Package ‘bgmm’

February 19, 2015

Type Package

Title Gaussian Mixture Modeling Algorithms And The Belief-based Mixture Modeling

Version 1.7

Date 2014-08-18

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Description Two partially supervised mixture modeling methods: soft-label and belief-based modeling are implemented. For completeness, we equipped the package also with the functionality of unsupervised, semi- and fully supervised mixture modeling. The package can be applied also to selection of the best-fitting from a set of models with different component numbers or constraints on their structures. For detailed introduction see:


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Depends R (>= 2.0), mvtnorm, car, lattice, combinat

URL http://bgmm.molgen.mpg.de/

License GPL-3

Repository CRAN

NeedsCompilation no

Date/Publication 2014-12-19 15:57:54

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bgmm-package Belief-based Gaussian mixture modeling

Description

This package implements partially supervised mixture modeling methods: soft-label and belief-based modeling, the semi-supervised methods and for completeness also unsupervised and fully supervised methods for mixture modeling.

Details

Package: bgmm
Type: Package
Version: 1.6
Date: 2013-11-18
License: GPL-3
LazyLoad: yes

For short overview see the webpage http://bgmm.molgen.mpg.de/rapBGMM/.

Author(s)

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References

See Also

Package for unsupervised learning of Gaussian mixture model link{mclust}, methods for supervised learning link{MASS::lda()}, link{MASS::qda()}. 

Examples

```r
## Do not run
## It could take more than one minute
demo(bgmm)
```

---

**CellCycle**  
*Data for clustering of 384 cell cycle genes into five clusters corresponding to cell cycle phases*

**Description**

Time course expression data for 384 cell cycle genes (Cho et al., 1998). Literature examples of genes that should, and of genes that should not peak at each time point are given. For each cycle phase, there is a characteristic binary profile, stating when the phase occurs.

**Usage**

data(CellCycle)

**Format**

- CellCycleData: list: 17x384
- CellCycleBeliefs: list: 17x (35x2)
- CellCycleCenters: matrix: 5x17
- CellCycleClass: vector: 384

**Details**

- **CellCycleData**: A list, where each entry corresponds to one time-point. A given time point entry contains a vector with expression ratios for 384 cell cycle genes measured in this time point.
- **CellCycleBeliefs**: A list, where each entry corresponds to one time-point. A given time point entry gives the certainty (belief/plausibility) for each out of 35 example genes. Out of the genes, seven are known to peak in this time point and the remaining 28 are known to peak in other cycle phases.
- **CellCycleCenters**: A matrix, where the columns are the 17 time-points and the rows to the five cell phase clusters. A given entry in the matrix is equal to 1 if the genes from the cluster should peak in the time point, and 0 otherwise.
- **CellCycleClass**: Gives the true cluster for each gene. Each cluster corresponds to a cell cycle phase.

**Author(s)**

Ewa Szczurek
chooseModels

References


See Also

miRNA, Ste12

Examples

```r
library(bgmm)
data(CellCycle)
print(CellCycleData)
print(CellCycleBeliefs)
print(CellCycleCenters)
print(CellCycleClass)
```

chooseModels is a function for selecting a subset of fitted models.

**Description**

The function `chooseModels` extracts a sublist of models that match constraints on the number of components or on the model structure. The function `chooseOptimal` returns the model which is the best according to the given model selection criteria.

**Usage**

```r
chooseModels(models, kList = NULL, struct = NULL)
chooseOptimal(models, penalty=2, ...)
```

**Arguments**

- `models`: an object of the class `mModellist` which represents a list of fitted models.
- `kList`: a vector which specifies the requested numbers of Gaussian components (constraints on the number of components).
- `struct`: a vector which specifies four letter abbreviations of names of the requested model structures (constraints on the model structure).
- `penalty`: a penalty parameter in the GIC criteria. This parameter can be a single number or a string, either "BIC" or "AIC".
- `...`: other arguments that will be passed to the `getGIC` function.
chooseModels

Details

The function chooseModels() extracts a sublist of models from the models argument. The returned sublist is also an object of the class mModellist and is composed of models that simultaneously satisfy the restrictions of the number of Gaussian components defined by klist and restrictions of the model structure defined by struct. If the argument klist is set to NULL then no restrictions of the number of components are applied, same with the argument struct.

The function chooseOptimal() returns an object of the class mModel which is the single model that has the best (smallest) GIC score.

Value

An object of the class mModellist or mModel.

