

Package ‘pivmet’

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Type Package

Title Pivotal methods for Bayesian relabelling and k-means clustering

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Description The package provides some pivotal algorithms for dealing with Bayesian Gaussian mixture models and relabelling the MCMC chains in order to undo the label switching problem. The same pivotal methods may be used for initializing the centers of the classical k-means algorithm in order to obtain a better clustering solution.

url <https://github.com/leoegidi/pivmet>

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R topics documented:

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MUS	<i>MUS algorithm</i>
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Description

Perform Maxima Units Search (MUS) algorithm on a large and sparse matrix in order to find a set of pivotal units through a sequential search in the given matrix.

Usage

MUS(C, clusters, prec_par = 5)

Arguments

C	$N \times N$ matrix with a non-negligible number of zeros. For instance, a similarity matrix estimated from a $N \times D$ data matrix whose rows are statistical units, or a co-association matrix resulting from clustering ensembles.
clusters	A vector of integers from 1:k (with $k \leq 4$) indicating a partition of the N units resulting from clustering.
prec_par	Optional argument. The maximum number of alternative pivots for each group.

Details

Consider H distinct partitions of a set of N d -dimensional statistical units into k groups determined by some clustering technique. A $N \times N$ co-association matrix C with generic element $c_{i,j} = n_{i,j}/H$ can be constructed, where $n_{i,j}$ is the number of times the i -th and the j -th unit are assigned to the same cluster with respect to the clustering ensemble. Units which are very distant from each other are likely to have zero co-occurrences; as a consequence, C is a square symmetric matrix expected to contain a non-negligible number of zeros. The main task of the MUS algorithm is to detect submatrices of small rank from the co-association matrix and extract those units—pivots—such that the $k \times k$ submatrix of C , determined by only the pivotal rows and columns indexes, is identical or nearly identical. Practically, the resulting units have the desirable property to be representative of the group they belong to.

Value

pivots	The k pivotal units
--------	---------------------

References

Egidi, L., Pappadà, R., Pauli, F., Torelli, N. (2018). Maxima Units Search(MUS) algorithm: methodology and applications. In: Perna, C. , Pratesi, M., Ruiz-Gazen A. (eds.) Studies in Theoretical and Applied Statistics, Springer Proceedings in Mathematics and Statistics 227, pp. 71–81.

Examples

```
# Data generated from a mixture of three bivariate Gaussian distributions

N <- 620
centers <- 3
n1 <- 20
n2 <- 100
n3 <- 500
x <- matrix(NA, N, 2)
truegroup <- c( rep(1,n1), rep(2, n2), rep(3, n3))

for (i in 1:n1){
  x[i,]=rmvnorm(1, c(1,5), sigma=diag(2))}
for (i in 1:n2){
  x[n1+i,]=rmvnorm(1, c(4,0), sigma=diag(2))}
for (i in 1:n3){
  x[n1+n2+i,]=rmvnorm(1, c(6,6), sigma=diag(2))}

# Build a similarity matrix from clustering ensembles

H <- 1000
a <- matrix(NA, H, N)

for (h in 1:H){
  a[h,] <- kmeans(x,centers)$cluster
}

sim_matr <- matrix(1, N,N)
for (i in 1:(N-1)){
  for (j in (i+1):N){
    sim_matr[i,j] <- sum(a[,i]==a[,j])/H
    sim_matr[j,i] <- sim_matr[i,j]
  }
}

# Obtain a clustering solution via kmeans with multiple random seeds

cl <- KMeans(x, centers)$cluster

# Find three pivots

mus_alg <- MUS(C = sim_matr, clusters = cl)
```

Description

Perform classical k-means clustering on a data matrix using pivots as initial centers.

Usage

```
piv_KMeans(x, centers, alg.type = c("KMeans", "hclust"),
  piv.criterion = c("MUS", "maxsumint", "maxsumnoint", "maxsumdiff"),
  H = 1000, ...)
```

Arguments

x	A $N \times D$ data matrix, or an object that can be coerced to such a matrix (such as a numeric vector or a dataframe with all numeric columns).
centers	The number of clusters in the solution.
alg.type	The clustering algorithm for the initial partition of the N units into the desired number of clusters. Possible choices are "KMeans" and "hclust".
piv.criterion	The pivotal criterion used for identifying one pivot for each group. Possible choices are: "MUS", "maxsumint", "maxsumnoint", "maxsumdiff". If <code>centers <= 4</code> , the default method is "MUS"; otherwise, the default method is "maxsumdiff" (see the details and the vignette).
H	If "MUS" is selected, this is the number of distinct k-means partitions used for building a $N \times N$ co-association matrix.
...	Optional arguments to be passed to MUS or KMeans.

