Package ‘mcclust’

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Description Implements methods for processing a sample of (hard) clusterings, e.g. the MCMC output of a Bayesian clustering model. Among them are methods that find a single best clustering to represent the sample, which are based on the posterior similarity matrix or a relabelling algorithm.
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R topics documented:

mcclust-package .................................................. 2
arandi .......................................................... 3
cls.draw1.5 ..................................................... 4
cls.draw2 ....................................................... 5
cltosim .......................................................... 5
comp.psm ....................................................... 6
maxpear .......................................................... 7
medv ............................................................. 8
minbinder ....................................................... 9
norm.label ..................................................... 12
relabel .......................................................... 12
vi.dist .......................................................... 14
mcclust-package

Ysim1.5 ................................................................. 15
Ysim2 ................................................................. 16

Index 17

mcclust-package  Process MCMC Sample of Clusterings.

Description

Implements methods for processing a sample of (hard) clusterings, e.g. the MCMC output of a Bayesian clustering model. Among them are methods that find a single best clustering to represent the sample, which are based on the posterior similarity matrix or a relabelling algorithm.

Details

Package: mcclust
Type: Package
Version: 1.0
Date: 2009-03-12
License: GPL (>= 2)
LazyLoad: yes

Most important functions:

comp.psm for computing posterior similarity matrix (PSM). Based on the PSM maxpear and minbinder provide several optimization methods to find a clustering with maximal posterior expected adjusted Rand index with the true clustering or one that minimizes the posterior expectation of a loss function by Binder (1978). minbinder provides the optimization algorithm of Lau and Green.

relabel contains the relabelling algorithm of Stephens (2000).

arandi and vi.dist compute distance functions for clusterings, the (adjusted) Rand index and the entropy-based variation of information distance.

Author(s)

Arno Fritsch
Maintainer: Arno Fritsch <arno.fritsch@tu-dortmund.de>

References


**Examples**

data(cls.draw2)
# sample of 500 clusterings from a Bayesian cluster model
tru.class <- rep(1:8,each=50)
# the true grouping of the observations
psm2 <- comp.psm(cls.draw2)
# posterior similarity matrix

# optimize criteria based on PSM
mbind2 <- minbinder(psm2)
mpear2 <- maxpear(psm2)

# Relabelling
k <- apply(cls.draw2,1, function(cl) length(table(cl)))
max.k <- as.numeric(names(table(k))[which.max(table(k))])
relab2 <- relabel(cls.draw2[k==max.k,])

# compare clusterings found by different methods with true grouping
arandi(mpear2$cl, tru.class)
arandi(mbind2$cl, tru.class)
arandi(relab2$cl, tru.class)

---

**arandi** *(Adjusted) Rand Index for Clusterings*

**Description**

Computes the adjusted or unadjusted Rand index between two clusterings/partitions of the same objects.

**Usage**

arandi(cl1, cl2, adjust = TRUE)

**Arguments**

c1l,c2l vectors of cluster memberships (need to have the same lengths).
adjust logical. Should index be adjusted? Defaults to TRUE.
Details

The Rand index is based on how often the two clusterings agree in the treatment of pairs of observations, where agreement means that two observations are in/not in the same cluster in both clusterings.

The adjusted Rand index adjusts for the expected number of chance agreements.

Formulas of Hubert and Arabie (1985) are used for the computation.

Author(s)

Arno Fritsch, <arno.fritsch@tu-dortmund.de>

References


See Also

*vi.dist*

Examples

```r
cl1 <- sample(1:3,10,replace=TRUE)
c12 <- c(cl1[1:5], sample(1:3,5,replace=TRUE))
arandi(cl1,cl2)
arandi(cl1,cl2,adjust=FALSE)
```

---

**cls.draw1.5**  
*Sample of Clusterings from Posterior Distribution of Bayesian Cluster Model*

Description

Output of a Dirichlet process mixture model with normal components fitted to the data set Ysim1.5. True clusters are given by `rep(1:8,each =50)`.

Usage

`data(cls.draw1.5)`

Format

matrix with 500 rows and 400 columns. Each row contains a clustering of the 400 observations.