Author(s)

Przemyslaw Biecek

References


Examples

```r
simulated = simulateData(d=2, k=3, n=100, m=50, cov="0", within="E", n.labels=2)
models3 = belieflist(X=simulated$X, knowns=simulated$knowns, B=simulated$B, kList=2:4, mean="D", within="D")
plotGIC(models3, penalty="BIC")
```

## Do not run
## It could take more than one minute
# simulated = simulateData(d=2, k=3, n=300, m=60, cov="0", within="E", n.labels=2)
# # models3 = belieflist(X=simulated$X, knowns=simulated$knowns, B=simulated$B, kList=2:7, mean="D")
# # plotGIC(models3, penalty="BIC")
# # models4 = chooseModels(models3, kList=2:5, struct=c("DDDD","DDED","DDE0"))
# plot(models4)
# # plotGIC(models4, penalty="BIC")
# # model4 = chooseOptimal(models3, "BIC")
# plot(model4)
```
crossval

\textit{k-fold cross-validation for the specified model}

Description

The function \texttt{crossval()} performs k-fold cross-validation.

Usage

\begin{verbatim}
crossval(model = NULL, X = NULL, knowns = NULL, class = NULL,
    k = length(unique(class)), B = NULL, P = NULL, model.structure = getModelStructure(),
    ..., folds = 2, fun = belief)
\end{verbatim}

Arguments

- \texttt{model} an object of the class \texttt{mModel}.
- \texttt{X} a data.frame with unknown realizations. If not supplied \texttt{X} is extracted from the \texttt{model} argument.
- \texttt{knowns} a data.frame with labeled realizations. If not supplied \texttt{knowns} is extracted from the \texttt{model} argument.
- \texttt{class, B, P} a vector of classes, beliefs and plausibilities. If not supplied they will be extracted from the \texttt{model} argument.
- \texttt{fun} function that will be used for modeling, one of \texttt{supervised, unsupervised, belief, soft, semisupervised}.
- \texttt{model.structure, k, ...} arguments that will be passed to \texttt{fun} function.
- \texttt{folds} number of folds in k-fold cross validation. Cannot be greater that number of labeled samples.

Details

The function \texttt{crossval()} divides the dataset into k equal subsets, the number of labeled cases versus number of unlabeled cases is keep as close to constant as possible (the subset are generated with stratification). Then each subset is used as test set against a train set build from all remaining sets. In total k new models are estimated thus this procedure is time consuming.

For each model the error is calculated as average absolute differences between the distribution of estimated posteriors and distribution of beliefs/plausibilities for labeled cases.

Value

The list with three vectors: errors calculated as mean absolute differences between estimated posteriors and initial beliefs for known cases, indexes of folds for both labeled and unlabeled cases.
**DEprobs**

**Author(s)**
Przemyslaw Biecek

**References**

**Examples**

```r
set.seed(1313)
simulated = simulateData(d=2, k=3, n=300, m=60, cov="0", within="E", n.labels=2)
amodel = belief(X=simulated$X, knowns=simulated$knowns, B=simulated$B, k=4)
str(crossval(model=amodel, folds=6))
```

---

**DEprobs**

*Signed probabilities of differential expression*

**Description**
The `DEprobs` function is an application of mixture modeling to differential gene expression analysis. The function takes as input a two- or three-component model of one-dimensional gene expression data. The data is assumed to represent log fold change expression values and be negative when the corresponding genes are down-regulated. The function calculates probabilities of differential expression for the data and gives them a sign according to the sign of the data.

**Usage**

```r
DEprobs(model, verbose=FALSE)
```

**Arguments**

- `model` an object of the class `mModel`,
- `verbose` indicates whether log messages should be printed out.

**Details**
Given the input `model`, the function identifies the component which corresponds to the differentially expressed genes as the one which looks differential according to the posterior probabilities.

For input models with two Gaussian components the differential component should be the one with a broader range (encompassing the other), or the one with higher deviation from 0 (we assume the data are centered around 0).

For input models with three Gaussian components there are two differential components: one corresponding to the down-regulated genes, and one corresponding to the up-regulated genes. Those components are identified as the ones with the lowest and the highest mean, respectively.

For `verbose=TRUE` the index of the differential component is printed out.
Genotypes

Value
An list with the following elements:

- `diff.p.X` a vector with the calculated signed differential expression probabilities for the unlabeled observations in the dataset `model$X`.
- `diff.p.knowns` a vector with the calculated signed differential expression probabilities for the unlabeled observations in the dataset `model$knowns`. For `model$knowns=NULL` `diff.p.knowns` is also NULL(null).
- `diff.c` the index (or two indexes, in case of a three-component input model) of the identified differential component.

Author(s)
Przemyslaw Biecek

References

Examples
```r
data(Ste12)
X = Ste12Data[ match(names(Ste12Data), rownames(Ste12Beliefs), nomatch = 0) == 0 ]
knowns = Ste12Data[rownames(Ste12Beliefs)]
model = belief(X=X, knowns=knowns, B=Ste12Beliefs)
dep=DEprobs(model)
str(dep)
```

Description
The genotypes dataset describes 333 SNPs. Each SNP is characterized by the presence of one of its two possible alleles (or the presence of both of them). Therefore, the SNPs can be divided into three types. The first type corresponds to the SNPs with the first possible allele, the second type with the second allele, and the third with both alleles. The presence of the alleles is measured experimentally with fluorescence intensities. The dataset contains the intensities in the slots `X` and `knowns`.

15 SNPs in the dataset are given their correct type. These 'known' SNPs can be used as the input for the semi-, partially and fully supervised modeling methods. They were selected by taking at random five SNPs per each type. Their intensities are contained in the slot `knowns`. Their belief/plausibility values (given in the slot `B`) of the most probable type (the slot `labels`) are set to 0.95, and of the other two types are equal 0.025. The remaining 318 SNPs are kept unlabeled.

Usage
data(genotypes)
**Format**

- \( X \) : a matrix with 318 rows (unlabeled SNPs) and 2 columns (alleles) knowns : a matrix with 15 rows (known SNPs) and 2 columns (alleles)
- \( B \) : a matrix with 15 rows (known SNPs) and 3 columns (types)
- \( \text{labels} \) : a vector of length 15 (types of the known SNPs)

**Details**

The rows of both the slots \( X \) and knowns correspond to the SNPs. For each SNP, the values in the columns represent the intensities of the fluorescence signal corresponding to the alleles of the SNP. The slot \( B \) corresponds to the belief matrix while \( \text{labels} \) contains the true types for the labeled SNPs.

**References**


**Examples**