Details

The function implements a modified version of k-means which aims at improving the clustering solution starting from a careful seeding. In particular, it performs a pivot-based initialization step using pivotal methods to find the initial centers for the clustering procedure. The starting point consists of multiple runs of the classical k-means (which uses random seeds via Kmeans function of the RcmdrMisc package) with a fixed number of clusters in order to build the co-association matrix of data units.

Value

A list with components

cluster	A vector of integers indicating the cluster to which each point is allocated.
centers	A matrix of cluster centres (centroids).
totss	The total sum of squares.
withinss	The within-cluster sum of squares for each cluster.
tot.withinss	The within-cluster sum of squares summed across clusters.
betweennss	The between-cluster sum of squared distances.
size	The number of points in each cluster.
iter	The number of (outer) iterations.
ifault	integer: indicator of a possible algorithm problem – for experts.
pivots	The pivotal units identified by the selected pivotal criterion.

Author(s)

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References

Egidi, L., Pappadà, R., Pauli, F., Torelli, N. (2018). K-means seeding via MUS algorithm. Conference Paper, Book of Short Papers, SIS2018, ISBN: 9788891910233.

Examples

```
# Data generated from a mixture of three bivariate Gaussian distributions

N <- 620
k <- 3
n1 <- 20
n2 <- 100
n3 <- 500
x <- matrix(NA, N, 2)
truegroup <- c( rep(1,n1), rep(2, n2), rep(3, n3))

for (i in 1:n1){
  x[i,]=rmvnorm(1, c(1,5), sigma=diag(2))}
for (i in 1:n2){
  x[n1+i,]=rmvnorm(1, c(4,0), sigma=diag(2))}
for (i in 1:n3){
  x[n1+n2+i,]=rmvnorm(1, c(6,6), sigma=diag(2))}

# Apply piv_KMeans with MUS as pivotal criterion

res <- piv_KMeans(x, k)

# Apply piv_KMeans with maxsumdiff as pivotal criterion

res2 <- piv_KMeans(x, k, piv.criterion ="maxsumdiff")

# Plot the data and the clustering solution

par(mfrow=c(1,2), pty="s")
colors_cluster <- c("grey", "darkolivegreen3", "coral")
colors_centers <- c("black", "darkgreen", "firebrick")
plot(x, col = colors_cluster[truegroup],
     bg= colors_cluster[truegroup], pch=21, xlab="x[,1]",
     ylab="x[,2]", cex.lab=1.5,
     main="True data", cex.main=1.5)

plot(x, col = colors_cluster[res$cluster],
     bg=colors_cluster[res$cluster], pch=21, xlab="x[,1]",
     ylab="x[,2]", cex.lab=1.5,
     main="piv_KMeans", cex.main=1.5)
points(x[res$pivots[1],1], x[res$pivots[1],2],
       pch=24, col=colors_centers[1],bg=colors_centers[1],
       cex=1.5)
points(x[res$pivots[2],1], x[res$pivots[2],2],
       pch=24, col=colors_centers[2], bg=colors_centers[2],
       cex=1.5)
```

```

points(x[res$pivots[3],1], x[res$pivots[3],2],
      pch=24, col=colors_centers[3], bg=colors_centers[3],
      cex=1.5)
points(res$centers, col = colors_centers[1:k],
      pch = 8, cex = 2)

```

piv_MCMC

JAGS Sampling for Gaussian Mixture Models and Clustering via Co-Association Matrix.

Description

Perform MCMC JAGS sampling for Gaussian mixture models, post-process the chains and apply a clustering technique to the MCMC sample. Pivotal units for each group are selected among four alternative criteria.

