Package ‘depmixS4’

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depmixS4-package

depmixS4 provides classes for specifying and fitting hidden Markov models

Description

depmixS4 is a framework for specifying and fitting dependent mixture models, otherwise known as hidden or latent Markov models. Optimization is done with the EM algorithm or optionally with Rdonlp2 when (general linear (in-)equality) constraints on the parameters need to be incorporated. Models can be fitted on (multiple) sets of observations. The response densities for each state may be chosen from the GLM family, or a multinomial. User defined response densities are easy to add; for the latter an example is given for the ex-gauss distribution as well as the multivariate normal distribution.

Mixture or latent class (regression) models can also be fitted; these are the limit case in which the length of observed time series is 1 for all cases.

Details

Model fitting is done in two steps; first, models are specified through the `depmix` function (or the `mix` function for mixture and latent class models), which both use standard `glm` style arguments to specify the observed distributions; second, the model needs to be fitted by using the `fit` function; imposing constraints is done through the `fit` function. Standard output includes the optimized parameters and the posterior densities for the states and the optimal state sequence.
For full control and the possibility to add new response distributions, check the `makeDepmix` help page.

**Author(s)**

Ingmar Visser & Maarten Speekenbrink  
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**References**


**See Also**

`depmix`, `fit`

**Examples**

```r
# create a 2 state model with one continuous and one binary response
data(speed)
mod <- depmix(list(rt~1,corr~1),data=speed,nstates=2,family=list(gaussian(),multinomial()))
# print the model, formulae and parameter values (ie the starting values)
mod
```

---

### Balance Scale Data

**Description**

Balance scale data of four distance items from 779 participants; participants ages are included.

**Usage**

```r
data(balance)
```

**Format**

A data frame with 779 observations on the following variables. The full dataset is described and analyzed extensively in Jansen & Van der Maas (2002). The trichotomous data are left, balance, right. The dichotomous version of the data is scored correct, incorrect.

- **sex** Participants sex.
- **agedays** Age in days.
age  Age in years.
t1  Trichotomously scored distance item.
t2  Trichotomously scored distance item.
t3  Trichotomously scored distance item.
t4  Trichotomously scored distance item.
d1  Dichotomously scored distance item.
d2  Dichotomously scored distance item.
d3  Dichotomously scored distance item.
d4  Dichotomously scored distance item.

Source


Examples

data(balance)

depmix
Dependent Mixture Model Specification

Description

depmix creates an object of class depmix, a dependent mixture model, otherwise known as hidden Markov model. For a short description of the package see depmixS4. See the vignette for an introduction to hidden Markov models and the package.

Usage

depmix(response, data=NULL, nstates, transition=~1, family=gaussian(), prior=~1, inidata=NULL, respstart=NULL, trstart=NULL, instart=NULL, ntimes=NULL,...)

Arguments

response  The response to be modeled; either a formula or a list of formulae (in the multivariate case); this interfaces to the glm and other distributions. See ‘Details’.
data     An optional data.frame to interpret the variables in the response and transition arguments.
nstates  The number of states of the model.
transition  A one-sided formula specifying the model for the transitions. See ‘Details’.
family
A family argument for the response. This must be a list of family's if the response is multivariate.

prior
A one-sided formula specifying the density for the prior or initial state probabilities.

initdata
An optional data.frame to interpret the variables occurring in prior. The number of rows of this data.frame must be equal to the number of cases being modeled, length(ntimes). See 'Details'.

respstart
Starting values for the parameters of the response models.

trstart
Starting values for the parameters of the transition models.

instart
Starting values for the parameters of the prior or initial state probability model.

ntimes
A vector specifying the lengths of individual, i.e. independent, time series. If not specified, the responses are assumed to form a single time series, i.e. ntimes=nrow(data). If the data argument has an attribute ntimes, then this is used. The first example in fit uses this argument.

... Not used currently.

details
The function depmix creates an S4 object of class depmix, which needs to be fitted using fit to optimize the parameters.

