

Package ‘mhsmm’

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Type Package

Title Parameter estimation and prediction for hidden Markov and semi-Markov models for data with multiple observation sequences.

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Description Parameter estimation and prediction for hidden Markov and semi-Markov models for data with multiple observation sequences. Suitable for equidistant time series data, with multivariate and/or missing data.

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LazyLoad yes

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addStates	<i>Adds a bar representing state sequence.</i>
-----------	--

Description

Add a colour coded horizontal bar representing the state sequence to a plot of (presumably time-series) data.

Usage

```
addStates(states, x=NULL,
          ybot = axTicks(2)[1], ytop=ybot + (axTicks(2)[2] - axTicks(2)[1]),
          dy = ytop - ybot,
          greyscale = FALSE, leg = NA, J = length(unique(states)),
          time.scale = 1, shiftx = 0)
```

Arguments

states	A vector of integers representing the states traversed
x	The time values where the states are observed ((1:length(states)-shiftx)/time.scale if NULL)
ybot	Vertical bottom limit of the bar.
ytop	Vertical top limit of the bar.
dy	Height of the bar.

greyscale	If TRUE produces a bar in greyscale.
leg	Array of state names, if present, produces a legend.
J	Number of states
time.scale	Resolution of the timescale
shiftx	Shift the bar forward or backwards horizontal by shiftx distance.

Author(s)

Soren Hojsgaard

See Also

addStates

Examples

```
plot(rnorm(100), type='l')
addStates(rep(c(1,2), each=50))

plot(seq(0.01, 1, .01), rnorm(100), type='l')
addStates(rep(c(1,2), each=50), seq(0.01, 1, .01))
```

dmvnorm.hsmm	<i>Emission ensity function for a multivariate normal emission distribu- tion</i>
--------------	---

Description

Calculates the density of observations x for state j given the parameters in `model`. This is used for a multivariate Gaussian emission distribution of a HMM or HSMM and is a suitable prototype for user's to make their own custom distributions.

Usage

```
dmvnorm.hsmm(x, j, model)
```

Arguments

x	Observed value
j	State
<code>model</code>	A <code>hsmmspec</code> or <code>hmmspec</code> object

Details

This is used by `hmm` and `hsmm` to calculate densities for use in the E-step of the EM algorithm. It can also be used as a template for users wishing to building their own emission distributions

Value

A vector of probability densities.

Author(s)

Jared O'Connell

See Also

mstep.mvnorm, rmvnorm.hsmm

Examples

```
J<-2
initial <- rep(1/J,J)
P <- matrix(c(.3, .5, .7, .5), nrow=J)
b <- list(mu=list(c(-3,0),c(1,2)), sigma=list(diag(2),matrix(c(4,2,2,3), ncol=2)))
model <- hmmspec(init=initial, trans=P, emission=b, f=dmvnorm.hsmm)
model
train <- simulate(model, nsim=300, seed=1234, r=rmvnorm.hsmm)
plot(train, xlim=c(0,100))
h1 = hmmfit(train,model,mstep=mstep.mvnorm)
```

dnorm.hsmm

Emission density function for normal emission distribution

Description

Calculates the density of observations x for state j given the parameters in `model`. This is used for the Gaussian emission distribution of a HMM or HSMM and is a suitable prototype for user's to make their own custom distributions.

Usage

```
dnorm.hsmm(x, j, model)
```

Arguments

<code>x</code>	Observed value
<code>j</code>	State
<code>model</code>	A <code>hsmmspec</code> or <code>hmmspec</code> object

Details

This is used by `hmm` and `hsmm` to calculate densities for use in the E-step of the EM algorithm. It can also be used as a template for users wishing to building their own emission distributions

Value

A vector of probability densities.

Author(s)

Jared O'Connell

dpois.hsmm

Emission density function for Poisson emission distribution

Description

Calculates the density of observations x for state j given the parameters in `model`. This is used for a Poisson emission distribution of a HMM or HSMM and is a suitable prototype for user's to make their own custom distributions.