Usage

```

piv_MCMC(y, k, priors, nMC, piv.criterion = c("MUS", "maxsumint",
      "maxsumnoint", "maxsumdiff"), clustering = c("diana", "hclust"))

```

Arguments

y	N-dimensional data vector/matrix.
k	Number of mixture components.
priors	Input prior hyperparameters (see Details).
nMC	Number of MCMC iterations for the JAGS function execution.
piv.criterion	The pivotal criterion used for identifying one pivot for each group. Possible choices are: "MUS", "maxsumint", "maxsumnoint", "maxsumdiff". If $k \leq 4$, the default method is "MUS"; otherwise, the default method is "maxsumdiff" (see the Details and the vignette).
clustering	The clustering technique adopted for partitioning the N observations into k groups. Possible choices: "diana" (default), "hclust".

Details

The function fits univariate and bivariate Bayesian Gaussian mixture models of the form (here for univariate only):

$$(Y_i | Z_i = j) \sim \mathcal{N}(\mu_j, \phi_j),$$

where the Z_i , $i = 1, \dots, N$, are i.i.d. random variables, $j = 1, \dots, k$, ϕ_j is the group variance, $Z_i \in 1, \dots, k$ are the latent group allocation, and

$$P(Z_i = j) = \pi_j.$$

The likelihood of the model is then

$$L(y; \mu, \pi, \phi) = \prod_{i=1}^N \sum_{j=1}^k \pi_j \mathcal{N}(\mu_j, \phi_j),$$

where $(\mu, \phi) = (\mu_1, \dots, \mu_k, \phi_1, \dots, \phi_k)$ are the component-specific parameters and $\pi = (\pi_1, \dots, \pi_k)$ the mixture weights. Let ν denote a permutation of $1, \dots, k$, and let $\nu(\mu) = (\mu_{\nu(1)}, \dots, \mu_{\nu(k)})$,

$\nu(\phi) = (\phi_{\nu(1)}, \dots, \phi_{\nu(k)})$, $\nu(\pi) = (\pi_{\nu(1)}, \dots, \pi_{\nu(k)})$ be the corresponding permutations of μ , ϕ and π . Denote by V the set of all the permutations of the indexes $1, \dots, k$, the likelihood above is invariant under any permutation $\nu \in V$, that is

$$L(y; \mu, \pi, \phi) = L(y; \nu(\mu), \nu(\pi), \nu(\phi)).$$

As a consequence, the model is unidentified with respect to an arbitrary permutation of the labels. When Bayesian inference for the model is performed, if the prior distribution $p_0(\mu, \pi, \phi)$ is invariant under a permutation of the indices, then so is the posterior. That is, if $p_0(\mu, \pi, \phi) = p_0(\nu(\mu), \nu(\pi), \nu(\phi))$, then

$$p(\mu, \pi, \phi|y) \propto p_0(\mu, \pi, \phi)L(y; \mu, \pi, \phi)$$

is multimodal with (at least) $k!$ modes.

Priors are chosen as weakly informative. For univariate mixtures, the specification is the same as the function `BMMmodel` of the `bayesmix` package:

$$\mu_j \sim \mathcal{N}(0, 1/B0inv)$$

$$\phi_j \sim \text{invGamma}(nu0Half, nu0S0Half)$$

$$\pi \sim \text{Dirichlet}(1, \dots, 1)$$

$$S0 \sim \text{Gamma}(g0Half, g0G0Half),$$

with default values: $B0inv = 0.1$, $nu0Half = 10$, $S0 = 2$, $nu0S0Half = nu0Half \times S0$, $g0Half = 5e - 17$, $g0G0Half = 5e - 33$, in accordance with the default specification:

```
priors=list(kind = "independence", parameter = "priorsFish", hierarchical = "tau")
```

(see `bayesmix` for further details and choices).

For bivariate mixtures, the prior specification is the following:

$$\boldsymbol{\mu}_j \sim \mathcal{N}_2(\boldsymbol{\mu}_0, S2)$$

$$1/\Sigma \sim \text{Wishart}(S3, 3)$$

$$\pi \sim \text{Dirichlet}(1, \dots, 1),$$

where $S2$ and $S3$ are diagonal matrices with diagonal elements (the variances) equal to $1e+05$. The user may specify other values for the hyperparameters $\boldsymbol{\mu}_0, S2, S3$ via `priors` argument in such a way:

```
priors =list(mu0 = c(1,1), S2 = ..., S3 = ...),
```

with the constraint for $S2$ and $S3$ to be positive definite.