```r
library(bgmm)
data(gnotypes)

getModelStructure()
```

---

**Description**

This function creates an object which describes constraints over the model parameters.

**Usage**

```r
getModelStructure(mean = "D", between = "D", within = "D", cov = "D")
```

**Arguments**

- `mean` : Each argument is a single character, by default equal "D" (Different). If argument is set to "E" (or "0" for the argument `cov`) then the given parameter is constrained. By default all arguments are set to "D".
  - `mean="E"` forces equality of the means between the components,
  - `between="E"` forces equality of the covariance matrices among the components,
  - `within="E"` forces equality of variances within each covariance matrix i to some constant \( v_i \) and equality of covariances to some constant \( w_i \)
  - `cov="0"` forces equality of covariances within each covariance matrix i to "0",
**Value**

List of four elements specifying the constraints on 1) relations between the component means, 2) relations between the covariance matrices of the model components, 3) relations within each covariance matrix and 4) the covariances within each matrix. By default, the function returns an unconstrained structure.

**Author(s)**

Ewa Szczurek

**References**


**Examples**

```r
getModelStructure()
```

```r
getModelStructure(mean="E")
```

---

**init.model.params**  
**Initiation of model parameters**

**Description**

Methods for the initiation of model parameters for the EM algorithm. Two initiation procedures are implemented. The first procedure is available by setting the argument `method='knowns'`. It takes into account only labeled observations and is thus suitable for datasets with a high percentage of labeled cases. The second is available by setting `method='all'` and does not take the labeling into account.

**Usage**