Usage

```
dpois.hsmm(x, j, model)
```

Arguments

<code>x</code>	Observed value
<code>j</code>	State
<code>model</code>	A <code>hsmmspec</code> or <code>hmmspec</code> object

Details

This is used by `hmm` and `hsmm` to calculate densities for use in the E-step of the EM algorithm. It can also be used as a template for users wishing to building their own emission distributions

Value

A vector of probability densities.

Author(s)

Jared O'Connell

See Also

`mstep.pois`, `rpois.hsmm`

Examples

```
J<-3
initial <- rep(1/J,J)
P <- matrix(c(.8, .5, .1, 0.05, .2, .5, .15, .3, .4), nrow=J)
b <- list(lambda=c(1,3,6))
model <- hmmspec(init=initial, trans=P, emission=b, f=dpois.hsmm)
model
train <- simulate(model, nsim=300, seed=1234, r=rpois.hsmm)
plot(train, xlim=c(0,100))
h1 = hmmfit(train,model,mstep=mstep.pois)
```

gammafit

Parameter estimation for the Gamma distribution

Description

Estimates parameters for the Gamma distribution using the Method of Maximum Likelihood, works with weighted data.

Usage

```
gammafit(x, wt = NULL)
```

Arguments

x	A vector of observations
wt	Optional set of weights

Value

shape	The shape parameter
scale	The scale parameter (equal to 1/rate)

Author(s)

Jared O'Connell

References

Choi, S. and Wette, R. (1969), Maximum likelihood estimation of the parameters of the gamma distribution and their bias, *Technometrics*, 11, 683-96-690.

Examples

```
gammafit(rgamma(1000, shape=10, scale=13))
```

hmmfit *fit a hidden Markov model*

Description

Estimates parameters of a HMM using the EM algorithm.

Usage

```
hmmfit(x, start.val, mstep=mstep.norm, lock.transition=FALSE, tol=1e-08, maxit=1000)
```

Arguments

<code>x</code>	A <code>hsmm.data</code> object (see Details)
<code>start.val</code>	Starting parameters for the model (see Details)
<code>mstep</code>	Re-estimates the parameters of density function on each iteration
<code>lock.transition</code>	If TRUE will not re-estimate the transition matrix
<code>maxit</code>	Maximum number of iterations
<code>tol</code>	Convergence tolerance

Value

<code>start</code>	A vector of the starting probabilities for each state
<code>a</code>	The transition matrix of the embedded Markov chain
<code>emission</code>	A list of the parameters of the emission distribution

Author(s)

Jared O'Connell

References

Rabiner, L. (1989), A tutorial on hidden Markov models and selected applications in speech recognition, Proceedings of the IEEE, 77, 257-286.

See Also

`sim.hmm`, `sim.mhmm`, `predict.hmm`

Examples

```

J<-3
initial <- rep(1/J,J)
P <- matrix(c(.8,.5,.1,0.05,.2,.5,.15,.3,.4),nrow=J)
b <- list(mu=c(-3,0,2),sigma=c(2,1,.5))
model <- hmmspec(init=initial, trans=P, emission=b,f=dnorm.hsmm)
model

train <- simulate(model, nsim=300, seed=1234, r=rnorm.hsmm)
plot(train,xlim=c(0,100))

init0 <- rep(1/J,J)
P0 <- matrix(1/J,nrow=J,ncol=J)
b0 <- list(mu=c(-3,1,3),sigma=c(1,1,1))
startval <- hmmspec(init=init0, trans=P0,emission=b0,f=dnorm.hsmm)
h1 = hmmfit(train,startval,mstep=mstep.norm)

plot(h1$loglik,type='b',ylab='Log-likelihood',xlab='Iteration')
summary(h1)

#proportion of incorrect states
mean(train$s!=predict(h1,train)$s)

#simulate a new test set
test <- simulate(model, nsim=c(100,200,300), seed=1234,r=rnorm.hsmm)
mean(test$s!=predict(h1,test)$s)

```

hmmspec

Specificatin of HMMs

Description

Creates a model specification for a hidden Markov model

Usage

```
hmmspec(init, trans, emission, f, r=NULL, mstep=NULL)
```

Arguments

init	Distribution of states at t=1 ie. $P(S=s)$ at t=1
trans	The transition matrix of the Markov chain
emission	A list containing the parameters of the emission distribution
f	Density function of the emission distribution.
r	The function used to generate observations from the emission distribution
mstep	Re-estimates the parameters of density function on each iteration

Value

A hmmspec object

Author(s)

Jared O'Connell

References

Rabiner, L. (1989), A tutorial on hidden Markov models and selected applications in speech recognition, Proceedings of the IEEE, 77, 257-286.