Source

Sample of Clusterings from Posterior Distribution of Bayesian Cluster Model

Description

Output of a Dirichlet process mixture model with normal components fitted to the data set Ysim2. True clusters are given by rep(1:8,each =50).

Usage

data(cls.draw2)

Format

matrix with 500 rows and 400 columns. Each row contains a clustering of the 400 observations.

Source


Compute Similarity Matrix for a Clustering and vice versa

Description

A similarity matrix is a symmetric matrix whose entry \([i, j]\) is 1 if observation \(i\) and \(j\) are in the same cluster and 0 otherwise.

Usage

cltoSim(cl)
Simtocl(Sim)

Arguments

- **cl**: vector of cluster memberships
- **Sim**: similarity matrix

Warning

Simtocl does not check whether Sim is a valid similarity matrix, e.g. that Sim\([i, j]\)==1 if Sim\([i, k]\]==1 and Sim\([j, k]\]==1.
**Author(s)**

Arno Fritsch, <arno.fritsch@tu-dortmund.de>

**See Also**

comp.psm for an average similarity matrix.

**Examples**

```r
cl <- c(3,3,1,2,2)
(Sim <- cltoSim(cl))
Simtocl(Sim)

# not a valid similarity matrix
(Sim2 <- matrix(c(1,0,1,0,1,1,1,1,1), ncol=3))
Simtocl(Sim2) # no warning
```

---

**comp.psm**

*Estimate Posterior Similarity Matrix*

**Description**

For a sample of clusterings of the same objects the proportion of clusterings in which observation $i$ and $j$ are together in a cluster is computed and a matrix containing all proportions is given out.

**Usage**

`comp.psm(cls)`

**Arguments**

- `cls` a matrix in which every row corresponds to a clustering of the `ncol(cls)` objects

**Details**

In Bayesian cluster analysis the posterior similarity matrix is a matrix whose entry $[i,j]$ contains the posterior probability that observation $i$ and $j$ are together in a cluster. It is estimated by the proportion of a posteriori clusterings in which $i$ and $j$ cluster together.

**Value**

a symmetric `ncol(cls)*ncol(cls)` matrix

**Author(s)**

Arno Fritsch, <arno.fritsch@tu-dortmund.de>
maxpear

See Also

cltoSim

Examples

```r
(cls <- rbind(c(1,1,2,2),c(1,1,2,2),c(1,2,2,2),c(2,2,1,1)))
comp.psm(cls)
```

maxpear

Maximize/Compute Posterior Expected Adjusted Rand Index

Description

Based on a posterior similarity matrix of a sample of clusterings maxpear finds the clustering that maximizes the posterior expected Rand adjusted index (PEAR) with the true clustering, while pear computes PEAR for several provided clusterings.

Usage

```r
maxpear(psm, cls.draw = NULL, method = c("avg", "comp", "draws", "all"), max.k = NULL)
pear(cls, psm)
```

Arguments

- `psm` a posterior similarity matrix, usually obtained from a call to `comp.psm`.
- `cls, cls.draw` a matrix in which every row corresponds to a clustering of the `ncol(cls)` objects. `cls.draw` refers to the clusterings that have been used to compute `psm`, `cls.draw` has to be provided if `method="draw"` or "all".
- `method` the maximization method used. Should be one of "avg", "comp", "draws" or "all". The default is "avg".
- `max.k` integer, if `method="avg"` or "comp" the maximum number of clusters up to which the hierarchical clustering is cut. Defaults to `ceiling(nrow(psm)/8)`.

Details

For `method="avg"` and "comp" 1-psm is used as a distance matrix for hierarchical clustering with average/complete linkage. The hierarchical clustering is cut for the cluster sizes 1:max.k and PEAR computed for these clusterings.

Method "draws" simply computes PEAR for each row of `cls.draw` and takes the maximum.

If `method="all"` all maximization methods are applied.
**Value**

- **cl**: clustering with maximal value of PEAR. If `method="all"` a matrix containing the clustering with the highest value of PEAR over all methods in the first row and the clusterings of the individual methods in the next rows.

