

Package ‘sams’

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Type Package

Title Merge-Split Samplers for Conjugate Bayesian Nonparametric Models

Version 0.4.3

Description Markov chain Monte Carlo samplers for posterior simulations of conjugate Bayesian non-parametric mixture models. Functionality is provided for Gibbs sampling as in Algorithm 3 of Neal (2000) <[DOI:10.1080/10618600.2000.10474879](https://doi.org/10.1080/10618600.2000.10474879)>, restricted Gibbs merge-split sampling as described in Jain & Neal (2004) <[DOI:10.1198/1061860043001](https://doi.org/10.1198/1061860043001)>, and sequentially-allocated merge-split sampling <[DOI:10.1080/00949655.2021.1998502](https://doi.org/10.1080/00949655.2021.1998502)>, as well as summary and utility functions.

License GPL-3

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R topics documented:

asCanonical	2
asClusterLabels	3
asSetPartition	3
clusterProportions	4
clusterTrace	5

clusterWithItem	6
createNewCluster	6
dCRP	7
getThetas	8
isCanonical	8
joinExistingCluster	9
nClusters	9
nealAlgorithm3	10
p18_bern	11
p18_corr_mvn	12
p18_mvn	12
p6_bern	13
p6_big_bern	13
p6_mvn	14
partitionEntropy	14
poch	15
psm	16
psmMergeSplit	17
psmMergeSplit_base	19
restrictedGibbsMergeSplit	20
seqAllocatedMergeSplit	22
seqAllocatedMergeSplit_base	24
simpleMergeSplit	25
sizeOfLargestCluster	27
transformedWeights	27

Index **28**

asCanonical	<i>Coerce a Vector of Cluster Labels to Canonical Form</i>
-------------	--

Description

Coerce a Vector of Cluster Labels to Canonical Form

Usage

asCanonical(partition)

Arguments

partition	A numeric vector representing a set partition of the integers 1, ..., n using cluster labels
-----------	--

Value

A numeric vector representing partition, but now in canonical form.

asClusterLabels	<i>Coerce a Set Partition in List Structure to Numeric Vectors of Cluster Label</i>
-----------------	---

Description

Coerce a Set Partition in List Structure to Numeric Vectors of Cluster Label

Usage

```
asClusterLabels(partition)
```

Arguments

partition A list representing a set partition of the integers 1, ..., n

Value

A numeric vector representing the set partition using cluster labels.

asSetPartition	<i>Coerce a Set Partition as Numeric Vectors of Cluster Labels to a List Structure</i>
----------------	--

Description

Coerce a Set Partition as Numeric Vectors of Cluster Labels to a List Structure

Usage

```
asSetPartition(partition)
```

Arguments

partition A numeric vector representing a partition of the integers 1, ..., n using cluster labels

Value

The set partition in a list structure.

clusterProportions *Compute the Proportion of Items in Each Cluster for All Partitions*

Description

Compute the Proportion of Items in Each Cluster for All Partitions

Usage

```
clusterProportions(partitions)
```

Arguments

partitions A matrix, with each row representing a set partition of the integers 1, ..., n as cluster labels

Value

A matrix whose columns represent the cumulative proportion of the data that correspond to that cluster.

Examples

```
# Neal (2000) model and data
nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log=TRUE)
  }
}

logPostPredict <- mkLogPosteriorPredictiveDensity()

nSamples <- 500L
partitions <- matrix(0, nrow=nSamples, ncol=length(nealData))
for ( i in 2:nSamples ) {
  partitions[i,] <- nealAlgorithm3(partitions[i-1,], logPostPredict, mass = 1.0, nUpdates = 2)
}
clusterProportions(partitions)
```

clusterTrace *Plot Traces of Cluster Sizes*

Description

Plot Traces of Cluster Sizes

Usage

```
clusterTrace(
  partitions,
  plot.cols = rep("black", ncol(partitions)),
  plot.title = ""
)
```

Arguments

partitions	A matrix, with each row a numeric vector cluster labels
plot.cols	A character vector of valid color names, whose length represents the maximum number of stacked traces to be plotted
plot.title	A character string to be used as the main title on the trace plot

