

# Package ‘CHMM’

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**Type** Package

**Title** Coupled Hidden Markov Models

**Version** 0.1.1

**Description** An exact and a variational inference for coupled Hidden Markov Models applied to the joint detection of copy number variations.

**Depends** R (>= 3.1.0)

**Imports** mclust

**License** GPL-2

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 6.0.1

**URL** <http://github.com/julieaubert/CHMM>

**BugReports** <http://github.com/julieaubert/CHMM/issues>

**NeedsCompilation** yes

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CHMM\_EM

*Perform exact inference of coupled hidden markov models.***Description**

Perform exact inference of coupled hidden markov models.

**Usage**

```
CHMM_EM(X, nb.states, S, omega, meth.init = "mclust", var.equal = TRUE,
        itmax = 500, threshold = 1e-07)
```

**Arguments**

X	a data matrix of observations. Columns correspond to individuals.
nb.states	a integer specifying the numbers of states.
S	a matrix of similarity between individuals.
omega	a value of omega.
meth.init	a string specifying the initialization method ("mclust" or "kmeans"). The default method is "mclust".
var.equal	a logical variable indicating whether to treat the variances as being equal.
itmax	an integer specifying the maximal number of iterations for the EM algorithm.
threshold	a value for the threshold used for the stopping criteria.

**Value**

a list of 10 components

- postPr a list containing for each series the posterior probabilities.
- initGb a numeric specifying the initial state probabilities.
- transGb a matrix of the state transition probabilities.
- emisGb a list containing for each series the emission probabilities.
- esAvg a numeric of the estimated mean for each state.
- esVar a numeric of the estimated variance for each state.
- ID.K a matrix containing all combination of possible state for nbI series.
- loglik a numeric with the value of the loglikelihood.
- RSS a numeric corresponding to the Residuals Sum of Squares.
- iterstop an integer corresponding to the total number of iterations.

**References**

Wang, X., Lebarbier, E., Aubert, J. and Robin, S., Variational inference for coupled Hidden Markov Models applied to the joint detection of copy number variations.

CHMM\_VEM

*Perform variational inference of coupled Hidden Markov Models.***Description**

Perform variational inference of coupled Hidden Markov Models.

**Usage**

```
CHMM_VEM(X, nb.states, S = NULL, omega = 0.7, meth.init = "mclust",
  var.equal = TRUE, itmax = 500, threshold = 1e-07)
```

**Arguments**

X	a data matrix of observations. Columns correspond to individuals.
nb.states	a integer specifying the numbers of states.
S	a matrix of similarity between individuals.
omega	a value of omega.
meth.init	a string specifying the initialization method ("mclust" or "kmeans"). The default method is "mclust".
var.equal	a logical variable indicating whether to treat the variances as being equal.
itmax	an integer specifying the maximal number of iterations for the EM algorithm.
threshold	a value for the threshold used for the stopping criteria.

**Value**

a list of 9 components

postPr a list containing for each series the posterior probabilities.

initPr a numeric specifying the initial state probabilities.

transPr a matrix of the state transition probabilities.

esAvg a numeric of the estimated mean for each state.

esVar a numeric of the estimated variance for each state.

emisPr a list containing for each series the emission probabilities.

emisPrW a list containing for each series the emission probabilities taking into account for the dependency structure.

RSS a numeric corresponding to the Residuals Sum of Squares.

iterstop an integer corresponding to the total number of iterations.

**References**

Wang, X., Lebarbier, E., Aubert, J. and Robin, S., Variational inference for coupled Hidden Markov Models applied to the joint detection of copy number variations.

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clusterseg	<i>Summarize the results of the coupled HMM.</i>
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**Description**

Summarize the results of the coupled HMM.

**Usage**

```
clusterseg(x)
```

**Arguments**

x a matrix of status. Columns corresponds to series (individuals).

**Value**

a data.frame with 4 columns

sample name of the sample (series).

posbegin beginning position.

posend ending position.

status status.

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coupledHMM	<i>Perform inference of coupled hidden markov models.</i>
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**Description**

Perform inference of coupled hidden markov models.

**Usage**

```
coupledHMM(X, nb.states = 3, S = NULL, omega.list = c(0.3, 0.7, 0.9),  
var.equal = TRUE, exact = FALSE, meth.init = "mclust", viterbi = TRUE,  
itmax = 500, threshold = 1e-07)
```

**Arguments**

<code>X</code>	a matrix of observations. Columns correspond to series (individuals).
<code>nb.states</code>	a integer specifying the numbers of states.
<code>S</code>	a matrix of similarity between individuals.
<code>omega.list</code>	a vector of omega values.
<code>var.equal</code>	a logical variable indicating whether to treat the variances as being equal ( <code>var.equal = TRUE</code> ).
<code>exact</code>	a logical variable indicating whether to use VEM ( <code>exact = FALSE</code> ) or EM ( <code>exact = TRUE</code> ) algorithm for the inference of the model.
<code>meth.init</code>	a string specifying the initialization method ("mclust" or "kmeans") for the (V)-EM algorithm. The default method is "mclust".
<code>viterbi</code>	a logical variable indicating whether to use Maximum A Posteriori method ( <code>FALSE</code> ) or Viterbi algorithm ( <code>TRUE</code> , by default) for recovering the most likely path.
<code>itmax</code>	an integer specifying the maximal number of iterations for the CHMM_(V)EM algorithm.
<code>threshold</code>	a value for the threshold used for the stopping criteria for the CHMM_(V)EM algorithm.