The response model(s) are by default created by call(s) to GLMresponse using the formula and the family arguments, the latter specifying the error distribution. See GLMresponse for possible values of the family argument for glm-type responses (ie a subset of the glm family options, and the multinomial). Alternative response distributions are specified by using the makeDepmix function. Its help page has examples of specifying a model with a multivariate normal response, as well as an example of adding a user-defined response model, in this case for the ex-gauss distribution.

If response is a list of formulae, the response's are assumed to be independent conditional on the latent state.

The transitions are modeled as a multinomial logistic model for each state. Hence, the transition matrix can be modeled using time-varying covariates. The prior density is also modeled as a multinomial logistic. Both of these models are created by calls to transInit.

Starting values for the initial, transition, and response models may be provided by their respective arguments. NB: note that the starting values for the initial and transition models as well as of the multinomial logit response models are interpreted as probabilities, and internally converted to multinomial logit parameters. The order in which parameters must be provided can be easily studied by using the setpars and getpars functions.

Linear constraints on parameters can be provided as argument to the fit function.

The print function prints the formulae for the response, transition and prior models along with their parameter values.

Missing values are allowed in the data, but missing values in the covariates lead to errors.

Value

depmix returns an object of class depmix which has the following slots:
response  A list of a list of response models; the first index runs over states; the second index runs over the independent responses in case a multivariate response is provided.

transition  A list of transInit models, ie multinomial logistic models with length the number of states.

prior  A multinomial logistic model for the initial state probabilities.

dens,trDens,init  See depmix-class help for details. For internal use.

stationary  Logical indicating whether the transitions are time-dependent or not; for internal use.

ntimes  A vector containing the lengths of independent time series.

nstates  The number of states of the model.

nresp  The number of independent responses.

npars  The total number of parameters of the model. Note: this is not the degrees of freedom because there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior probabilities.

Note

Models are not fitted; the return value of depmix is a model specification without optimized parameter values. Use the fit function to optimize parameters, and to specify additional constraints.

Author(s)

Ingmar Visser & Maarten Speekenbrink

References


See Also

fit, transInit, GLMresponse, depmix-methods for accessor functions to depmix objects.

For full control see the makeDepmix help page and its example section for the possibility to add user-defined response distributions.

Examples

```r
# create a 2 state model with one continuous and one binary response
# ntimes is used to specify the lengths of 3 separate series
data(speed)
mod <- depmix(list(rt~1,corr~1),data=speed,nstates=2,
  family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
```
# print the model, formulae and parameter values
mod
set.seed(1)
# fit the model by calling fit
fm <- fit(mod)

# Volatility of S & P 500 returns
# (thanks to Chen Haibo for providing this example)
data(sp500)

# fit some models
msp <- depmix(logret~1,nstates=2,data=sp500)
set.seed(1)
fmsp <- fit(msp)

# plot posterior state sequence for the 2-state model
plot(ts(posterior(fmsp, type="smoothing")[,1], start=c(1950,2),deltat=1/12),ylab="probability",
main="Posterior probability of state 1 (volatile, negative markets).",
frame=FALSE)

## Not run:
# this creates data with a single change point with Poisson data
set.seed(3)
y1 <- rpois(50,1)
y2 <- rpois(50,2)
ydf <- data.frame(y=c(y1,y2))

# fit models with 1 to 3 states
m1 <- depmix(y~1,ns=1,family=poisson(),data=ydf)
set.seed(1)
fm1 <- fit(m1)
m2 <- depmix(y~1,ns=2,family=poisson(),data=ydf)
set.seed(1)
fm2 <- fit(m2)
m3 <- depmix(y~1,ns=3,family=poisson(),data=ydf)
set.seed(1)
fm3 <- fit(m3,em=em.control(maxit=500))