Examples

```
# Neal (2000) model and data
nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log=TRUE)
  }
}

logPostPredict <- mkLogPosteriorPredictiveDensity()

nSamples <- 500L
partitions <- matrix(0, nrow=nSamples, ncol=length(nealData))
for ( i in 2:nSamples ) {
  partitions[i,] <- nealAlgorithm3(partitions[i-1,], logPostPredict, mass = 1.0, nUpdates = 2)
}

clusterTrace(partitions, plot.title = "Neal (2000) Data")
```

clusterWithItem *Identify Which Cluster Contains a Given Item*

Description

Identify Which Cluster Contains a Given Item

Usage

```
clusterWithItem(i, partition)
```

Arguments

i	Item index as an integer vector of length one
partition	Set partition of the integers 1, ..., <i>n</i> represented as either a numeric vector of cluster labels, or a list containing subsets of these integers

Value

A list consisting of

which An integer representing which cluster *i* belongs to

cluster The subset of indices that correspond to the same cluster as *i*

createNewCluster *Create a New Cluster with Given Item*

Description

Create a New Cluster with Given Item

Usage

```
createNewCluster(i, partition)
```

Arguments

i	Item index as an integer vector of length one
partition	Set partition of the integers 1, ..., <i>n</i> represented as either a numeric vector of cluster labels, or a list containing subsets of these integers

Value

Updated partition with a new cluster.

dCRP	<i>Compute Probability Mass of a Partition Under the Two Parameter Chinese Restaurant Process (CRP)</i>
------	---

Description

Compute Probability Mass of a Partition Under the Two Parameter Chinese Restaurant Process (CRP)

Usage

```
dCRP(partition, mass = 1, discount = 0, log = FALSE)
```

Arguments

partition	A numeric vector of cluster labels, or a matrix whose rows are numeric vectors of cluster labels
mass	A numeric value indicating the mass parameter in the CRP, which must be greater than the <code>-discount</code> argument
discount	A numeric value on the interval $[0,1)$, indicating the discount parameter of the two parameter CRP
log	A logical value indicating whether results should be returned on the log scale

Value

A numeric vector of probabilities, or log probabilities if `log = TRUE`.

Examples

```
partitions <- matrix(c(0,0,0,0,0,
                      0,0,0,0,1,
                      0,0,0,1,2,
                      0,0,1,2,3,
                      0,1,2,3,4), ncol = 5, nrow = 5, byrow = TRUE)

# discount = 0 shows higher probability for lower quantity of components
dCRP(partitions, mass = 1, discount = 0, log = FALSE)

# discount = 0.5 shows higher probability for higher quantity of components
dCRP(partitions, mass = 1, discount = 0.5, log = FALSE)
```

getThetas	<i>Get theta Parameters from a Numeric Vector of Cluster Labels and Unique phi Values</i>
-----------	---

Description

Get theta Parameters from a Numeric Vector of Cluster Labels and Unique phi Values

Usage

```
getThetas(partition, phi)
```

Arguments

partition	A numeric vector representing a partition of the integers 1, ..., n using cluster labels
phi	A list of unique model parameters whose length must equal the number of unique cluster labels in partition

Value

A numeric vector of model parameters $\theta_1, \dots, \theta_n$.

isCanonical	<i>Check if a Vector of Cluster Labels is in Canonical Form</i>
-------------	---

Description

Check if a Vector of Cluster Labels is in Canonical Form

Usage

```
isCanonical(partition)
```

Arguments

partition	A numeric vector representing a partition of the integers 1, ..., n using cluster labels
-----------	--

Value

Logical, indicating whether partition is in canonical form.

joinExistingCluster *Join Item to an Existing Cluster*

Description

Join Item to an Existing Cluster

Usage

```
joinExistingCluster(i, join, partition)
```

Arguments

i	Item index as an integer vector of length one
join	Label or index of cluster that i must join
partition	Set partition of the integers 1, ..., n represented as either a numeric vector of cluster labels, or a list containing subsets of these integers

Value

Updated partition.

nClusters *Count the Number of Clusters in a Set Partition*

Description

Count the Number of Clusters in a Set Partition

Usage

```
nClusters(partition)
```

Arguments

partition	A numeric vector representing a partition of the integers 1, ..., n using cluster labels
-----------	--

Value

The number of clusters in the given set partition as a numeric vector of length one.