The function performs JAGS sampling using the `bayesmix` package for univariate Gaussian mixtures, and the `runjags` package for bivariate Gaussian mixtures. After MCMC sampling, this function clusters the units in k groups, calls the `piv_sel()` function and yields the pivots obtained from one among four different methods (the user may specify one among them via `piv.criterion` argument): "maxsumint", "maxsumnoint", "maxsumdiff" and "MUS" (available only if $k \leq 4$) (see the vignette for thorough details). Due to computational reasons clarified in the Details section of the function `piv_rel`, the length of the MCMC chains will be minor or equal than the input argument `nMC`; this length, corresponding to the value `true.iter` returned by the procedure, is the number of MCMC iterations for which the number of JAGS groups exactly coincides with the prespecified number of groups k .

Value

The function gives the MCMC output, the clustering solutions and the pivotal indexes. Here there is a complete list of outputs.

<code>Freq</code>	$k \times 2$ matrix where: the first column reports the number of units allocated to each group as given by JAGS program; the second column reports the same number of units as given by the chains' post-processing.
<code>true.iter</code>	The number of MCMC iterations for which the number of JAGS groups exactly coincides with the prespecified number of groups k .
<code>z</code>	$N \times k \times \text{true.iter}$ array with values: 1, if the i -th unit belongs to the j -th group at the h -th iteration; 0, otherwise.
<code>ris</code>	MCMC output matrix as provided by JAGS.
<code>groupPost</code>	$\text{true.iter} \times N$ matrix with values from $1:k$ indicating the post-processed group allocation vector.
<code>mu_switch</code>	If y is a vector, a $\text{true.iter} \times k$ matrix with the post-processed MCMC chains for the mean parameters; if y is a matrix, a $\text{true.iter} \times 2 \times k$ array with the post-processed MCMC chains for the mean parameters.
<code>mu_raw</code>	If y is a vector, a $nMC \times k$ matrix with the raw MCMC chains for the mean parameters as given by JAGS; if y is a matrix, a $nMC \times 2 \times k$ array with the raw MCMC chains for the mean parameters as given by JAGS.
<code>C</code>	Co-association matrix constructed from the MCMC sample.
<code>grr</code>	Group vector allocation as provided by "diana" or "hclust".
<code>pivots</code>	The pivotal units identified by the selected pivotal criterion.
<code>piv.criterion</code>	Gives the pivotal criterion used for identifying the pivots.

Author(s)

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References

Egidi, L., Pappadà, R., Pauli, F. and Torelli, N. (2018). Relabelling in Bayesian Mixture Models by Pivotal Units. *Statistics and Computing*, 28(4), 957-969.

Examples

```
# Bivariate simulation

N <- 200
k <- 4
nMC <- 1000
M1 <- c(-.5, 8)
M2 <- c(25.5, .1)
M3 <- c(49.5, 8)
M4 <- c(63.0, .1)
Mu <- matrix(rbind(M1, M2, M3, M4), c(4, 2))
stdev <- cbind(rep(1, k), rep(200, k))
Sigma.p1 <- matrix(c(stdev[1, 1], 0, 0, stdev[1, 1]), nrow=2, ncol=2)
Sigma.p2 <- matrix(c(stdev[1, 2], 0, 0, stdev[1, 2]), nrow=2, ncol=2)
W <- c(0.2, 0.8)
```



```

sim <- piv_sim(N,k,Mu, stdev, Sigma.p1,Sigma.p2,W)
res <- piv_MCMC(y = sim$y, k =k, nMC = nMC)
#changing priors
res2 <- piv_MCMC(y = sim$y,
                 priors = list (
                   mu0=c(1,1),
                   S2 = matrix(c(0.002,0,0, 0.1),2,2, byrow=TRUE),
                   S3 = matrix(c(0.1,0,0,0.1), 2,2, byrow =TRUE)),
                 k = k, nMC = nMC)

# Fishery data (bayesmix package)

data(fish)
y <- fish[,1]
k <- 5
nMC <- 5000
res <- piv_MCMC(y = y, k = k, nMC = nMC)
# changing priors
res2 <- piv_MCMC(y = y,
                 priors = list(kind = "condconjugate",
                               parameter = "priorsRaftery",
                               hierarchical = "tau"), k =k, nMC = nMC)

```

piv_plot

*Plotting outputs from pivotal relabelling***Description**

Plot and visualize MCMC outputs, posterior relabelled chains and estimates and diagnostics.