# plot the BICs to select the proper model
plot(1:3,c(BIC(fm1),BIC(fm2),BIC(fm3)),ty="b")

## End(Not run)

## Not run:
# similar to the binomial model, data may also be entered in
# multi-column format where the n for each row can be different
dt <- data.frame(y1=c(0,1,1,2,4,5),y2=c(1,0,1,0,1,0),y3=c(4,4,3,2,1,1))
# specify a mixture model ...
m2 <- mix(cbind(y1,y2,y3)~1,data=dt,ns=2,family=multinomial("identity"))
set.seed(1)
fm2 <- fit(m2)
# ... or dependent mixture model
dm2 <- depmix(cbind(y1, y2, y3)~1, data=dt, ns=2, family=multinomial("identity"))
set.seed(1)
fm2 <- fit(dm2)

## End(Not run)

depmix-class  
Class "depmix"

Description
A depmix model.

Slots
response: List of list of response objects.
transition: List of transInit objects.
prior: transInit object.
dens: Array of dimension sum(ntimes)*nresp*nstates providing the densities of the observed responses for each state.
trDens: Array of dimension sum(ntimes)*nstates providing the probability of a state transition depending on the predictors.
init: Array of dimension length(ntimes)*nstates with the current predictions for the initial state probabilities.
homogeneous: Logical indicating whether the transitions are time-dependent or not; for internal use.
ntimes: A vector containing the lengths of independent time series; if data is provided, sum(ntimes) must be equal to nrow(data).
nstates: The number of states of the model.
nresp: The number of independent responses.
npars: The total number of parameters of the model. This is not the degrees of freedom, ie there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior.

Accessor Functions
The following functions should be used for accessing the corresponding slots:
npar: The number of parameters of the model.
nresp: The number of responses.
nstates: The number of states.
n-times: The vector of independent time series lengths.
Author(s)
Ingmar Visser & Maarten Speekenbrink

depmix-methods
'depmix' and 'mix' methods.

Description
Various methods for depmix and mix objects.

Usage

```r
## S4 method for signature 'depmix'
logLik(object, method=c("fb","lystig","classification"), na.allow=TRUE)
## S4 method for signature 'mix'
logLik(object, method=c("fb","lystig","classification"), na.allow=TRUE)

## S4 method for signature 'depmix.fitted.classLik'
logLik(object, method=c("classification","fb","lystig"), na.allow=TRUE)
## S4 method for signature 'mix.fitted.classLik'
logLik(object, method=c("classification","fb","lystig"), na.allow=TRUE)

## S4 method for signature 'depmix'
nobs(object, ...)
## S4 method for signature 'mix'
nobs(object, ...)

## S4 method for signature 'depmix'
npar(object)
## S4 method for signature 'mix'
npar(object)

## S4 method for signature 'depmix'
freepars(object)
## S4 method for signature 'mix'
freepars(object)

## S4 method for signature 'depmix'
setpars(object, values, which="pars", ...)
## S4 method for signature 'mix'
setpars(object, values, which="pars", ...)

## S4 method for signature 'depmix'
getpars(object, which="pars", ...)
## S4 method for signature 'mix'
```
getpars(object, which="pars", ...)

## S4 method for signature 'depmix'
getmodel(object, which="response", state=1, number=1)

Arguments

- **object**: A depmix or mix object.
- **values**: To be used in setpars to set new parameter values; see the example.
- **method**: The log likelihood can be computed by either the forward backward algorithm (Rabiner, 1989), or by the method of Lystig and Hughes, 2002. The former is the default and implemented in a fast C routine. The forward-backward routine also computes the state and transition smoothed probabilities, which are not directly necessary for the log likelihood. Those smoothed variables, and the forward and backward variables are accessible through the forwardbackward function. When method="classification", the classification likelihood is computed, which is the likelihood of the data assuming the state sequence is known and equal to the maximum a posteriori state sequence. The MAP state sequence is available through the viterbi function. The classification likelihood is computed by default when calling the logLik method on a a model fitted by maximising the classification likelihood.
- **na.allow**: Allow missing observations? When set to FALSE, the logLik method will return NA in the presence of missing observations. When set to TRUE, missing values will be ignored when computing the likelihood. When observations are partly missing (when a multivariate observation has missing values on only some of its dimensionis), this may give unexpected results.
- **which**: getpars function: The default "pars" returns a vector of all parameters of a depmix object; the alternative value "fixed" returns a logical vector of the same length indicating which parameters are fixed. The setpars functions sets parameters to new values; setpars also recomputes the dens, trans and init slots of depmix objects. Note that the getpars and setpars functions for depmix objects simply call the functions of the same name for the response and transition models.
- **state**: In getmodel this determines the submodel to be returned (together with number in the case of response models); when which="transition", getmodel returns the transition submodel for state=state.
- **number**: In getmodel this determines the "response" model to be returned from state state.
- **...**: Not used currently.