Examples

```
p <- c(0,1,1,2,3,2,4,4,2)
nClusters(p)
```

 nealAlgorithm3

Conjugate Gibbs Sampler for a Partition

Description

Algorithm 3 from Neal (2000) to update the state of a partition based on the "Chinese Restaurant Process" (CRP) prior and a user-supplied log posterior predictive density function, with additional functionality for the two parameter CRP prior.

Usage

```
nealAlgorithm3(
  partition,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  mass = 1,
  discount = 0,
  nUpdates = 1L
)
```

Arguments

partition	A numeric vector of cluster labels representing the current partition.
logPosteriorPredictiveDensity	A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i y_{subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).
mass	A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the <code>-discount</code> argument.
discount	A numeric value on the interval [0,1) corresponding to the discount parameter in the two parameter CRP prior. Set to zero for the usual, one parameter CRP prior.
nUpdates	An integer giving the number of Gibbs scans before returning. This has the effect of thinning the Markov chain.

Value

A numeric vector giving the updated partition encoded using cluster labels.

References

Neal, R. M. (2000). Markov chain sampling methods for Dirichlet process mixture models. *Journal of computational and graphical statistics*, 9(2), 249-265.

Examples

```

nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log=TRUE)
  }
}

logPostPredict <- mkLogPosteriorPredictiveDensity()

nSamples <- 1000L
partitions <- matrix(0, nrow = nSamples, ncol = length(nealData))
for (i in 2:nSamples) {
  partitions[i,] <- nealAlgorithm3(partitions[i-1,], logPostPredict, mass = 1.0, nUpdates = 1)
}

# convergence and mixing diagnostics
nSubsets <- apply(partitions, 1, function(x) length(unique(x)))
mean(nSubsets)
sum(acf(nSubsets)$acf) - 1 # Autocorrelation time

entropy <- apply(partitions, 1, partitionEntropy)
plot.ts(entropy)

```

p18_bern

Multivariate Independent Bernoulli Data (p = 18)

Description

Multivariate categorical data, generated as five clusters of independent variables with different success probabilities.

Usage

p18_bern_1

p18_bern_2

p18_bern_3

Format

An object of class `data.frame` with 100 rows and 18 columns.

An object of class `data.frame` with 100 rows and 18 columns.

An object of class `data.frame` with 100 rows and 18 columns.

p18_corr_mvn	<i>Correlated Multivariate Normal Data (p = 18)</i>
--------------	---

Description

Multivariate continuous data, generated as five clusters with an AR(1) correlation structure determining the common covariance matrix for each observation.

Usage

p18_corr_mvn_1

p18_corr_mvn_2

p18_corr_mvn_3

Format

An object of class `matrix` (inherits from `array`) with 100 rows and 18 columns.

An object of class `matrix` (inherits from `array`) with 100 rows and 18 columns.

An object of class `matrix` (inherits from `array`) with 100 rows and 18 columns.

p18_mvn	<i>Independent Multivariate Normal Data (p = 18)</i>
---------	--

Description

Multivariate continuous data, generated as five clusters of independent variables with different means.

Usage

p18_mvn_1

p18_mvn_2

p18_mvn_3

Format

An object of class `matrix` (inherits from `array`) with 100 rows and 18 columns.

An object of class `matrix` (inherits from `array`) with 100 rows and 18 columns.

An object of class `matrix` (inherits from `array`) with 100 rows and 18 columns.

p6_bern

*Multivariate Independent Bernoulli Data ($p = 6$)***Description**

Multivariate categorical data, generated as five clusters of independent variables with different success probabilities.

Usage

`p6_bern_1`

`p6_bern_2`

`p6_bern_3`

Format

An object of class `data.frame` with 100 rows and 6 columns.

An object of class `data.frame` with 100 rows and 6 columns.