Usage

```
piv_plot(y, mcmc, rel_est, type = c("chains", "estimates", "hist"))
```

Arguments

y	Data vector or matrix.
mcmc	The ouptut of the raw MCMC sampling, as provided by piv_MCMC.
rel_est	Pivotal estimates as provided by piv_rel.
type	Type of plots required. Choose among: "chains", "estimates", "hist".

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Examples

```
# Fishery data

data(fish)
y <- fish[,1]
N <- length(y)
k <- 5
nMC <- 5000
res <- piv_MCMC(y = y, k = k, nMC = nMC)
rel <- piv_rel(mcmc=res, nMC = nMC)
piv_plot(y, res, rel, "chains")
piv_plot(y, res, rel, "estimates")
piv_plot(y, res, rel, "hist")
```

piv_rel	<i>Performing the pivotal relabelling step and computing the relabelled posterior estimates</i>
---------	---

Description

This function allows to perform the pivotal relabelling procedure described in Egidi et al. (2018) and to obtain the relabelled posterior estimates.

Usage

```
piv_rel(mcmc, nMC)
```

Arguments

mcmc	The output of the MCMC sampling from piv_MCMC.
nMC	The number of total MCMC iterations (given in input to the piv_MCMC function, or any function suited for MCMC sampling).

Details

Prototypical models in which the label switching problem arises are mixture models, as explained in the Details section of the piv_MCMC function.

These models are unidentified with respect to an arbitrary permutation of the labels $1, \dots, k$. Relabelling means permuting the labels at each iteration of the Markov chain in such a way that the relabelled chain can be used to draw inferences on component-specific parameters.

We assume here that a MCMC sample is obtained for the posterior distribution of a Gaussian mixture model—for instance via piv_MCMC function—with a prior distribution which is labelling invariant. Furthermore, suppose that we can find k units, one for each group, which are (pairwise) separated with (posterior) probability one (that is, the posterior probability of any two of them being in the same group is zero). It is then straightforward to use the k units, called pivots in what follows and denoted by the indexes i_1, \dots, i_k , to identify the groups and to relabel the chains: for each MCMC iteration $h = 1, \dots, H$ (H corresponds to the argument nMC) and group $j = 1, \dots, k$, set

$$[\mu_j]_h = [\mu_{[Z_{i_j}]_h}]_h;$$

$$[Z_i]_h = j \text{ for } i : [Z_i]_h = [Z_{i_j}]_h.$$

The applicability of this strategy is limited by the existence of the pivots, which is not guaranteed. The existence of the pivots is a requirement of the method, meaning that its use is restricted to those chains—or those parts of a chain—for which the pivots are present. First, although the model is based on a mixture of k components, each iteration of the chain may imply a different number of non-empty groups. Let then $[k]_h \leq k$ be the number of non-empty groups at iteration h ,

$$[k]_h = \#\{j : [Z_i]_h = j \text{ for some } i\},$$

where $\#A$ is the cardinality of the set A . Hence, the relabelling procedure outlined above can be used only for the subset of the chain for which $[k]_h = k$; let it be

$$\mathcal{H}_k = \{h : [k]_h = k\},$$

which correspond to the argument `true.iter` given by `piv_MCMC`. This means that the resulting relabelled chain is not a sample (of size H) from the posterior distribution, but a sample (of size $\#\mathcal{H}_k$) from the posterior distribution conditional on there being (exactly) k non-empty groups. Even if k non-empty groups are available, however, there may not be k perfectly separated units. Let us define

$$\mathcal{H}_k^* = \{h \in \mathcal{H}_k : \exists r, s \text{ s.t. } [Z_{i_r}]_h = [Z_{i_s}]_h\}$$

that is, the set of iterations where (at least) two pivots are in the same group. In order for the pivot method to be applicable, we need to exclude iterations \mathcal{H}_k^* ; that is, we can perform the pivot relabelling on $\mathcal{H}_k - \mathcal{H}_k^*$, corresponding to the argument `final_it`.

Value

This function gives the relabelled posterior estimates—both mean and medians—obtained from the Markov chains of the MCMC sampling.