```r
init.model.params(X = NULL, knowns = NULL, class = NULL,
                  k = length(unique(class)), method = "all", B = P, P = NULL)
```

**Arguments**

- `X`  
a data.frame with the unlabeled observations, its rows correspond to the observations while the columns correspond to variables/data dimensions.

- `knowns`  
a data.frame with the labeled observations, rows correspond to the observations while the columns correspond to variables/data dimensions.

- `B`  
a beliefs matrix with the distribution of beliefs for the labeled observations. If not specified and the argument `P` is given, the beliefs matrix is set to the value of `P`. 
\[
\text{init.model.params}
\]

\[P\]  a matrix of plausibilities, specified only for the labeled observations. The function assumes that the remaining observations are unlabeled and gives them uniformly distributed plausibilities by default. If not specified and the argument B is given, the plausibilities matrix is set to the value of B.

class  class is a vector of labels for the known observations. If not specified, it is derived from either the argument B or P with the use of the MAP rule.

\[k\]  the desired number of model components.

\[\text{method}\]  a method for parameter initialization, one of following c("knowns", "all"), see the section Details.

**Details**

For \[\text{method}='\text{knowns}'\], the initialization is based only on the labeled observations. i.e. those observations which have certain or probable components assigned. The initial model parameters for each component are estimated in one step from the observations that are assigned to this component (as in fully supervised learning).

If \[\text{method}='\text{all}\] (default), the initialization is based on all observations. In this case, to obtain the initial set of model components, we start by clustering the data using the k-means algorithm (repeated 10 times to get stable results). The only exception is for one dimensional data. In such a case the clusters are identified by dividing the data into k equal subsets of observations, where the subsets are separated by empirical quantiles \[c(1/2k, 3/2k, 5/2k, \ldots, (2k-1)/2k)\]. After this initial clustering each cluster is linked to one model component and initial values for the model parameters are derived from the clustered observations.

For the partially and semi-supervised methods, correspondence of labels from the initial clustering algorithm and labels for the observations in the knowns dataset rises a technical problem. The cluster corresponding to component \[y\] should be as close as possible to the set of labeled observations with label \[y\].

Note that for the unsupervised modeling this problem is irrelevant and any cluster may be used to initialize any component.

To match the cluster labels with the labels of model components a greedy heuristic is used. The heuristic calculates weighted distances between all possible pairs of cluster centers and sets of observations grouped by their labels. In each step, the pair with a minimal distance is chosen (the pair: a group of observations with a common label and a cluster, for which the center of the group is the closest to the center of the cluster). For the chosen pair, the cluster is labeled with the same label as the group of observations. Then, this pair is removed and the heuristic repeats for the reduced set of pairs.

**Value**

A list with the following elements:

\[\pi\]  a vector of length k with the initial values for the mixing proportions.

\[\mu\]  a matrix with the means' vectors with the initial values for k components.

\[\text{cvar}\]  a three-dimensional matrix with the covariance matrices with the initial values for k components.
miRNA

Author(s)
Przemyslaw Biecek

References

Examples

data(genotypes)
initNparams = init.model.params(X=genotypes$X, knowns=genotypes$knowns, class = genotypes$labels)
str(initNparams)

miRNA  miRNA transfection data for miR1 and miR124 target genes

Description
miRNA transfection data (Lim et al., 2005) and knowledge from computational miRNA target predictions.

Usage
data(miRNA)

Format
miR1Data vector: 117, miR124Data vector: 117, miRNABeliefs matrix of example certainty: 26 x 2, miRNAClass vector: 117

Details
miR1Data Log2 expression ratios of miR1 transfection versus wild type, for 117 genes. miR124Data Log2 expression ratios of miR124 transfection versus wild type, for 117 genes. miRNABeliefs Gives the certainty (belief/plausibility) for each out of 26 example miRNA targets to belong to their cluster. miRNAClass Gives the true cluster for each gene. Cluster 1 corresponds to the experimentally verified targets of miR1. Cluster 2 corresponds to the targets of miR124.

Author(s)
Ewa Szczurek

References
See Also

Ste12.CellCycle

Examples

library(bgmm)
data(miRNA)
print(miR1Data)
print(miR24Data)
print(miRNABeliefs)
print(miRNAClass)

mModel Fitting Gaussian mixture model

Description

These functions fit different variants of Gaussian mixture models. These variants differ in the fraction of knowledge utilized into the fitting procedure.

Usage

belief(X, knowns, B = NULL, k = ifelse(!is.null(B), ncol(B),
  ifelse(is.null(P), ncol(P), length(unique(class)))), P = NULL,
  class = map(B), init.params = init.model.params(X, knowns,
  B = B, P = P, class = class, k = k), model.structure = getModelStructure(),
  stop.likelihood.change = 10^-5, stop.max.nsteps = 100, trace = FALSE,
  b.min = 0.025,
  all.possible.permutations=FALSE)

soft(X, knowns, P = NULL, k = ifelse(!is.null(P), ncol(P),
  ifelse(is.null(B), ncol(B), length(unique(class)))), B = NULL,
  class = NULL, init.params = init.model.params(X, knowns,
  class = class, B = P, k = k),
  model.structure = getModelStructure(), stop.likelihood.change = 10^-5,
  stop.max.nsteps = 100, trace = FALSE, b.min = 0.025,
  all.possible.permutations=FALSE)

semisupervised(X, knowns, class = NULL, k = ifelse(!is.null(class),
  length(unique(class)), ifelse(!is.null(B), ncol(B), ncol(P)))),
  B = NULL, P = NULL, ...
  all.possible.permutations=FALSE)

supervised(knowns, class = NULL, k = length(unique(class)), B = NULL, P = NULL,
  model.structure = getModelStructure(), ...)

unsupervised(X, k, init.params=init.model.params(X, knowns=NULL, k=k),
model.structure=getModelStructure(), stop.likelihood.change=10^-5, stop.max.nsteps=100, trace=FALSE)

Arguments

X  a data.frame with the unlabeled observations. The rows correspond to the observations while the columns to variables/dimensions of the data.

knowns  a data.frame with the labeled observations. The rows correspond to the observations while the columns to variables/dimensions of the data.

B  a beliefs matrix which specifies the distribution of beliefs for the labeled observations. The number of rows in B should equal the number of rows in the data.frame knowns. It is assumed that both the observations in B and in knowns are given in the same order. Columns correspond to the model components. If matrix B is provided, the number of columns has to be less or equal k. Internally, the matrix B is completed to k columns.

P  a matrix of plausibilities, i.e., weights of the prior probabilities for the labeled observations. If matrix P is provided, the number of columns has to be less or equal k. The same conditions as for B apply.

class  a vector of classes/labels for the labeled observations. The number of its unique values has to be less or equal k.

k  a number of components, by default equal to the number of columns of B.

init.params  initial values for the estimates of the model parameters (means, variances and mixing proportions), by default derived with the use of the init.model.params function.

stop.likelihood.change, stop.max.nsteps  the parameters for the EM algorithms defining the stop criteria, i.e., the minimum required improvement of loglikelihood and the maximum number of steps.

trace  if trace=TRUE the loglikelihoods for every step of EM algorithm are printed out.

model.structure  an object returned by the getModelStructure function, which specifies constraints for the model to be fitted.

b.min  this argument is passed to the init.model.params function.

...  these arguments will be passed to the init.model.params function.

all.possible.permutations  If equal TRUE, all possible initial parameters’ permutations of components are considered. Since there is kList! permutations, model fitting is repeated kList! times. As a result, only the model with the highest likelihood is returned.

Details

In the belief() function, if the argument B is not provided, it is by default initialized from the argument P. If the argument P is not provided, B is derived from the class argument with the use of the function get.simple.beliefs() which assigns 1/(k-1)*b.min to the component given by class and b.min to all remaining components.