Value

- **logLik**: returns a logLik object with attributes df and nobs.
depmix.fitted-class

nobs returns the number of observations (used in computing the BIC).
npar returns the number of parameters of a model.
freepars returns the number of non-redundant parameters.
setpars returns a (dep-)mix object with new parameter values.
getpars returns a vector with the current parameter values.
getmodel returns a submodel of a (dep-)mix model; the prior model, one of the transition models (determined by argument state) or one of the response models (determined by arguments state and number).

Author(s)
Ingmar Visser & Maarten Speekenbrink

Examples

# create a 2 state model with one continuous and one binary response
data(speed)
mod <- depmix(list(rt~1,corr~1),data=speed,nstates=2,family=list(gaussian(),multinomial()))

getmodel(mod,"response",2,1)
getmodel(mod,"prior")

# get the loglikelihood of the model
logLik(mod)

# to see the ordering of parameters to use in setpars
mod <- setpars(mod, value=1:npar(mod))
mod

# to see which parameters are fixed (by default only baseline parameters in
# the multinomial logistic models for the transition models and the initial
# state probabilities model)
mod <- setpars(mod, getpars(mod,which="fixed"))
mod


---

depmix.fitted-class Class "depmix.fitted" (and "depmix.fitted.classLik")

Description
A fitted depmix model.
Slots

A depmix.fitted object is a depmix object with three additional slots, here is the complete list:

response: List of list of response objects.
transition List of transInit objects.
prior: transInit object.
dens: Array of dimension sum(ntimes)*nresp*nstates providing the densities of the observed responses for each state.
trDens: Array of dimension sum(ntimes)*nstates providing the probability of a state transition depending on the predictors.
init: Array of dimension length(ntimes)*nstates with the current predictions for the initial state probabilities.
stationary: Logical indicating whether the transitions are time-dependent or not; for internal use.
ntimes: A vector containing the lengths of independent time series; if data is provided, sum(ntimes) must be equal to nrow(data).
nstates: The number of states of the model.
nresp: The number of independent responses.
npars: The total number of parameters of the model. This is not the degrees of freedom, ie there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior.
message: This provides some information on convergence, either from the EM algorithm or from Rdonlp2.
conMat: The linear constraint matrix, which has zero rows if there were no constraints.
lin.lower The lower bounds on the linear constraints.
lin.upper The upper bounds on the linear constraints.
posterior: Posterior (Viterbi) state sequence.

Details

The print function shows some convergence information, and the summary method shows the parameter estimates.

Extends

depmix.fitted extends the "depmix" class directly. depmix.fitted.classLik is similar to depmix.fitted, the only difference being that the model is fitted by maximising the classification likelihood.

Author(s)

Ingmar Visser & Maarten Speekenbrink
**Description**

A `depmix.sim` model. The `depmix.sim` class directly extends the `depmix` class, and has an additional slot for the true states. A `depmix.sim` model can be generated by `simulate(mod,...)`, where `mod` is a `depmix` model.