- **value**: value of PEAR. A vector corresponding to the rows of `cl` if `method="all"`.

- **method**: the maximization method used.

**Author(s)**

Arno Fritsch, <arno.fritsch@tu-dortmund.de>

**References**


**See Also**

- `comp.psm` for computing posterior similarity matrix, `minbinder`, `medv`, `relabel` for other possibilities for processing a sample of clusterings.

**Examples**

```r
data(cls.draw1.5)
# sample of 500 clusterings from a Bayesian cluster model
tru.class <- rep(1:8,each=50)
# the true grouping of the observations
psm1.5 <- comp.psm(cls.draw1.5)
mpear1.5 <- maxpear(psm1.5)
table(mpear1.5$cl, tru.class)

# Does hierarchical clustering with Ward's method lead
# to a better value of PEAR?
hclust.ward <- hclust(as.dist(1-psm1.5), method="ward")
cls.ward <- t(apply(matrix(1:20),1, function(k) cutree(hclust.ward,k=k)))
ward1.5 <- pear(cls.ward, psm1.5)
max(ward1.5) > mpear1.5$value
```

**Description**

Based on a posterior similarity matrix of a sample of clusterings `medv` obtains a clustering by using `1-psm` as distance matrix for hierarchical clustering with complete linkage. The dendrogram is cut at a value `h` close to 1.
**minbinder**

Usage

```r
medv(psm, h=0.99)
```

Arguments

- `psm`: a posterior similarity matrix, usually obtained from a call to `comp.psm`.
- `h`: The height at which the dendrogram is cut.

Value

vector of cluster memberships.

Author(s)

Arno Fritsch, <arno.fritsch@tu-dortmund.de>

References


See Also

- `comp.psm` for computing posterior similarity matrix, `maxpear`, `minbinder`, `relabel` for other possibilities for processing a sample of clusterings.

Examples

```r
data(cls.draw1.5)
# sample of 500 clusterings from a Bayesian cluster model
tru.class <- rep(1:8,each=50)
# the true grouping of the observations
psm1.5 <- comp.psm(cls.draw1.5)
medv1.5 <- medv(psm1.5)
table(medv1.5, tru.class)
```

---

**minbinder**

*Minimize/Compute Posterior Expectation of Binders Loss Function*

Description

Based on a posterior similarity matrix of a sample of clusterings `minbinder` finds the clustering that minimizes the posterior expectation of Binders loss function, while `binder` computes the posterior expected loss for several provided clusterings.
Usage

\[ \text{minbinder}(\text{psm}, \text{cls.draw} = \text{NULL}, \text{method} = \text{c("avg", "comp", "draws", "laugreen","all"); max.k = NULL, include.lg = FALSE, start.cl = NULL, tol = 0.001}) \]

\[ \text{binder}(\text{cls},\text{psm}) \]

\[ \text{laugreen}(\text{psm}, \text{start.cl}, \text{tol}=0.001) \]

Arguments

- **psm**: a posterior similarity matrix, usually obtained from a call to `comp.psm`.
- **cls, cls.draw**: a matrix in which every row corresponds to a clustering of the `ncol(cls)` objects. `cls.draw` refers to the clusterings that have been used to compute `psm`, `cls.draw` has to be provided if `method="draw"` or "all".
- **method**: the maximization method used. Should be one of "avg", "comp", "draws", "laugreen" or "all". The default is "avg".
- **max.k**: integer, if `method="avg"` or "comp" the maximum number of clusters up to which the hierarchical clustering is cut. Defaults to `ceiling(nrow(psm)/4)`.
- **include.lg**: logical, should method "laugreen" be included when `method="all"`? Defaults to FALSE.
- **start.cl**: clustering used as starting point for `method="laugreen"`. If `NULL start.cl=1:nrow(psm)` is used.
- **tol**: convergence tolerance for `method="laugreen"`.

Details

The posterior expected loss is the sum of the absolute differences of the indicator function of observation \( i \) and \( j \) clustering together and the posterior probability that they are in one cluster.