In the `soft()` function, if the argument `P` is not provided, it is by default initialized from the argument `B`. If the argument `B` is not provided, `P` is derived from the `class` argument as in the `belief()` function.

In the `supervised()` function, if the argument `class` is not provided, it is by default initialized from argument `B` or `P`, taking the label of each observation as its most believed or plausible component (by the MAP rule).

The number of columns in the beliefs matrix `B` or in the matrix of plausibilities `P` may be smaller than the number of model components defined by the argument `k`. Such situation corresponds to the scenario when the user does not know any examples for some component. In other words, this component is not used as a label for any observation, and thus can be omitted from the beliefs matrix. An equivalent would be to include a column for this component and fill it with beliefs/plausibilities equal 0.

Slots in the returned object are listed in section Value. The returned object differs slightly with respect to the used function. Namely, the `belief()` function returns an object with the slot `b`. The function `soft()` returns an object with a slot `p`, while the functions `supervised()` and `semisupervised()` return objects with a slot `class` instead.

The object returned by the function `supervised()` does not have the slot `X`.

**Value**

An object of the class `mModel`, with the following slots:

- `pi`: a vector with the fitted mixing proportions
- `mu`: a matrix with the means’ vectors, fitted for all components
- `cvar`: a three-dimensional matrix with the covariance matrices, fitted for all components
- `X`: the unlabeled observations
- `knowns`: the labeled observations
- `B`: the beliefs matrix
- `n`: the number of all observations
- `m`: the number of the unlabeled observations
- `k`: the number of fitted model components
- `d`: the data dimension
- `likelihood`: the log-likelihood of the fitted model
- `n.steps`: the number of steps performed by the EM algorithm
- `model.structure`: the set of constraints kept during the fitting process.

**Author(s)**

Przemyslaw Biecek

**References**

Examples

data(genotypes)

modelSupervised = supervised(knowns=genotypes$knowns, class=genotypes$labels)
plot(modelSupervised)

modelSemiSupervised = semisupervised(X=genotypes$X, knowns=genotypes$knowns, class = genotypes$labels)
plot(modelSemiSupervised)

modelBelief = belief(X=genotypes$X, knowns=genotypes$knowns, B=genotypes$B)
plot(modelBelief)

modelSoft = soft(X=genotypes$X, knowns=genotypes$knowns, P=genotypes$P)
plot(modelSoft)

modelUnSupervised = unsupervised(X=genotypes$X, k=3)
plot(modelUnSupervised)

mModelList

Fitting Gaussian mixture model or collection of models

Description

These functions fit collection of models of one particular variant/class. Models to be fitted may differ in the requested number of Gaussian components or in the requested model structure.

Usage

mModelList(X, knowns, B = NULL, P = NULL, class = NULL, kList = ncol(B),
           init.params = NULL, stop.likelihood.change = 10^-5, stop.max.nsteps = 100,
           trace = FALSE, mean = c("D", "E"), between = c("D", "E"),
           within = c("D", "E"), cov = c("D", "0"), funct = belief, all.possible.permutations = FALSE)

beliefList(..., funct=belief)

softList(..., funct=soft)

semisupervisedList(..., funct=semisupervised)

unsupervisedList(X, kList = 2, ...)
Arguments

X
a data.frame with the unlabeled observations. The rows correspond to the observations while the columns to variables/dimensions of the data.

knowns
a data.frame with the labeled observations. The rows correspond to the observations while the columns to variables/dimensions of the data.

B
a beliefs matrix which specifies the distribution of beliefs for the labeled observations. The number of rows in B should equal the number of rows in the data.frame knowns. It is assumed that both the observations in B and in knowns are given in the same order. Columns correspond to the model components. If matrix B is provided, the number of columns has to be less or equal k. Internally, the matrix B is completed to k columns.

P
a matrix of plausibilities, i.e., weights of the prior probabilities for the labeled observations. If matrix P is provided, the number of columns has to be less or equal k. The same conditions as for B apply.

class
a vector of classes/labels for the labeled observations. The number of its unique values has to be less or equal min(kList).

kList
a vector or a list with numbers of Gaussian components to fit. By default it is one number equal to the number of columns of B.

init.params
initial values for the estimates of the model parameters (means, variances and mixing proportions). The initial parameters are internally passed to the funct function.

stop.likelihood.change, stop.max.nsteps, trace
the parameters for the EM algorithm. Internally, these parameters are passed to the funct function.

mean, between, within, cov
four vectors which define the model structures for models to be fitted. For example, if mean="E", only models with constrained means are considered (means of Gaussian components are forced to be equal). On the other hand if mean=c("E", "D"), both models with constrained means and models without constraint on the means are fitted.

funct
a function which fits a variant of Gaussian mixture model, one of the: belief, soft, semisupervised or unsupervised functions.

... arguments that are passed to function funct.

all.possible.permutations
If equal TRUE, all possible initial parameters’ permutations of components are considered. Since there is kList! permutations, model fitting is repeated kList! times. As a result only the model with the highest likelihood is returned.

Details

Arguments kList, as well as mean, between, within, and cov define the list of models to be fitted. All combinations of specified model sizes and model structures are considered. List of fitted models is returned as a result.

The argument funct defines which variant of Gaussian mixture models should be used for model fitting. One can use the wrappers beliefList(), softList(), semisupervisedList(), unsupervisedList() which call the mModelList() function and have a prespecified argument funct.
Value
An object of the class mModelList, with the following slots:

- **models**: a list of models, each of the class mModel
- **loglikelihoods**: a vector with log likelihoods of the models from list models
- **names**: a vector with names of the models from list models
- **params**: a vector with the number of parameters of models from list models
- **klist**: equals the input argument klist

Author(s)
Przemyslaw Biecek

References

See Also
- mModel, getModelStructure

Examples
simulated = simulateData(d=2, k=3, n=100, m=60, cov="0", within="E", n.labels=2)

models1 = Modellist(X=simulated$X, knowns=simulated$knowns, B=simulated$B,
                      klist=3:4, mean=c("D","E"), between="D", within="D",
                      cov="0", funct=belief)

plot(models1)
plotGIC(models1, penalty="BIC")

# Do not run
# It could take more than one minute
# simulated = simulateData(d=2, k=3, n=300, m=60, cov="0", within="E", n.labels=2)
# models1 = Modellist(X=simulated$X, knowns=simulated$knowns, B=simulated$B,
#                      klist=3, mean=c("D","E"), between=c("D","E"), within=c("D","E"),
#                      cov=c("D","0"), funct=belief)
# plot(models1)
# plotGIC(models1, penalty="BIC")
# models2 = beliefList(X=simulated$X, knowns=simulated$knowns, B=simulated$B,
#                       klist=2:7, mean="D", between="D", within="E", cov="0")
# plot(models2)
# plotGIC(models2, penalty="BIC")
# models3 = beliefList(X=simulated$X, knowns=simulated$knowns, B=simulated$B,
#                       klist=2:7, mean="D")
# plotGIC(models3, penalty="BIC")
plot.mModel

Plotting a graphical visualization of a Gaussian model or a list of models

Description

The generic function plot is used to visualize the data set and Gaussian model components fitted to this data. On the resulting plot the observations without labels are presented with black points, whereas the labeled observations are marked by different colors and different symbols. The fitted Gaussian components are represented by ellipses into the two-dimensional case and by densities in the one dimensional case. If data has more than two dimensions thus graphs are presented on the subspace generated by first two PCA components. Note that the estimation is done in higher dimension and the reduction to 2D is done only for illustration. That gives different results than data reduction prior to modeling process.

Usage