<code>final_it</code>	The final number of valid MCMC iterations, as explained in Details
<code>final_it_p</code>	<code>final_it/nMC</code>
<code>mu_rel_mean</code>	k -vector (in case of univariate mixture) or $k \times 2$ matrix (in case of bivariate mixture) of estimated posterior means for the mean parameters.
<code>mu_rel_median</code>	k -vector (in case of univariate mixture) or $k \times 2$ matrix (in case of bivariate mixture) of estimated posterior medians for the mean parameters.
<code>mu_rel</code>	<code>final_it</code> \times k matrix (in case of univariate mixtures) or <code>final_it</code> \times $2 \times k$ array (in case of bivariate mixtures) for the complete relabelled chains

Author(s)

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References

Egidi, L., Pappada, R., Pauli, F. and Torelli, N. (2018). Relabelling in Bayesian Mixture Models by Pivotal Units. *Statistics and Computing*, 28(4), 957-969.

Examples

```
#Univariate simulation

N <- 250
nMC <- 2500
k <- 3
p <- rep(1/k,k)
x <- 3
stdev <- cbind(rep(1,k), rep(200,k))
Mu <- seq(-trunc(k/2)*x, trunc(k/2)*x, length=k)
W <- c(0.2, 0.8)
sim <- piv_sim(N,k,Mu,stdev,W=W)
res <- piv_MCMC(y = sim$y, k =k, nMC = nMC)
rel <- piv_rel(mcmc=res, nMC = nMC)


#Bivariate simulation

N <- 200
k <- 3
nMC <- 5000
M1 <- c(-.5,8)
M2 <- c(25.5,.1)
M3 <- c(49.5,8)
Mu <- matrix(rbind(M1,M2,M3),c(k,2))
stdev <- cbind(rep(1,k), rep(200,k))
Sigma.p1 <- matrix(c(stdev[1,1],0,0,stdev[1,1]),
  nrow=2, ncol=2)
Sigma.p2 <- matrix(c(stdev[1,2],0,0,stdev[1,2]),
  nrow=2, ncol=2)
W <- c(0.2,0.8)
sim <- piv_sim(N,k,Mu,stdev,Sigma.p1,Sigma.p2,W)
res <- piv_MCMC(y = sim$y, k = k, nMC = nMC)
rel <- piv_rel(mcmc = res, nMC = nMC)
piv_plot(y=sim$y, mcmc=res, rel_est = rel, type="chains")
piv_plot(y=sim$y, mcmc=res, rel_est = rel,
  type="hist")
```

piv_sel

Pivotal Selection via Co-Association Matrix

Description

Finding the pivots according to three different methods involving a co-association matrix C.

Usage

```
piv_sel(C, clusters)
```

Arguments

C	A $N \times N$ co-association matrix, i.e. a matrix whose elements are co-occurrences of pair of units in the same cluster among H distinct partitions.
clusters	A vector of integers indicating a partition of the N units into, say, k groups.

Details

Given a set of N observations (y_1, y_2, \dots, y_N) (y_i may be a d -dimensional vector, $d \geq 1$), consider clustering methods to obtain H distinct partitions into k groups. The matrix C is the co-association matrix, where $c_{i,p} = n_{i,p}/H$, with $n_{i,p}$ the number of times the pair (y_i, y_p) is assigned to the same cluster among the H partitions.

Let j be the group containing units \mathcal{J}_j , the user may choose $i^* \in \mathcal{J}_j$ that maximizes one of the quantities:

$$\sum_{p \in \mathcal{J}_j} c_{i^*p}$$

or

$$\sum_{p \in \mathcal{J}_j} c_{i^*p} - \sum_{j \notin \mathcal{J}_j} c_{i^*p}.$$

These methods give the unit that maximizes the global within similarity ("maxsumint") and the unit that maximizes the difference between global within and between similarities ("maxsumdiff"), respectively. Alternatively, we may choose $i^* \in \mathcal{J}_j$, which minimizes:

$$\sum_{p \notin \mathcal{J}_j} c_{i^*p},$$

obtaining the most distant unit among the members that minimize the global dissimilarity between one group and all the others ("maxsumnoint"). See the vignette for further details.

Value

pivots	A matrix with k rows and three columns containing the indexes of the pivotal units for each method.
--------	---

Author(s)

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References

Egidi, L., Pappadà, R., Pauli, F. and Torelli, N. (2018). Relabelling in Bayesian Mixture Models by Pivotal Units. *Statistics and Computing*, 28(4), 957-969.