For `method="avg"` and "comp" \(1-psm\) is used as a distance matrix for hierarchical clustering with average/complete linkage. The hierarchical clustering is cut for the cluster sizes \(1:max.k\) and the posterior expected loss is computed for these clusterings.

Method "draws" simply computes the posterior expected loss for each row of `cls.draw` and takes the minimum.

Method "laugreen" implements the algorithm of Lau and Green (2007), which is based on binary integer programming. Since the method can take some time to converge it is only used if explicitly demanded with `method="laugreen"` or `method="all" and include.lg=TRUE`. If `method="all"` all minimization methods except "laugreen" are applied.

Value

- **cl**: clustering with minimal value of expected loss. If `method="all"` a matrix containing the clustering with the smallest value of the expected loss over all methods in the first row and the clusterings of the individual methods in the next rows.
- **value**: value of posterior expected loss. A vector corresponding to the rows of `cl` if `method="all"`. 
method the maximization method used.
iter.lg if method="laugreen" the number of iterations the method needed to converge.

Author(s)
Arno Fritsch, <arno.fritsch@tu-dortmund.de>

References

See Also
*comp.psm* for computing posterior similarity matrix, *maxpear, medv, relabel* for other possibilities for processing a sample of clusterings. *lp* for the linear programming.

Examples
```r
data(cls.draw2)
# sample of 500 clusterings from a Bayesian cluster model
tru.class <- rep(1:8,each=50)
# the true grouping of the observations
psm2 <- comp.psm(cls.draw2)
mbind2 <- minbinder(psm2)
table(mbind2$cl, tru.class)

# Does hierachical clustering with Ward's method lead
# to a lower value of Binders loss?
hclust.ward <- hclust(as.dist(1-psm2), method="ward")
cls.ward <- t(apply(matrix(1:20),1, function(k) cutree(hclust.ward,k=k)))
ward2 <- binder(cls.ward, psm2)
min(ward2) < mbind2$value

# Method laugreen is applied to 40 randomly selected observations
ind <- sample(1:400, 40)
mbind.lg <- minbinder(psm2[ind, ind],cls.draw2[,ind], method="all",
                      include.lg=TRUE)
mbind.lg$value
```
norm.label  

Norm Labelling of a Clustering

Description
Cluster labels of a clusterings are replaced by 1:length(table(cl)).

Usage
norm.label(cl)

Arguments
c1  vector of cluster memberships

Value
the clustering with normed labels.

Author(s)
Arno Fritsch, <arno.fritsch@tu-dortmund.de>

See Also
relabel for labelling a sample of clusterings the same way

Examples
(cl <- sample(c(13,12,34), 13, replace=TRUE))
norm.label(cl)

(cl <- sample(c("a","b","f31"), 13, replace=TRUE))
norm.label(cl)

relabel  

Stephens' Relabelling Algorithm for Clusterings

Description
For a sample of clusterings in which corresponding clusters have different labels the algorithm attempts to bring the clusterings to a unique labelling.

Usage
relabel(cls, print.loss = TRUE)
Arguments

- cls: a matrix in which every row corresponds to a clustering of the ncol(cls) objects.
- print.loss: logical, should current value of loss function be printed after each iteration? Defaults to TRUE.

Details

The algorithm minimizes the loss function

$$\sum_{m=1}^{M} \sum_{i=1}^{n} \sum_{j=1}^{K} -\log \hat{p}_{ij} \cdot I\{z_{i}^{(m)} = j\}$$

over the $M$ clusterings, $n$ observations and $K$ clusters, where $\hat{p}_{ij}$ is the estimated probability that observation $i$ belongs to cluster $j$ and $z_{i}^{(m)}$ indicates to which cluster observation $i$ belongs in clustering $m$. $I\{\cdot\}$ is an indicator function.

Minimization is achieved by iterating the estimation of $\hat{p}_{ij}$ over all clusterings and the minimization of the loss function in each clustering by permuting the cluster labels. The latter is done by linear programming.