```
## S3 method for class 'mModel'
plot(x, ...)  
```

Arguments

- `x` an object of the class `mModel`.
- `...` graphical arguments that are passed to the underlying `plot()` function.

Details

For one dimensional data the width of the density corresponds to standard deviation of the fitted Gaussian component. Fitted means are marked by vertical dashed lines.

For two dimensional data ellipses represents covariances for the corresponding model components.

For more dimensional points and ellipses are projected into 2D subspace spanned by first two PCA components.

Author(s)

Przemyslaw Biecek

References

plot.mModelList

Plotting a graphical visualization of a model or a list of models

Description

The function `plot.mModelList()` creates a grid of panels and then plots a set of input fitted models in the consecutive panels. The `plot.mModel()` function is used to plot each single model.

Usage

```r
## S3 method for class 'mModelList'
plot(x, ...)  
```

Arguments

- `x` an object of the class `mModelList`.
- `...` graphical arguments that are passed to underlying `plot()` function.

Details

The argument `x` is a list of models. If these models differ both by component numbers and by the model structures, in the resulting grid of panels columns correspond to the different model structures while rows correspond to the different component numbers.

If considered models differ only by component numbers or only by the model structures, the grid of panels is as close to square as possible and consecutive panels contain consecutive models from the list of models `x`.

Examples

```r
data(genotypes)
modelSupervised = supervised(knowns=genotypes$knowns, class=genotypes$labels)
plot(modelSupervised)

# semi-supervised modeling
modelSemiSupervised = semisupervised(X=genotypes$X, knowns=genotypes$knowns, class = genotypes$labels)
plot(modelSemiSupervised)