See Also

simulate.hmmspec, simulate.hmm, hmm, predict.hmm

 hsmmfit

fit a hidden semi-Markov model

Description

Estimates parameters of a HSMM using the EM algorithm.

Usage

```
hsmmfit(x,model,mstep=NULL,M=NA,maxit=100,lock.transition=FALSE,lock.d=FALSE,graphical=FALSE)
```

Arguments

<code>x</code>	A hsmm.data object (see Details)
<code>model</code>	Starting parameters for the model (see hmmspec)
<code>mstep</code>	Re-estimates the parameters of density function on each iteration
<code>maxit</code>	Maximum number of iterations
<code>M</code>	Maximum number of time spent in a state (truncates the waiting distribution)
<code>lock.transition</code>	If TRUE will not re-estimate the transition matrix
<code>lock.d</code>	If TRUE will not re-estimate the sojourn time density
<code>graphical</code>	If TRUE will plot the sojourn densities on each iteration

Value

<code>start</code>	A vector of the starting probabilities for each state
<code>a</code>	The transition matrix of the embedded Markov chain
<code>emission</code>	A list of the parameters of the emission distribution
<code>waiting</code>	A list of the parameters of the waiting distribution

Author(s)

Jared O'Connell

References

Guedon, Y. (2003), Estimating hidden semi-Markov chains from discrete sequences, *Journal of Computational and Graphical Statistics*, Volume 12, Number 3, page 604-639 - 2003

See Also

hsmmspec, simulate.hsmmspec, predict.hsmm

Examples

```
J <- 3
init <- c(0,0,1)
P <- matrix(c(0, .1, .4, .5, 0, .6, .5, .9, 0), nrow=J)
B <- list(mu=c(10,15,20), sigma=c(2,1,1.5))
d <- list(lambda=c(10,30,60), shift=c(10,100,30), type='poisson')
model <- hsmmspec(init,P,emission=B,sojourn=d,f=dnorm.hsmm)
train <- simulate(model,r=rnorm.hsmm,nsim=100,seed=123456)
plot(train,xlim=c(0,400))
start.poisson <- hsmmspec(init=rep(1/J,J),transition=matrix(c(0, .5, .5, .5, 0, .5, .5, .5, 0), nrow=J,
  sigma=c(1,1,1)),sojourn=list(lambda=c(9,25,40),shift=c(5,95,45),type='poisson'))

M=500
h.poisson <- hsmmfit(train,start.poisson,mstep=mstep.norm,M=M)
plot(h.poisson$loglik,type='b',ylab='Log-likelihood',xlab='Iteration') ##has it converged?
summary(h.poisson)
predicted <- predict(h.poisson,train)
table(train$s,predicted$s) ##classification matrix
mean(predicted$s!=train$s) ##misclassification rate

d <- cbind(dunif(1:M,0,50),dunif(1:M,100,175),dunif(1:M,50,130))
start.np <- hsmmspec(init=rep(1/J,J),transition=matrix(c(0, .5, .5, .5, 0, .5, .5, .5, 0), nrow=J,em
  sigma=c(1,1,1)),sojourn=list(d=d,type='nonparametric'),f=dnorm.hsmm)
h.np <- hsmmfit(train,start.np,mstep=mstep.norm,M=M,graphical=TRUE)
mean(predicted$s!=train$s) ##misclassification rate
```

hsmmspec

Hidden semi-Markov model specification

Description

Creates a model specification of a hidden semi-Markov model.