An object of class `data.frame` with 100 rows and 6 columns.

p6_big_bern

*Large Sample Multivariate Independent Bernoulli Data ($p = 6$)***Description**

Multivariate categorical data, generated as five clusters of independent variables with different success probabilities.

Usage

`p6_big_bern_1`

`p6_big_bern_2`

`p6_big_bern_3`

Format

An object of class `data.frame` with 1000 rows and 6 columns.

An object of class `data.frame` with 1000 rows and 6 columns.

An object of class `data.frame` with 1000 rows and 6 columns.

p6_mvn

Independent Multivariate Normal Data (p = 6)

Description

Multivariate continuous data, generated as five clusters of independent variables with different means.

Usage

p6_mvn_1

p6_mvn_2

p6_mvn_3

Format

An object of class `matrix` (inherits from `array`) with 100 rows and 6 columns.

An object of class `matrix` (inherits from `array`) with 100 rows and 6 columns.

An object of class `matrix` (inherits from `array`) with 100 rows and 6 columns.

partitionEntropy

Calculate the Entropy of a Set Partition

Description

Calculate the Entropy of a Set Partition

Usage

partitionEntropy(partition)

Arguments

`partition` A numeric vector representing a partition of the integers 1, ..., n using cluster labels

Value

Calculated partition entropy as a numeric vector of length one

Examples

```
p <- c(0,0,0,1,1,2) # n = 6, 3 unique clusters
partitionEntropy(p)
```

poch

Compute the Pochhammer Symbol (Rising Factorials) With Increment

Description

Compute the Pochhammer Symbol (Rising Factorials) With Increment

Usage

```
poch(x, y = NULL, n = 1, log = FALSE)
```

Arguments

x	Non-negative numeric value
y	Non-negative real value representing increment parameter for Pochhammer function. If NULL, there is no increment (i.e. y=1).
n	Non-negative integer representing subscript in Pochhammer symbol
log	Logical value indicating whether to return results on log scale

Value

A numeric value indicating the result of Pochhammer function.

Examples

```
# effect of increment parameter
poch(5, y = NULL, n = 3, log = FALSE)
poch(5, y = 1, n = 3, log = FALSE)
poch(5, y = 1:4, n = 3, log = FALSE)

# increment being NULL is equivalent to ratio of gamma functions
a <- 7
b <- 3
out1 <- poch(a, y = NULL, n = b, log = FALSE)
out2 <- gamma(a + b) / gamma(a)
```

psm

*Compute the Posterior Pairwise Similarity for All Pairs of Items***Description**

Compute the Posterior Pairwise Similarity for All Pairs of Items

Usage

```
psm(partitions)
```

Arguments

`partitions` A matrix, with each row a numeric vector cluster labels

Value

A symmetric matrix of pairwise similarities based on the partitions given.

Examples

```
# Neal (2000) model and data
nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log=TRUE)
  }
}

logPostPredict <- mkLogPosteriorPredictiveDensity()

nSamples <- 500L
partitions <- matrix(0, nrow=nSamples, ncol=length(nealData))
for ( i in 2:nSamples ) {
  partitions[i,] <- nealAlgorithm3(partitions[i-1,], logPostPredict, mass = 1.0, nUpdates = 2)
}

psm(partitions)
```

psmMergeSplit	<i>Merge-Split Sampling for a Partition Based on Sequential Allocation Informed by Pairwise Similarities</i>
---------------	--

Description

Merge-split proposals for conjugate "Chinese Restaurant Process" (CRP) mixture models using sequentially-allocated elements. Allocation is performed with weights derived from a previously-calculated pairwise similarity matrix, and optionally complemented with "restricted Gibbs" scans as discussed in Jain & Neal (2004).