# belief-based modeling
modelBelief = belief(X=genotypes$X, knowns=genotypes$knowns, B=genotypes$B)
plot(modelBelief)

# soft-label modeling
modelSoft = soft(X=genotypes$X, knowns=genotypes$knowns, P=genotypes$P)
plot(modelSoft)

# unsupervised modeling
modelUnSupervised = unsupervised(X=genotypes$X, k=3)
plot(modelUnSupervised)
```
plotGIC

Description

The function plotGIC() plots the GIC scores for an input collection of models. The function getGIC() extracts GIC for given model and penalty function. The function getDF() extracts the number of degree of freedom for model parameters.
Usage

plotGIC(models, penalty = 2, plot.it = TRUE, ...)  

getGIC(model, p = 2, whichobs="unlabeled")  

getDF(model)

Arguments

models an object of the class mModelList or a matrix of GIC scores.  
model an object of the class mModel.  
penalty a penalty for the GIC criteria. This parameter can be a single number or a string,  
on of the "BIC", "AIC", "AIC3", "AIC4", "AICe", "AICu", "CAIC", "BIC",  
"MDL", "CLC", "ICL-BIC", "AWE".  
p same as penalty,  
whichobs one of "unlabeled", "labeled", "all". This parameter specify which observations  
should be used in the likelihood and gic score calculation,  
plot.it a logical value, if TRUE then the chart with the GIC scores will be plotted.  
... other arguments that will be passed to the getGIC function.

Details

The function plotGIC() calculates the GIC scores for each model from the models list and, given  
plot.it=TRUE, plots a dotchart with the calculated GIC scores.  

As a result the function plotGIC() returns a matrix with the calculated GIC scores. This matrix or  
its submatrix can be used in next call of the plotGIC() function as models argument. The columns  
of the matrix correspond to different component numbers of the models, while the rows correspond  
to their structures. The structures are coded with four-letter strings. The letters refer, in order from  
left to right: first, the relation between the means' vectors of the components, which can either  
be equal (letter "E") or unconstrained ("D"). Second, the relation between covariance matrices,  
which can all either be equal ("E"), or unconstrained ("D"). Third, the relation between the data  
vector components (corresponding to data dimensions) within each covariance matrix, i.e. each  
covariance matrix can either have all variances equal to some constant and all covariances equal  
to some constant ("E") or can be unconstrained ("D"). Fourth, the covariances in each covariance  
matrix, which can either all be forced to equal 0 ("0") or be unconstrained ("D").  

The best model, i.e. model with the smallest GIC score is marked with a star on the plotted chart.

Value

The matrix with GIC scores calculated for the list of models specified by the models argument.

Author(s)

Przemyslaw Biecek
predict.mModel

Predictions for fitted Gaussian component model

Description

For every row in the matrix X the posterior probability of belonging to class i is calculated.

Usage

```r
# S3 method for class 'mModel'
predict(object, X, knowns = NULL, B = NULL, P = NULL, ...)
```

Arguments

- `object`: an object of the class `mModel`
- `X`: a matrix or `data.frame` in which number of columns is equal to `object$d`
- `knowns`: a `data.frame` or `matrix` with the labeled observations. If the argument `knowns` is specified then either `B` or `P` need to be specified.
- `P`: a matrix with plausibilities for `knowns`. 

References

B a matrix with beliefs for object knowns.

... all other arguments will be neglected.

Details

The matrix $t_{ij}$ of posterior probabilities is calculated as normalized products of priors $p_i$'s and density of model components in values specified by rows of the matrix $X$.

If arguments `knowns` and `B` are specified then the priors's for objects in `knowns` are replaced by belief matrix $B$. If arguments `knowns` and `P` are specified then the priors's for objects in `knowns` are multiplied by plausibility matrix $P$.

Value

An list with the following elements:

- `tij.X`, `tij.knowns` the matrix $t_{ij}X$ is a matrix with number of rows equal to number of rows in the matrix $X$ and the number of columns equal to the number of components in model defined by argument `object`. Values in this matrix are posterior probabilities that observation $i$ belongs to component $j$. The slot `tij.knowns` is equal to NULL if neither `B` nor `P` are specified, otherwise it is a matrix with number of rows equal to number of rows in the matrix `knowns` and contains posterior probabilities for observations with specified belief or plausibilities matrix.

- `class.X`, `class.knowns` vectors of labels/classes obtained with the MAP rule. The vector `class.X` corresponds to observations in $X$ while the vector `class.knowns` corresponds to observations in `knowns`.

Author(s)


References

http://bgmm.molgen.mpg.de

See Also

- `belief`

Examples

data(genotypes)

modelSoft = soft(X=genotypes$X, knowns=genotypes$knowns, P=genotypes$B)

preds = predict(modelSoft, X = genotypes$X)
str(preds)
**Description**

The function `simulateData` generates an artificial dataset from a mixture of Gaussian components with a given set of parameters.

**Usage**