Usage

```
hsmmspec(init,transition,emission,sojourn,f,r=NULL,mstep=NULL)
```

Arguments

<code>init</code>	Distribution of states at $t=1$ ie. $P(S=s)$ at $t=1$
<code>transition</code>	The transition matrix of the embedded Markov chain (diagonal must be 0)
<code>emission</code>	A list containing the parameters of the emission distribution
<code>sojourn</code>	A list containing the parameters and type of sojourn distribution (see Details)
<code>f</code>	Density function of the emission distribution
<code>r</code>	The function used to generate observations from the emission distribution
<code>mstep</code>	Re-estimates the parameters of density function on each iteration

Details

The sojourn argument provides a list containing the parameters for the available sojourn distributions. Available sojourn distributions are shifted Poisson, Gamma and non-parametric.

In the case of the Gamma distribution, sojourn is a list with vectors shape and scale (the Gamma parameters in `dgamma`), both of length J . Where J is the number of states. See `reprocows` for an example using Gamma sojourn distributions.

In the case of shifted Poisson, sojourn is list with vectors shift and lambda, both of length J . See `hsmmfit` for an example using shifted Poisson sojourn distributions.

In the case of non-parametric, sojourn is a list containing a $M \times J$ matrix. Where entry (i,j) is the probability of a sojourn of length i in state j . See `hsmmfit` for an example using shifted non-parametric sojourn distributions.

Value

An object of class `hsmmspec`

Author(s)

Jared O'Connell

References

Guedon, Y. (2003), Estimating hidden semi-Markov chains from discrete sequences, Journal of Computational and Graphical Statistics, Volume 12, Number 3, page 604-639 - 2003

See Also

`hsmm`, `simulate.hsmmspec`, `predict.hsmm`

mstep.mvnorm	<i>Performs re-estimation (the M-step) for a multivariate normal emission distribution</i>
--------------	--

Description

Re-estimates the parameters of a multivariate normal emission distribution as part of the EM algorithm for HMMs and HSMMs. This is called by the `hmm` and `hsmm` functions. It is a suitable prototype function for users wishing to design their own emission distributions.

Usage

```
mstep.mvnorm(x, wt)
```

Arguments

<code>x</code>	A vector of observed values
<code>wt</code>	A T x J matrix of weights. Column entries are the weights for respective states.

Details

Users may write functions that take the same arguments and return the same values for their own custom emission distributions.

Value

Returns the `emission` slot of a `hmmspec` or `hsmmspec` object

<code>mu</code>	A list of length J contain the mean vectors
<code>sigma</code>	A list of length J containing the covariance matrices

Author(s)

Jared O'Connell

See Also

`dmvnorm.hsmm`, `rmvnorm.hsmm`

Examples

```
J<-2
initial <- rep(1/J,J)
P <- matrix(c(.3, .5, .7, .5),nrow=J)
b <- list(mu=list(c(-3,0),c(1,2)),sigma=list(diag(2),matrix(c(4,2,2,3), ncol=2)))
model <- hmmspec(init=initial, trans=P, emission=b,f=dmvnorm.hsmm)
model
train <- simulate(model, nsim=300, seed=1234, r=rmvnorm.hsmm)
plot(train,xlim=c(0,100))
h1 = hmmfit(train,model,mstep=mstep.mvnorm)
```

mstep.norm	<i>Performs re-estimation (the M-step) for a normal emission distribution</i>
------------	---

Description

Re-estimates the parameters of a normal emission distribution as part of the EM algorithm for HMMs and HSMMs. This is called by the `hmm` and `hsmm` functions. It is a suitable prototype function for users wishing to design their own emission distributions.

Usage

```
mstep.norm(x, wt)
```

Arguments

<code>x</code>	A vector of observed values
<code>wt</code>	A T x J matrix of weights. Column entries are the weights for respective states.

Details

Users may write functions that take the same arguments and return the same values for their own custom emission distributions.

Value

Returns the `emission` slot of a `hmmspec` or `hsmmspec` object

<code>mu</code>	Vector of length J contain the means
-----------------	--------------------------------------

<code>mu</code>	Vector of length J containing the variances
-----------------	---

...