**Slots**

- **response**: List of list of response objects.
- **transition**: List of `transInit` objects.
- **prior**: `transInit` object.
- **dens**: Array of dimension `sum(ntimes)*nresp*nstates` providing the densities of the observed responses for each state.
- **trDens**: Array of dimension `sum(ntimes)*nstates` providing the probability of a state transition depending on the predictors.
- **init**: Array of dimension `length(ntimes)*nstates` with the current predictions for the initial state probabilities.
- **homogeneous**: Logical indicating whether the transitions are time-dependent or not; for internal use.
- **ntimes**: A vector containing the lengths of independent time series; if data is provided, `sum(ntimes)` must be equal to `nrow(data)`.
- **nstates**: The number of states of the model.
- **nresp**: The number of independent responses.
- **npars**: The total number of parameters of the model. This is not the degrees of freedom, i.e., there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior.
- **states**: A matrix with the true states.

**Accessor Functions**

The following functions should be used for accessing the corresponding slots:

- **npar**: The number of parameters of the model.
- **nresp**: The number of responses.
- **nstates**: The number of states.
- **ntimes**: The vector of independent time series lengths.

**Author(s)**

Maarten Speekenbrink & Ingmar Visser
em.control

Control parameters for the EM algorithm

Description

Set control parameters for the EM algorithm.

Usage

em.control(maxit = 500, tol = 1e-08, crit = c("relative","absolute"),
random.start = TRUE, classification = c("soft","hard"))

Arguments

maxit The maximum number of iterations.
tol The tolerance level for convergence. See Details.
crit Sets the convergence criterion to "relative" or "absolute" change of the log-
likelihood. See Details.
random.start This is used for a (limited) random initialization of the parameters. See Details.
classification Type of classification to states used. See Details.

details

The argument crit sets the convergence criterion to either the relative change in the log-likelihood
or the absolute change in the log-likelihood. The relative likelihood criterion (the default) assumes
convergence on iteration $i$ when
\[
\frac{\log L(i) - \log L(i-1)}{\log L(i-1)} < \text{tol}.
\]
The absolute likelihood criterion assumes convergence on iteration $i$ when
\[
\log L(i) - \log L(i-1) < \text{tol}.
\]
Use crit="absolute" to invoke the latter convergence criterion. Note that in that case, optimal values of the tolerance parameter
tol scale with the value of the log-likelihood (and these are not changed automagically).

Argument random.start This is used for a (limited) random initialization of the parameters. In
particular, if random.start=TRUE, the (posterior) state probabilities are randomized at iteration 0
(using a uniform distribution), i.e. the $\gamma$ variables (Rabiner, 1989) are sampled from the Dirichlet
distribution with a (currently fixed) value of $\alpha = 0.1$; this results in values for each row of $\gamma$ that are
quite close to zero and one; note that when these values are chosen at zero and one, the initialization
is similar to that used in kmeans. Random initialization is useful when no initial parameters can be
given to distinguish between the states. It is also useful for repeated estimation from different
starting values.

Argument classification is used to choose either soft (default) or hard classification of observations to states. When using soft classification, observations are assigned to states with a weight
equal to the posterior probability of the state. When using hard classification, observations are
assigned to states according to the maximum a posteriori (MAP) states (i.e., each observation is
assigned to one state, which is determined by the Viterbi algorithm in the case of depmixx models).
As a result, the EM algorithm will find a local maximum of the classification likelihood (Celeux
& Govaert, 1992). Warning: hard classification is an experimental feature, especially for hidden Markov models, and its use is currently not advised.

Value

em.control returns a list of the control parameters.

Author(s)

Maarten Speekenbrink & Ingmar Visser

References


Examples

# using "hard" assignment of observations to the states, we can maximise the classification likelihood instead of the usual marginal likelihood
data(speed)
mod <- depmix(list(rt~1,corr~1),data=speed,nstates=2,
               family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
set.seed(1)
# fit the model by calling fit
fmod <- fit(mod,emcontrol=em.control(classification="hard"))
# can get rather different solutions with different starting values...
set.seed(3)
fmod2 <- fit(mod,emcontrol=em.control(classification="hard"))

fit

Fit 'depmix' or 'mix' models

Description

fit optimizes parameters of depmix or mix models, optionally subject to general linear (in)equality constraints.

