears in the document, but is offset by the frequency of the word in the corpus, which helps to control for the fact that some words are generally more common than others.

Usage

```
data(USA)
```

Value

A list with the following elements:

data	TF-IDF data. A matrix with 86 rows and 834 columns.
year	Year index. A vector with 86 elements.
president	President index. A vector with 86 elements.

References

Cacciatore S, Luchinat C, Tenori L
Knowledge discovery by accuracy maximization.
Proc Natl Acad Sci U S A 2014;111(14):5117-22.

Cacciatore S, Tenori L, Luchinat C, Bennett PR, MacIntyre DA
KODAMA: an updated R package for knowledge discovery and data mining.
Bioinformatics. Submitted.

Examples

```
# Here is reported the analysis on the State of the Union  
# of USA president as shown in Cacciatore, et al. (2014)  
#  
# data(USA)  
# kk=KODAMA(USA$data)  
# cc=cmdscale(kk$dissimilarity)  
# par(cex=0.5,mar=c(15,6,2,2));  
# plot(USA$year,cc[,1],axes=F,pch=20,xlab="",ylab="First Component");  
# axis(1,at=USA$year,labels=rownames(USA$data),las=2);  
# axis(2,las=2);
```

```
# box()  
#
```

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