Author(s)

Jared O'Connell

mstep.pois	<i>Performs re-estimation (the M-step) for a Poisson emission distribution</i>
------------	--

Description

Re-estimates the parameters of a Poisson emission distribution as part of the EM algorithm for HMMs and HSMMs. This is called by the `hmm` and `hsmm` functions. It is a suitable prototype function for users wishing to design their own emission distributions.

Usage

```
mstep.pois(x, wt)
```

Arguments

<code>x</code>	A vector of observed values
<code>wt</code>	A T x J matrix of weights. Column entries are the weights for respective states.

Details

Users may write functions that take the same arguments and return the same values for their own custom emission distributions.

Value

Returns the emission slot of a `hmmspec` or `hsmmspec` object

<code>lambda</code>	Vector of length J containing the Poisson parameters for each state j
---------------------	---

...

Author(s)

Jared O'Connell

See Also

`rpois.hsmm`, `dpois.hsmm`

Examples

```
J<-3
initial <- rep(1/J,J)
P <- matrix(c(.8, .5, .1, 0.05, .2, .5, .15, .3, .4), nrow=J)
b <- list(lambda=c(1, 3, 6))
model <- hmmspec(init=initial, trans=P, emission=b, f=dpois.hsmm)
model
train <- simulate(model, nsim=300, seed=1234, r=rpois.hsmm)
```

```
plot(train,xlim=c(0,100))
h1 = hmmfit(train,model,mstep=mstep.pois)
```

plot.hsmm *Plot function for hsmms*

Description

Displays the densities for the sojourn distributions of each state.

Usage

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

Arguments

x A hsmm object
... Arguments passed to plot

Author(s)

Jared O'Connell

plot.hsmm.data *Plot function for hsmm data*

Description

Produces a plot of the observed sequences, and displays a coloured bar signifying the hidden states (if available)

Usage

```
## S3 method for class 'hsmm.data':
plot(x, ...)
```

Arguments

x A hsmm.data object
... Arguments passed to plot.ts

Author(s)

Jared O'Connell

See Also

addStates

Examples

```
J<-3
initial <- rep(1/J,J)
P <- matrix(c(.8, .5, .1, 0.05, .2, .5, .15, .3, .4), nrow=J)
b <- list(mu=c(-3, 0, 2), sigma=c(2, 1, .5))
model <- hmmspec(init=initial, trans=P, emission=b, f=dnorm.hsmm)

train <- simulate(model, nsim=300, seed=1234, r=rnorm.hsmm)
plot(train, xlim=c(0, 100))
```

predict.hmm

Prediction function for hmm

Description

Predicts the underlying state sequence for an observed sequence `x` given a `hmm` model

Usage

```
## S3 method for class 'hmm':
predict(object, x, method = "viterbi", ...)
```

Arguments

<code>object</code>	An object of class <code>hmm</code>
<code>x</code>	A vector or <code>data.frame</code> of observations
<code>method</code>	Prediction method (see details)
<code>...</code>	further arguments passed to or from other methods.

Details

If `method="viterbi"`, this technique applies the Viterbi algorithm for HMMs, producing the most likely sequence of states given the observed data. If `method="smoothed"`, then the individually most likely (or smoothed) state sequence is produced, along with a matrix with the respective probabilities for each state.

Value

Returns a `hsmm.data` object, suitable for plotting.

<code>x</code>	A vector or <code>data.frame</code> of observations
<code>s</code>	A vector containing the reconstructed state sequence
<code>N</code>	The lengths of each sequence
<code>p</code>	A matrix where the rows represent time steps and the columns are the probability for the respective state (only produced when <code>method="smoothed"</code>)

Author(s)

Jared O'Connell

References

Rabiner, L. (1989), A tutorial on hidden Markov models and selected applications in speech recognition, Proceedings of the IEEE, 77, 257-286.