Usage

```
psmMergeSplit(
  partition,
  psm,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  t = 1,
  mass = 1,
  discount = 0,
  nUpdates = 1L,
  selectionWeights = NULL
)
```

Arguments

partition	A numeric vector of cluster labels representing the current partition.
psm	A matrix of previously-calculated pairwise similarity probabilities for each pair of data indices.
logPosteriorPredictiveDensity	A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i y_{subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).
t	A non-negative integer indicating the number of restricted Gibbs scans to perform for each merge/split proposal.
mass	A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the <code>-discount</code> argument.
discount	A numeric value on the interval $[0,1)$ corresponding to the discount parameter in the two-parameter CRP prior.
nUpdates	An integer giving the number of merge-split proposals before returning. This has the effect of thinning the Markov chain.
selectionWeights	A matrix or data frame whose first two columns are the unique pairs of data indices, along with a column of weights representing how likely each pair is to be selected at the beginning of each merge-split update.

Value

partition A numeric vector giving the updated partition encoded using cluster labels.

accept The acceptance rate of the Metropolis-Hastings proposals, i.e. the number of accepted proposals divided by nUpdates.

References

Jain, S., & Neal, R. M. (2004). A split-merge Markov chain Monte Carlo procedure for the Dirichlet process mixture model. *Journal of computational and Graphical Statistics*, 13(1), 158-182.

Examples

```
# Neal (2000) model and data
nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log=TRUE)
  }
}

logPostPredict <- mkLogPosteriorPredictiveDensity()

nSamples <- 1100L
nBurn <- 100
partitions <- matrix(0, nrow=nSamples, ncol=length(nealData))

# initial draws to inform similarity matrix
for ( i in 2:nBurn ) {
  partitions[i,] <- nealAlgorithm3(partitions[i-1,],
                                  logPostPredict,
                                  mass = 1,
                                  nUpdates = 1)
}

# Generate pairwise similarity matrix from initial draws
psm.mat <- psm(partitions[1:nBurn,])

accept <- 0
for ( i in (nBurn+1):nSamples ) {
  ms <- psmMergeSplit(partitions[i-1,],
                      psm.mat,
                      logPostPredict,
                      t = 1,
                      mass = 1.0,
```

```

        nUpdates = 1)
    partitions[i,] <- ms$partition
    accept <- accept + ms$accept
}

accept / (nSamples - nBurn) # post burn-in M-H acceptance rate
nSubsets <- apply(partitions, 1, function(x) length(unique(x)))
mean(nSubsets)
sum(acf(nSubsets)$acf)-1 # Autocorrelation time

entropy <- apply(partitions, 1, partitionEntropy)
plot.ts(entropy)

```

psmMergeSplit_base *Base Functionality for the psmMergeSplit Function*

Description

Merge-split proposals for conjugate "Chinese Restaurant Process" (CRP) mixture models using sequentially-allocated elements. Allocation is performed with weights derived from a previously-calculated pairwise similarity matrix.

Usage

```

psmMergeSplit_base(
  partition,
  psm,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  mass = 1,
  discount = 0,
  nUpdates = 1L,
  selectionWeights = NULL
)

```

Arguments

partition	A numeric vector of cluster labels representing the current partition.
psm	A matrix of previously-calculated pairwise similarity probabilities for each pair of data indices.
logPosteriorPredictiveDensity	A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i y_{subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).
mass	A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the <code>-discount</code> argument.

<code>discount</code>	A numeric value on the interval [0,1) corresponding to the discount parameter in the two-parameter CRP prior.
<code>nUpdates</code>	An integer giving the number of merge-split proposals before returning. This has the effect of thinning the Markov chain.
<code>selectionWeights</code>	A matrix or data frame whose first two columns are the unique pairs of data indices, along with a column of weights representing how likely each pair is to be selected at the beginning of each merge-split update.

Value

partition A numeric vector giving the updated partition encoded using cluster labels.

accept The acceptance rate of the Metropolis-Hastings proposals, i.e. the number of accepted proposals divided by `nUpdates`.

See Also

[psmMergeSplit](#)

`restrictedGibbsMergeSplit`

Merge-Split Sampling for a Partition Based on Restricted Gibbs Scans

Description

Merge-split proposals for conjugate "Chinese Restaurant Process" (CRP) mixture models using restricted Gibbs scans from a uniformly random launch state, as presented in Jain & Neal (2004), with additional functionality for the two parameter CRP prior.