**Value**

A list of 4 objets.

`omega` an integer corresponding to the selected value among the `omega.list`.

`model` a list corresponding to the output of the CHMM-EM or CHMM-VEM function for the selected model.

`status` a matrix with status associated to each series in column and each position in row.

`RSS.omega` a dataframe with omega values and the associated Residuals Sum of Squares.

**References**

Wang, X., Lebarbier, E., Aubert, J. and Robin, S., Variational inference for coupled Hidden Markov Models applied to the joint detection of copy number variations.

**See Also**

[CHMM\\_VEM](#), [CHMM\\_EM](#)

**Examples**

```
data(toyexample)
# Variational inference of a coupled hidden Markov Chains
resCHMM <- coupledHMM(X = toydata, nb.states = 3, S = cor(toystatus),
                      omega.list = c(0.3, 0.5, 0.7, 0.9))
# Breakpoints positions and status of segments
info <- clusterseg(resCHMM$status)
# head(info)
```

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init.EM                      *Initialization step of the CHMM\_EM function.*

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### Description

Initialization step of the CHMM\_EM function.

### Usage

```
init.EM(X, nb.states, meth.init, var.equal, nbI, nbT)
```

### Arguments

X	a matrix of observations. Columns correspond to series (individuals).
nb.states	an integer specifying the numbers of states.
meth.init	a string specifying the initialization method ("mclust" or "kmeans"). The default method is "mclust".
var.equal	a logical variable indicating whether to treat the variances as being equal (TRUE, value by default) or not (FALSE).
nbI	an integer specifying the number of series.
nbT	an integer specifying the length of one series.

### Details

By default, an initialization with the `meth.init="mclust"` is performed with homogeneous variances.

### Value

A list of 6 objects.

`esAvgGb` a matrix of  $\text{nbK}(\text{nb.states}^{\text{nbI}})$  rows and `nbI` columns of estimated mean.

`esVarGb` a matrix of  $\text{nbK}(\text{nb.states}^{\text{nbI}})$  rows and `nbI` columns of estimated variance.

`esAvg` a numeric of the estimated mean for each state.

`esVar` a numeric of the estimated variance for each state.

`transGb` a matrix of the state transition probabilities.

`initGb` a numeric specifying the initial state probabilities.

### See Also

[CHMM\\_EM](#)

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init.VEM	<i>Initialization step of the CHMM_VEM function.</i>
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**Description**

Initialization step of the CHMM\_VEM function.

**Usage**

```
init.VEM(X, nb.states, meth.init, var.equal, nbI, nbT)
```

**Arguments**

X	a matrix of observations. Columns correspond to series (individuals).
nb.states	an integer specifying the numbers of states.
meth.init	a string specifying the initialization method ("mclust" or "kmeans"). The default method is "mclust".
var.equal	a logical variable indicating whether to treat the variances as being equal (TRUE, value by default) or not (FALSE).
nbI	an integer specifying the number of series.
nbT	an integer specifying the length of one series.

**Value**

A list containing the parameters of the model

esAvg a numeric of the estimated mean for each state.

esVar a numeric of the estimated variance for each state.

transPr a matrix of the state transition probabilities

postPr a list containing for each series the posterior probabilities.

initPr a numeric specifying the initial state probabilities.

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toydata	<i>Toy example - observations for 5 correlated samples.</i>
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**Description**

A matrix containing the observations for the 1,000 positions of 5 correlated samples.

**Format**

A simulated matrix with 1000 rows and 5 columns. Each column is a series

**Examples**

```
data(toyexample)
# Variational inference of a coupled hidden Markov Chains
resCHMM <- coupledHMM(X = toydata, nb.states = 3, S = cor(toystatus),
                      omega.list = c(0.3, 0.5, 0.7, 0.9))
# Breakpoints positions and status of segments
info <- clusterseg(resCHMM$status)
# head(info)
```

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toystatus

*Toy example - status for 5 correlated samples.*

---

**Description**

A matrix containing the hidden status for the 1,000 positions of 5 correlated samples.

**Format**

A matrix of the hidden status corresponding to the toydata matrix.



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