See Also

hmm

Examples

```
##See examples in 'hmm'
```

predict.hsmm	<i>Prediction for hsmms</i>
--------------	-----------------------------

Description

Predicts the underlying state sequence for an observed sequence `x` given a `hsmm` model

Usage

```
## S3 method for class 'hsmm':
predict(object, x, method = "viterbi", ...)
```

Arguments

<code>object</code>	An object of type <code>hsmm</code>
<code>x</code>	A vector or dataframe of observations
<code>method</code>	Prediction method (see details)
<code>...</code>	further arguments passed to or from other methods.

Details

If `method="viterbi"`, this technique applies the Viterbi algorithm for HSMMS, producing the most likely sequence of states given the observed data. If `method="smoothed"`, then the individually most likely (or smoothed) state sequence is produced, along with a matrix with the respective probabilities for each state.

Value

Returns a `hsmm.data` object, suitable for plotting.

<code>x</code>	A vector or <code>data.frame</code> of observations
<code>s</code>	A vector containing the reconstructed state sequence
<code>N</code>	The lengths of each sequence
<code>p</code>	A matrix where the rows represent time steps and the columns are the probability for the respective state (only produced when <code>method="smoothed"</code>)

Author(s)

Jared O'Connell

References

Guedon, Y. (2003), Estimating hidden semi-Markov chains from discrete sequences, *Journal of Computational and Graphical Statistics*, Volume 12, Number 3, page 604-639 - 2003

See Also

`hsmm`

Examples

```
##See 'hsmm' for examples
```

```
print.hmm          Print method for hmm objects
```

Description

Prints the slots of a `hmm` object

Usage

```
## S3 method for class 'hmm':
print(x, ...)
```

Arguments

<code>x</code>	An object of type <code>hmm</code>
<code>...</code>	further arguments passed to or from other methods.

Author(s)

Jared O'Connell

`print.hmmspec` *Print function for hmmspec*

Description

Prints the parameters contained in the object

Usage

```
## S3 method for class 'hmmspec':  
print(x, ...)
```

Arguments

`x` An object of type `hmmspec`
`...` further arguments passed to or from other methods.

Author(s)

Jared O'Connell

`print.hsmmspec` *Print function for hsmmspec*

Description

Prints the parameters contained in the object

Usage

```
## S3 method for class 'hsmmspec':  
print(x, ...)
```

Arguments

`x` An object of type `hsmmspec`
`...` further arguments passed to or from other methods.

Author(s)

Jared O'Connell

`reproai`*Artificial insemination times for seven cows*

Description

This is an auxiliary data set to the `cows` data set containing times of artificial insemination for respective cows. Only the day of insemination was recorded so time of day is always midday.

Usage`reproai`**Format**

`reproai` is a dataframe with 12 rows and `id` being the cow's id and `days.from.calving` recording the number of days from calving when insemination occurred.

Source

Danish Cattle Research Centre

References

Peters, A. and Ball, P. (1995), "Reproduction in Cattle," 2nd ed.

`reprocows`*Reproductive data from seven dairy cows*

Description

This data set contains hourly observations on progesterone and an activity index at hourly intervals since calving on seven dairy cows.

Usage`reprocows`**Format**

`reprocows` is a data frame containing 13040 rows. `id` is the cow ID, `progesterone` is a measurement of the hormone in ng/L taken from a milk sample, `activity` is a relative measure of activity calculated from a pedometer.

There are a large number of missing values as progesterone is measured only at milking time (and at a farm manager's discretion). Missing values in activity occur due to hardware problems can occur with pedometers.

Source

Danish Cattle Research Centre

References

Peters, A. and Ball, P. (1995), "Reproduction in Cattle," 2nd ed.