Usage

```
restrictedGibbsMergeSplit(
  partition,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  t = 1,
  mass = 1,
  discount = 0,
  nUpdates = 1L,
  selectionWeights = NULL
)
```

Arguments

`partition` A numeric vector of cluster labels representing the current partition.

logPosteriorPredictiveDensity	A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i y_{subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).
t	A non-negative integer indicating the number of restricted Gibbs scans to perform for each merge/split proposal.
mass	A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the <code>-discount</code> argument.
discount	A numeric value on the interval $[0,1)$ corresponding to the discount parameter in the two parameter CRP prior.
nUpdates	An integer giving the number of merge-split proposals before returning. This has the effect of thinning the Markov chain.
selectionWeights	A matrix or data frame whose first two columns are the unique pairs of data indices, along with a column of weights representing how likely each pair is to be selected at the beginning of each merge-split update.

Value

partition An integer vector giving the updated partition encoded using cluster labels.

accept The acceptance rate of the Metropolis-Hastings proposals, i.e. the number accepted proposals divided by nUpdates.

References

Jain, S., & Neal, R. M. (2004). A split-merge Markov chain Monte Carlo procedure for the Dirichlet process mixture model. *Journal of computational and Graphical Statistics*, 13(1), 158-182.

Examples

```
# Neal (2000) model and data
nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log = TRUE)
  }
}

logPostPredict <- mkLogPosteriorPredictiveDensity()
```

```

nSamples <- 1000L
partitions <- matrix(0, nrow = nSamples, ncol = length(nealData))
accept <- 0
for ( i in 2:nSamples ) {
  ms <- restrictedGibbsMergeSplit(partitions[i-1,],
                                logPostPredict,
                                t = 1,
                                mass = 1.0,
                                nUpdates = 2)

  partitions[i,] <- ms$partition
  accept <- accept + ms$accept
}

accept / nSamples # M-H acceptance rate

# convergence and mixing diagnostics
nSubsets <- apply(partitions, 1, function(x) length(unique(x)))
mean(nSubsets)
sum(acf(nSubsets)$acf)-1 # Autocorrelation time

entropy <- apply(partitions, 1, partitionEntropy)
plot.ts(entropy)

```

```
seqAllocatedMergeSplit
```

Merge-split Sampling for a Partition Based on Sequential Allocation of Items

Description

Merge-split proposals for conjugate "Chinese Restaurant Process" (CRP) mixture models using sequential allocation of items, as originally described in Dahl (2003), with additional functionality for the two parameter CRP prior, as well as complementing these allocations with restricted Gibbs scans such as those discussed in Jain & Neal (2004).

Usage

```

seqAllocatedMergeSplit(
  partition,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  t = 1,
  mass = 1,
  discount = 0,
  nUpdates = 1L,
  selectionWeights = NULL
)

```

Arguments

partition	A numeric vector of cluster labels representing the current partition.
logPosteriorPredictiveDensity	A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i y_{subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).
t	A non-negative integer indicating the number of restricted Gibbs scans to perform for each merge/split proposal.
mass	A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the <code>-discount</code> argument.
discount	A numeric value on the interval $[0,1)$ corresponding to the discount parameter in the two parameter CRP prior.
nUpdates	An integer giving the number of merge-split proposals before returning. This has the effect of thinning the Markov chain.
selectionWeights	A matrix or data frame whose first two columns are the unique pairs of data indices, along with a column of weights representing how likely each pair is to be selected at the beginning of each merge-split update.

Value

partition An integer vector giving the updated partition encoded using cluster labels.

accept The acceptance rate of the Metropolis-Hastings proposals, i.e. the number accepted proposals divided by nUpdates.

References

Dahl, D. B. (2003). An improved merge-split sampler for conjugate Dirichlet process mixture models. Technical Report, 1, 086. Jain, S., & Neal, R. M. (2004). A split-merge Markov chain Monte Carlo procedure for the Dirichlet process mixture model. *Journal of computational and Graphical Statistics*, 13(1), 158-182.