Usage

## S4 method for signature 'mix'
fit(object, fixed=NULL, equal=NULL, conrows=NULL, conrows.upper=NULL, conrows.lower=NULL,
    method=NULL, verbose=FALSE, emcontrol=em.control(),
## S4 method for signature 'mix.fitted'
summary(object, which="all")

## S4 method for signature 'depmix.fitted'
summary(object, which="all")

### Arguments

- **object**: An object of class (dep-)mix.
- **fixed**: Vector of mode logical indicating which parameters should be fixed.
- **equal**: Vector indicating equality constraints; see Details.
- **conrows**: Rows of a general linear constraint matrix; see Details.
- **conrows.upper**, **conrows.lower**: Upper and lower bounds for the linear constraints; see Details.
- **method**: The optimization method; mostly determined by constraints.
- **verbose**: Should optimization information be displayed on screen?
- **emcontrol**: Named list with control parameters for the EM algorithm (see `em.control`).
- **solnpcontrol**: Control parameters passed to the 'rsolnp' optimizer; see `solnp` for explanation and defaults used there.
- **donlpcontrol**: Control parameters passed to 'donlp' optimizer; see `?donlp2Control` for explanation and defaults used there; this can be used to tweak optimization but note that extra output is not returned.
- **which**: Should summaries be provided for "all" submodels? Options are "prior", "response", and for fitted depmix models also "transition".
- ... Further arguments passed on to the optimization methods.

### Details

Models are fitted by the EM algorithm if there are no constraints on the parameters. Aspects of the EM algorithm can be controlled through the `emcontrol` argument; see details in `em.control`. Otherwise the general optimizers `solnp`, the default (from package Rsolnp) or `donlp2` (from package Rdonlp2) are used which handle general linear (in-)equality constraints. These optimizers are selected by setting `method='rsolnp'` or `method='donlp'` respectively.

Three types of constraints can be specified on the parameters: fixed, equality, and general linear (in-)equality constraints. Constraint vectors should be of length `npar(object)`; note that this hence includes redundant parameters such as the base category parameter in multinomial logistic models which is always fixed at zero. See help on `getpars` and `setpars` about the ordering of parameters.

The `equal` argument is used to specify equality constraints: parameters that get the same integer number in this vector are estimated to be equal. Any integers can be used in this way except 0 and 1, which indicate fixed and free parameters, respectively.
Using solnp (or donlp2), a Newton-Raphson scheme is employed to estimate parameters subject to linear constraints by imposing:

\[ bl \leq A \cdot x \leq bu, \]

where \( x \) is the parameter vector, \( bl \) is a vector of lower bounds, \( bu \) is a vector of upper bounds, and \( A \) is the constraint matrix.

The conrows argument is used to specify rows of \( A \) directly, and the conrows.lower and conrows.upper arguments to specify the bounds on the constraints. conrows must be a matrix of npar(object) columns and one row for each constraint (a vector in the case of a single constraint). Examples of these three ways of constraining parameters are provided below.

Note that when specifying constraints that these should respect the fixed constraints inherent in e.g. the multinomial logit models for the initial and transition probabilities. For example, the baseline category coefficient in a multinomial logit model is fixed on zero.

llratio performs a log-likelihood ratio test on two fitted models; the first object should have the largest degrees of freedom (find out by using freepars).

Value

fit returns an object of class \texttt{depmix.fitted} which contains the original depmix object, and further has slots:

message: Convergence information.

conMat: The constraint matrix \( A \), see Details.

posterior: The posterior state sequence (computed with the viterbi algorithm), and the posterior probabilities (delta probabilities in Rabiner, 1989, notation).

The print method shows the message along with the likelihood and AIC and BIC; the summary method prints the parameter estimates.

Posterior densities and the viterbi state sequence can be accessed through posterior.