Examples

```
data(reprocows)
data(reproai)
data(reproppa)
tm = 1600

J <- 3
init <- c(1,0,0)
trans <- matrix(c(0,0,0,1,0,1,0,1,0),nrow=J)
emis <- list(mu=c(0,2.5,0),sigma=c(1,1,1))

N <- as.numeric(table(reprocows$id))
train <- list(x=reprocows$activity,N=N)
class(train) <- "hsmm.data"
tmp <- gammafit(reproppa * 24)
M <- max(N)
d <- cbind(dgamma(1:M,shape=tmp$shape,scale=tmp$scale), #ppa sojourn directly estimated from
           dunif(1:M,4,30), #oestrus between 4 and 30 hours
           dunif(1:M,15*24,40*24)) #not-oestrus between 15 and 40 days

startval <- hsmmspec(init,trans,emis,list(d=d,type='gamma'),f=dnorm.hsmm)
h.activity <- hsmmfit(train,startval,mstep=mstep.norm,maxit=10,M=M,lock.transition=TRUE)
```

reproppa

Observed lengths of post-partum anoestrus for 73 dairy cows

Description

This data set contains the observed length of post-partum anoestrus (in days) for 73 dairy cattle.

Usage

```
reproppa
```

Format

reproppa a vector containing 73 integers.

Source

Danish Cattle Research Centre

References

Peters, A. and Ball, P. (1995), "Reproduction in Cattle," 2nd ed.

rmvnorm.hsmm	<i>Random number generation from a multivariate normal distributed emission distribution</i>
--------------	--

Description

This generates values from a multivariate normal distributed emission state j given parameters in `model`.

Usage

```
rmvnorm.hsmm(j, model)
```

Arguments

<code>j</code>	An integer representing the state
<code>model</code>	A <code>hmm</code> spec or <code>hsmm</code> spec object

Details

This is essentially a wrapper for `rnorm`. Users may build functions with the same arguments and return values so they can use their own custom emission distributions.

Value

A single value from the emission distribution.

Author(s)

Jared O'Connell

See Also

`dmvnorm.hsmm`, `mstep.mvnorm`

Examples

```
J<-2
initial <- rep(1/J,J)
P <- matrix(c(.3, .5, .7, .5), nrow=J)
b <- list(mu=list(c(-3,0),c(1,2)), sigma=list(diag(2),matrix(c(4,2,2,3), ncol=2)))
model <- hmm-spec(init=initial, trans=P, emission=b, f=dmvnorm.hsmm)
train <- simulate(model, nsim=300, seed=1234, r=rmvnorm.hsmm)
plot(train, xlim=c(0,100))
h1 = hmmfit(train,model,mstep=mstep.mvnorm)
```

rnorm.hsmm	<i>Random number generation from a normally distributed emission distribution</i>
------------	---

Description

This generates values from a normally distributed emission state `j` given parameters in `model`.

Usage

```
rnorm.hsmm(j, model)
```

Arguments

<code>j</code>	An integer representing the state
<code>model</code>	A <code>hmmspec</code> or <code>hsmmspec</code> object

Details

This is essentially a wrapper for `rnorm`. Users may build functions with the same arguments and return values so they can use their own custom emission distributions.

Value

A single value from the emission distribution.

Author(s)

Jared O'Connell

rpois.hsmm	<i>Random number generation from a Poisson distributed emission distribution</i>
------------	--

Description

This generates values from a Poisson distributed emission state `j` given parameters in `model`.

Usage

```
rpois.hsmm(j, model)
```

Arguments

<code>j</code>	An integer representing the state
<code>model</code>	A <code>hmmspec</code> or <code>hsmmspec</code> object

Details

This is essentially a wrapper for `rpois`. Users may build functions with the same arguments and return values so they can use their own custom emission distributions.

Value

A single value from the emission distribution.

Author(s)

Jared O'Connell

See Also

`mstep.pois`, `dpois.hsmm`

Examples

```
J<-3
initial <- rep(1/J,J)
P <- matrix(c(.8, .5, .1, 0.05, .2, .5, .15, .3, .4),nrow=J)
b <- list(lambda=c(1,3,6))
model <- hmmspec(init=initial, trans=P, emission=b,f=dpois.hsmm)
model
train <- simulate(model, nsim=300, seed=1234, r=rpois.hsmm)
plot(train,xlim=c(0,100))
h1 = hmmfit(train,model,mstep=mstep.pois)
```

sim.mc

Markov chain simulation

Description

Simulates a Markov chain

Usage

```
sim.mc(init, transition, N)
```

Arguments

<code>init</code>	The distribution of states at the first time step
<code>transition</code>	The transition probability matrix of the Markov chain
<code>N</code>	The number of observations to simulate

Value

A vector of integers representing the state sequence.