Examples

```
# Neal (2000) model and data
nealData <- c(-1.48, -1.40, -1.16, -1.08, -1.02, 0.14, 0.51, 0.53, 0.78)
mkLogPosteriorPredictiveDensity <- function(data = nealData,
                                             sigma2 = 0.1^2,
                                             mu0 = 0,
                                             sigma02 = 1) {
  function(i, subset) {
    posteriorVariance <- 1 / ( 1/sigma02 + length(subset)/sigma2 )
    posteriorMean <- posteriorVariance * ( mu0/sigma02 + sum(data[subset])/sigma2 )
    posteriorPredictiveSD <- sqrt(posteriorVariance + sigma2)
    dnorm(data[i], posteriorMean, posteriorPredictiveSD, log=TRUE)
  }
}
```

```

    }
  }

logPostPredict <- mkLogPosteriorPredictiveDensity()

nSamples <- 1000L
partitions <- matrix(0, nrow = nSamples, ncol = length(nealData))
accept <- 0
for ( i in 2:nSamples ) {
  ms <- seqAllocatedMergeSplit(partitions[i-1,],
                              logPostPredict,
                              t = 1,
                              mass = 1.0,
                              nUpdates = 2)

  partitions[i,] <- ms$partition
  accept <- accept + ms$accept
}

accept / nSamples # M-H acceptance rate

# convergence and mixing diagnostics
nSubsets <- apply(partitions, 1, function(x) length(unique(x)))
mean(nSubsets)
sum(acf(nSubsets)$acf)-1 # Autocorrelation time

entropy <- apply(partitions, 1, partitionEntropy)
plot.ts(entropy)

```

```
seqAllocatedMergeSplit_base
```

Base Functionality for the seqAllocatedMergeSplit Function

Description

Merge-split proposals for conjugate "Chinese Restaurant Process" (CRP) mixture models using sequential allocation of items, as originally described in Dahl (2003), with additional functionality for the two parameter CRP prior.

Usage

```

seqAllocatedMergeSplit_base(
  partition,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  mass = 1,
  discount = 0,
  nUpdates = 1L,
  selectionWeights = NULL
)

```

Arguments

partition	A numeric vector of cluster labels representing the current partition.
logPosteriorPredictiveDensity	A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i y_{subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).
mass	A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the <code>-discount</code> argument.
discount	A numeric value on the interval $[0,1)$ corresponding to the discount parameter in the two-parameter CRP prior.
nUpdates	An integer giving the number of merge-split proposals before returning. This has the effect of thinning the Markov chain.
selectionWeights	A matrix or data frame whose first two columns are the unique pairs of data indices, along with a column of weights representing how likely each pair is to be selected at the beginning of each merge-split update.

Value

- partition** An integer vector giving the updated partition encoded using cluster labels.
- accept** The acceptance rate of the Metropolis-Hastings proposals, i.e. the number accepted proposals divided by `nUpdates`.

References

Dahl, D. B. (2003). An improved merge-split sampler for conjugate Dirichlet process mixture models. Technical Report, 1, 086.

See Also

[seqAllocatedMergeSplit](#)

simpleMergeSplit	<i>Merge-Split Sampling for a Partition Using Uniformly Random Allocation</i>
------------------	---

Description

Merge-split proposals for conjugate "Chinese Restaurant Process" (CRP) mixture models using uniformly random allocation of items, as presented in Jain & Neal (2004), with additional functionality for the two parameter CRP prior.

Usage

```
simpleMergeSplit(
  partition,
  logPosteriorPredictiveDensity = function(i, subset) 0,
  mass = 1,
  discount = 0,
  nUpdates = 1L,
  selectionWeights = NULL
)
```

Arguments

partition A numeric vector of cluster labels representing the current partition.

logPosteriorPredictiveDensity A function taking an index i (as a numeric vector of length one) and a subset of integers $subset$, and returning the natural logarithm of $p(y_i|y_{s subset})$, i.e., that item's contribution to the log integrated likelihood given a subset of the other items. The default value "turns off" the likelihood, resulting in prior simulation (rather than posterior simulation).

mass A specification of the mass (concentration) parameter in the CRP prior. Must be greater than the `-discount` argument.

discount A numeric value on the interval $[0,1)$ corresponding to the discount parameter in the two parameter CRP prior.

nUpdates An integer giving the number of merge-split proposals before returning. This has the effect of thinning the Markov chain.

selectionWeights A matrix or data frame whose first two columns are the unique pairs of data indices, along with a column of weights representing how likely each pair is to be selected at the beginning of each merge-split update.