```r
simulateData(d = 2, k = 4, n = 100, m = 10, mu = NULL, cvar = NULL,
             s.pi = rep(1/k, k), b.min = 0.02, mean = "D", between = "D",
             within = "D", cov = "D", n.labels = k)
```

**Arguments**

- **d** — the dimension of the data set,
- **k** — the number of the model components,
- **n** — the total number of observations, both labeled and unlabeled,
- **mu** — a matrix with `k` rows and `d` columns, which defines the means' vectors for the corresponding model components. If not specified, by default its values are generated from a normal distribution N(0,49),
- **cvar** — a three-dimensional array with the dimensions `(k, d, d)`. If not specified, each covariance matrix is generated in three steps: first, `2*d` samples from a `d`-dimensional normal distribution N(0,Id) are generated. Next, a covariance matrix `d x d` for these samples is calculated. Finally, the resulting sample covariance matrix is scaled by a factor generated from an exponential distribution Exp(1),
- **s.pi** — a vector of `k` probabilities, i.e. the mixing proportions of the model. The mixing proportions specify a multinomial distribution over the components, from which the numbers of observations in each cluster are generated. By default a uniform distribution is used.
- **mean**, **between**, **within**, **cov** — constraints on the model structure. By default all are equal to "D". If other values are set, the parameters `mu` and `cvar` are adjusted to match the specified constraints,
- **m** — the number of the observations, for which the beliefs are to be calculated,
- **b.min** — the belief that an observation does not belong to a component. Formally, the belief `b_{ij}` for the observation `i` to belong to component `j` is equal `b.min` if `i` is not generated from component `j`. Thus, the belief that `i` belongs to its true component is set to `1 - b.min*(n.labels-1)`, and `b.min` is constrained that `b.min < 1/n.labels`. By default `b.min`=0.02,
- **n.labels** — the number of components used as labels, defining the number of columns in the resulting beliefs matrix. By default `n.labels` equals `k`, but the user can specify a smaller number. Using this argument the user can define a scenario in which the data are generated from a mixture of three components, but only two of them are used as labels in the beliefs matrix (applied in the example below).
Value

An list with the following elements:

- **X** the matrix of size n-m rows and d columns with generated values of unlabeled observations,
- **knowns** the matrix of size m rows and d columns with generated values of labeled observations,
- **B** the belief matrix of the size m rows and k columns derived for knowns matrix,
- **model.params** the list of model parameters,
- **Ytrue** indexes of the true Gaussian components from which each observation was generated.

Author(s)

Przemyslaw Biecek

References


Examples

```r
simulated = simulateData(d=2, k=3, n=300, m=60, cov="E", within="E", n.labels=2)
model = belief(X = simulated$X, knowns = simulated$knowns, B=simulated$B)
plot(model)
```

```r
simulated = simulateData(d=1, k=2, n=300, m=60, n.labels=2)
model = belief(X = simulated$X, knowns = simulated$knowns, B=simulated$B)
plot(model)
```

---

Ste12

*Ste12 knockout data under pheromone treatment versus wild type; Examples of Ste12 targets; Binding p-values of Ste12 to those targets.*

Description

Ste12 knockout expression data (Roberts et al., 2002) and knowledge from a Ste12 binding experiment (Harbison et al., 2004) used for identifying Ste12 target genes under pheromone treatment.

Usage

data(Ste12)

Format

Ste12Data vector: 601 Ste12Beliefs matrix of example certainty: 42 x 2 Ste12Binding vector: 42
Details

Ste12Data Log2 expression ratios of Ste12 knockout versus wild type, both under 50nM alpha-factor treatment for 30min. This data is for 601 genes that had more than 1.5 fold change in expression after pheromone treatment versus wild type. Ste12Beliefs: Gives the certainty (belief/plausibility) for each out of 42 example Ste12 targets to belong to their cluster. Ste12Beliefs: Gives the certainty (belief/plausibility) for each out of 42 example Ste12 targets to belong to their cluster. The 42 examples were chosen to meet two criteria: (1) Had a binding p-value <0.0001 (see Ste12Binding), and (2) Had a 2-fold change in response to pheromone treatment (versus wild-type) Ste12Binding: Gives the binding p-value for each example Ste12 target (see Ste12Belief).

Author(s)

Ewa Szczurek

References


See Also

mirna, CellCycle

Examples

data("Ste12")
print(Ste12Data)
print(Ste12Beliefs)
print(Ste12Binding)
Supplementary functions

Usage

```r
## S3 method for class 'numeric'
determinant(x, logarithm = TRUE, ...)

map(B)

loglikelihood.mModel(model, X)
```

Arguments

- `x` a single number.
- `X` a data.frame with the unlabeled observations, the rows correspond to the observations and the columns to the dimensions of the data.
- `B` a beliefs matrix with the distribution of beliefs for the labeled observations.
- `model` an object of the class mModel.
- `logarithm, ...` these arguments are ignored.

Author(s)

Przemyslaw Biecek

References


Examples

```r
data(genotypes)

map(genotypes$B)
```
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