Value

- cls: the input cls with unified labelling.
- P: an $n \times K$ matrix, where entry $[i, j]$ contains the estimated probability that observation $i$ belongs to cluster $j$.
- loss.val: value of the loss function.
- cl: vector of cluster memberships that have the highest probabilities $\hat{p}_{ij}$.

Warning

The algorithm assumes that the number of clusters $K$ is fixed. If this is not the case $K$ is taken to be the most common number of clusters. Clusterings with other numbers of clusters are discarded and a warning is issued.

Note

The implementation is a variant of the algorithm of Stephens which is originally applied to draws of parameters for each observation, not to cluster labels.

Author(s)

Arno Fritsch, <arno.fritsch@tu-dortmund.de>

References

See Also

`lp.transport` for the linear programming, `maxpear`, `minbinder`, `medv` for other possibilities of processing a sample of clusterings.

Examples

```r
(cls <- rbind(c(1,1,2,2),c(1,1,2,2),c(1,2,2,2),c(2,2,1,1)))
# group 2 in clustering 4 corresponds to group 1 in clustering 1-3.
cls.relab <- relabel(cls)
cls.relab$cls
```

---

**vi.dist**

*Variation of Information Distance for Clusterings*

**Description**

Computes the 'variation of information' distance of Meila (2007) between two clusterings/partitions of the same objects.

**Usage**

```r
vi.dist(cl1, cl2, parts = FALSE, base = 2)
```

**Arguments**

- `cl1, cl2`: vectors of cluster memberships (need to have the same lengths).
- `parts`: logical; should the two conditional entropies also be returned?
- `base`: base of logarithm used for computation of entropy and mutual information.

**Details**

The variation of information distance is the sum of the two conditional entropies of one clustering given the other. For details see Meila (2007).

**Value**

The VI distance. If `parts=TRUE` the two conditional entropies are appended.

**Author(s)**

Arno Fritsch, <arno.fritsch@tu-dortmund.de>

**References**

See Also

arandi

Examples

cl1 <- sample(1:3,10,replace=TRUE)
c2 <- c(cl1[1:5], sample(1:3,5,replace=TRUE))
vi.dist(cl1,c2)
vi.dist(cl1,c2, parts=TRUE)
Simulated 3-dimensional Normal Data Containing 8 Clusters

Description
Cluster means are given by the 8 possible values of $(\pm 2, \pm 2, \pm 2)$ to which standard normal noise was added. True clusters are given by `rep(1:8, each =50)`.

Usage
data(Ysim2)

Format
matrix with 400 rows and 3 columns.

Source
Simulated by
2 * matrix(c(rep(c(1,1,1),50), rep(c(1,1,-1),50), rep(c(1,-1,1),50), rep(c(-1,1,1),50), 
rep(c(1,-1,-1),50), rep(c(-1,1,-1),50), rep(c(-1,-1,1),50), rep(c(-1,-1,-1),50)),
byrow=TRUE, ncol=3) + matrix(rnorm(400*3),ncol=3)

References
Index

* cluster
  arandi, 3
  cltoSim, 5
  comp.psm, 6
  maxpear, 7
  medv, 8
  minbinder, 9
  norm.label, 12
  relabel, 12
  vi.dist, 14

* datasets
  cls.draw1.5, 4
  cls.draw2, 5
  Ysim1.5, 15
  Ysim2, 16

* optimize
  maxpear, 7
  minbinder, 9

* package
  mcclust-package, 2

arandi, 3, 15

binder (minbinder), 9

cls.draw1.5, 4
cls.draw2, 5
cltoSim, 5, 7
comp.psm, 6, 6, 8, 9, 11

laugreen (minbinder), 9
lp, 11
lp.transport, 14

maxpear, 7, 9, 11, 14
mcclust (mcclust-package), 2
mcclust-package, 2
medv, 8, 8, 11, 14
minbinder, 8, 9, 9, 14

norm.label, 12

pear (maxpear), 7
relabel, 8, 9, 11, 12, 12
Simtocl (cltoSim), 5
vi.dist, 4, 14
Ysim1.5, 15
Ysim2, 16