Examples

```
# Iris data

data(iris)
x<- iris[,1:4]
N <- length(iris[,1])
H <- 1000
a <- matrix(NA, H, N)
```

```

# Perform H k-means partitions

for (h in 1:H){
  a[h,] <- kmeans(x, centers = 3)$cluster
}
# Build the co-association matrix

C <- matrix(1, N,N)
for (i in 1:(N-1)){
  for (j in (i+1):N){
    C[i,j] <- sum(a[i]==a[j])/H
    C[j,i] <- C[i,j]
  }
}

km <- kmeans(x, centers =3)

# Find the pivots according to the three possible pivotal criterion

ris <- piv_sel(C, clusters = km$cluster)

plot(iris[,1], iris[,2], xlab = "Sepal.Length", ylab= "Sepal.Width",
col = km$cluster)

# Add the pivots according to maxsumdiff criterion

points( x[ris$pivot[,3]], c( "Sepal.Length", "Sepal.Width" )], col = 1:3,
cex =2, pch = 8 )

```

piv_sim

*Generate Data from a Gaussian Nested Mixture***Description**

Simulate N observations from a nested Gaussian mixture model with k pre-specified components under uniform group probabilities $1/k$, where each group is in turn drawn from a further level consisting of two subgroups.

Usage

```

piv_sim(N, k, Mu, stdev, Sigma.p1 = matrix(c(1, 0, 0, 1), 2, 2, byrow =
  TRUE), Sigma.p2 = matrix(c(100, 0, 0, 100), 2, 2, byrow = TRUE),
  W = c(0.5, 0.5))

```

Arguments

N	The desired sample size.
k	The desired number of mixture components.
Mu	The input mean vector/matrix.
stdev	A k x2 matrix of input standard deviations, one for each group (from 1 to k) and for each subgroup (from 1 to 2). For univariate mixtures only.
Sigma.p1	The covariance matrix for the first subgroup. For bivariate mixtures only.
Sigma.p2	The covariance matrix for the second subgroup. For bivariate mixtures only.
W	The vector for the mixture weights of the two subgroups,

Details

The functions allows to simulate values from a double (nested) univariate Gaussian mixture:

$$(Y_i|Z_i = j) \sim \sum_{s=1}^2 p_{js} \mathcal{N}(\mu_j, \sigma_{js}^2),$$

or from a bivariate nested Gaussian mixture:

$$(Y_i|Z_i = j) \sim \sum_{s=1}^2 p_{js} \mathcal{N}_2(\boldsymbol{\mu}_j, \Sigma_s),$$

where σ_{js}^2 is the variance for the group j and the subgroup s (stdev is the argument for specifying the $k \times 2$ standard deviations for univariate mixtures); Σ_s is the covariance matrix for the subgroup s , $s = 1, 2$, where the two matrices are specified by Sigma.p1 and Sigma.p2 respectively; μ_j and $\boldsymbol{\mu}_j$, $j = 1, \dots, k$ are the mean input vector and matrix respectively, specified by the argument Mu; W is a vector of dimension 2 for the subgroups weights.

Value

y	The N simulated observations.
true.group	A vector of integers from 1:k indicating the values of the latent variables Z_i .
subgroups	A 2 x N matrix with values 1 or 2 indicating the subgroup to which each observation is drawn from.

Examples

```
# Bivariate mixture simulation with three components

N <- 2000
k <- 3
M1 <- c(-45,8)
M2 <- c(45,.1)
M3 <- c(100,8)
Mu <- matrix(rbind(M1,M2,M3),c(k,2))
stdev <- cbind(rep(1,k), rep(200,k))
Sigma.p1 <- matrix(c(stdev[1,1],0,0,stdev[1,1]),
  nrow=2, ncol=2)
Sigma.p2 <- matrix(c(stdev[1,2],0,0,stdev[1,2]),
  nrow=2, ncol=2)
W <- c(0.2,0.8)
sim <- piv_sim(N, k, Mu, Sigma.p1 = Sigma.p1,
  Sigma.p2 = Sigma.p2, W)
plot(sim$y, xlab="y[,1]", ylab="y[,2]")
```

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