Value

partition An integer vector giving the updated partition encoded using cluster labels.

accept The acceptance rate of the Metropolis-Hastings proposals, i.e. the number accepted proposals divided by `nUpdates`.

References

Jain, S., & Neal, R. M. (2004). A split-merge Markov chain Monte Carlo procedure for the Dirichlet process mixture model. *Journal of computational and Graphical Statistics*, 13(1), 158-182.

sizeOfLargestCluster *Calculate the Number of Items in the Largest Cluster of a Set Partition*

Description

Calculate the Number of Items in the Largest Cluster of a Set Partition

Usage

```
sizeOfLargestCluster(partition)
```

Arguments

partition	A numeric vector representing a partition of the integers 1, ..., n using cluster labels
-----------	--

Value

The number of items in the largest cluster of the given partition as a numeric vector of length one.

Examples

```
p <- c(0,1,1,1,1,1,2)
sizeOfLargestCluster(p)
```

transformedWeights *Enumerate Transformed Weights for Choosing i and j Non-Uniformly*

Description

Enumerate Transformed Weights for Choosing i and j Non-Uniformly

Usage

```
transformedWeights(m, fn = function(x) x, eps = 1e-12)
```

Arguments

m	A square matrix of pairwise similarities between items
fn	A function that maps pairwise similarities. Default is the identity function.
eps	A numeric value close to 0 to give some nonzero weight to pairs of items with 0 or 1 pairwise similarity

Index

* datasets

- p18_bern, [11](#)
 - p18_corr_mvn, [12](#)
 - p18_mvn, [12](#)
 - p6_bern, [13](#)
 - p6_big_bern, [13](#)
 - p6_mvn, [14](#)
- asCanonical, [2](#)
asClusterLabels, [3](#)
asSetPartition, [3](#)
- clusterProportions, [4](#)
clusterTrace, [5](#)
clusterWithItem, [6](#)
createNewCluster, [6](#)
- dCRP, [7](#)
- getThetas, [8](#)
- isCanonical, [8](#)
- joinExistingCluster, [9](#)
- nClusters, [9](#)
nealAlgorithm3, [10](#)
- p18_bern, [11](#)
p18_bern_1 (p18_bern), [11](#)
p18_bern_2 (p18_bern), [11](#)
p18_bern_3 (p18_bern), [11](#)
p18_corr_mvn, [12](#)
p18_corr_mvn_1 (p18_corr_mvn), [12](#)
p18_corr_mvn_2 (p18_corr_mvn), [12](#)
p18_corr_mvn_3 (p18_corr_mvn), [12](#)
p18_mvn, [12](#)
p18_mvn_1 (p18_mvn), [12](#)
p18_mvn_2 (p18_mvn), [12](#)
p18_mvn_3 (p18_mvn), [12](#)
p6_bern, [13](#)
p6_bern_1 (p6_bern), [13](#)
p6_bern_2 (p6_bern), [13](#)
p6_bern_3 (p6_bern), [13](#)
p6_big_bern, [13](#)
p6_big_bern_1 (p6_big_bern), [13](#)
p6_big_bern_2 (p6_big_bern), [13](#)
p6_big_bern_3 (p6_big_bern), [13](#)
p6_mvn, [14](#)
p6_mvn_1 (p6_mvn), [14](#)
p6_mvn_2 (p6_mvn), [14](#)
p6_mvn_3 (p6_mvn), [14](#)
partitionEntropy, [14](#)
poch, [15](#)
psm, [16](#)
psmMergeSplit, [17](#), [20](#)
psmMergeSplit_base, [19](#)
restrictedGibbsMergeSplit, [20](#)
seqAllocatedMergeSplit, [22](#), [25](#)
seqAllocatedMergeSplit_base, [24](#)
simpleMergeSplit, [25](#)
sizeOfLargestCluster, [27](#)
transformedWeights, [27](#)