Author(s)

Jared O'Connell

Examples

```
p <- matrix(c(.1, .3, .6, rep(1/3, 3), 0, .5, .5), ncol=3, byrow=TRUE)
init <- rep(1/3, 3)
sim.mc(init, p, 10)
```

simulate.hmmspec *Simulation of hidden Markov models*

Description

Simulates data from a hidden Markov model

Usage

```
## S3 method for class 'hmmspec':
simulate(object, nsim, seed = NULL, r=NULL, ...)
```

Arguments

object	A hmmspec object
nsim	An integer or vector of integers (for multiple sequences) specifying the length of the sequence(s)
seed	seed for the random number generator
r	The function used to generate observations from the emission distribution
...	further arguments passed to or from other methods.

Details

If `nsim` is a single integer then a HMM of that length is produced. If `nsim` is a vector of integers, then `length(nsim)` sequences are generated with respective lengths.

Value

An object of class `hmmdata`

x	A vector of length <code>sum(N)</code> - the sequence(s) of observed values
s	A vector of length <code>sum(N)</code> - the sequence(s) of hidden states
N	A vector of the length of each observation sequence (used to segment x and s)

Author(s)

Jared O'Connell

References

Rabiner, L. (1989), A tutorial on hidden Markov models and selected applications in speech recognition, Proceedings of the IEEE, 77, 257-286.

See Also

hsmmspec, simulate.hmm, hmm, predict.hmm

Examples

```
J<-3
initial <- rep(1/J,J)
P <- matrix(c(.8,.5,.1,0.05,.2,.5,.15,.3,.4),nrow=J)
b <- list(mu=c(-3,0,2),sigma=c(2,1,.5))
model <- hsmmspec(init=initial, trans=P, emission=b,f=dnorm.hsmm)
train <- simulate(model, nsim=100, seed=1234, r=rnorm.hsmm)
plot(train)
```

simulate.hsmmspec *Simulation for HSMMs*

Description

Simulates values for a specified hidden semi-Markov model

Usage

```
## S3 method for class 'hsmmspec':
simulate(object, nsim, seed = NULL, r=NULL, ...)
```

Arguments

object	A hsmmspec object
nsim	An integer or vector of integers (for multiple sequences) specifying the length of the sequence(s)
seed	seed for the random number generator
r	The function used to generate observations from the emission distribution
...	further arguments passed to or from other methods.

Details

If `nsim` is a single integer then a HSMM of that length is produced. If `nsim` is a vector of integers, then `length(nsim)` sequences are generated with respective lengths.

Value

An object of class `hmmdata`

<code>x</code>	A vector of length <code>sum(N)</code> - the sequence(s) of observed values
<code>s</code>	A vector of length <code>sum(N)</code> - the sequence(s) of hidden states
<code>N</code>	A vector of the length of each observation sequence (used to segment <code>x</code> and <code>s</code>)

Author(s)

Jared O'Connell

References

Guedon, Y. (2003), Estimating hidden semi-Markov chains from discrete sequences, *Journal of Computational and Graphical Statistics*, Volume 12, Number 3, page 604-639 - 2003

See Also

`hsmm`, `simulate.hsmm`, `predict.hsmm`

summary.hmm

Summary method for hmm objects

Description

Prints the estimated parameters of a `hmm` object

Usage

```
## S3 method for class 'hmm':
summary(object, ...)
```

Arguments

<code>object</code>	A <code>hmm</code> object
<code>...</code>	further arguments passed to or from other methods.

Value

An object of class `'summary.hmm'`

Author(s)

Jared O'Connell

summary.hsmm *Summary function for hsmm*

Description

Returns a summary object for a hsmm object

Usage

```
## S3 method for class 'hsmm':  
summary(object, ...)
```

Arguments

object	An object of type hsmm
...	further arguments passed to or from other methods.

Author(s)

Jared O'Connell

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