As fitted models are depmixS4 models, they can be used as starting values for new fits, for example with constraints added. Note that when refitting already fitted models, the constraints, if any, are not added automatically, they have to be added again.

Author(s)

Ingmar Visser & Maarten Speekenbrink

References

Some of the below models for the speed data are reported in:

Examples

data(speed)

# 2-state model on rt and corr from speed data set
# with Pacc as covariate on the transition matrix
# ntimes is used to specify the lengths of 3 separate time-series
mod1 <- depmix(list(rt~1,corr~1),data=speed,transition=~Pacc,nstates=2,
                family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
# fit the model
set.seed(3)
fmod1 <- fit(mod1)
fmod1 # to see the logLik and optimization information
# to see the parameters
summary(fmod1)

# to obtain the posterior most likely state sequence, as computed by the
# Viterbi algorithm
pst_global <- posterior(fmod1, type = "global")
# local decoding provides a different method for state classification:
pst_local <- posterior(fmod1,type="local")
identical(pst_global, pst_local)
# smoothing probabilities are used for local decoding, and may be used as
# easily interpretable posterior state probabilities
pst_prob <- posterior(fmod1, type = "smoothing")

# testing viterbi states for new data
df <- data.frame(corr=c(1,0,1),rt=c(6.4,5.5,5.3),Pacc=c(0.6,0.1,0.1))
# define model with new data like above
modNew <- depmix(list(rt~1,corr~1),data=df,transition=~Pacc,nstates=2,
                family=list(gaussian(),multinomial("identity")))
# get parameters from estimated model
modNew <- setpars(modNew,getpars(fmod1))
# check the state sequence and probabilities
pst_new <- posterior(modNew, type="global")

# same model, now with missing data
## Not run:
speed[2,1] <- NA
speed[3,2] <- NA

# 2-state model on rt and corr from speed data set
# with Pacc as covariate on the transition matrix
# ntimes is used to specify the lengths of 3 separate series
mod1ms <- depmix(list(rt~1,corr~1),data=speed,transition=~Pacc,nstates=2,
                 family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
# fit the model
set.seed(3)
fmod1ms <- fit(mod1ms)
## End(Not run)
# instead of the normal likelihood, we can also maximise the "classification" likelihood
# this uses the maximum a posteriori state sequence to assign observations to states
# and to compute initial and transition probabilities.

fmod1b <- fit(mod1, emcontrol = em.control(classification = "hard"))
fmod1b # to see the logLik and optimization information

# FIX SOME PARAMETERS

# get the starting values of this model to the optimized
# values of the previously fitted model to speed optimization

pars <- c(unlist(getpars(fmod1)))

# constrain the initial state probs to be 0 and 1
# also constrain the guessing probs to be 0.5 and 0.5
# (ie the probabilities of corr in state 1)
# change the ones that we want to constrain
pars[1] = 0
pars[2] = 1 # this means the process will always start in state 2
pars[13] = 0.5
pars[14] = 0.5 # the corr parameters are now both 0.5
mod2 <- setpars(mod1, pars)

# fix the parameters by setting:
free <- c(0, 0, rep(c(0, 1), 4), 1, 1, 0, 0, 1, 1, 1, 1)
# fit the model
fmod2 <- fit(mod2, fixed = !free)

# likelihood ratio insignificant, hence fmod2 better than fmod1
llratio(fmod1, fmod2)

# ADDING SOME GENERAL LINEAR CONSTRAINTS

# set the starting values of this model to the optimized
# values of the previously fitted model to speed optimization

## Not run:

pars <- c(unlist(getpars(fmod2)))
mod3 <- setpars(mod2, pars)

# start with fixed and free parameters
conpat <- c(0, 0, rep(c(0, 1), 4), 1, 1, 0, 0, 1, 1, 1, 1)
# constrain the beta's on the transition parameters to be equal
fmod3 <- fit(mod3, equal = conpat)
llratio(fmod2, fmod3)

# above constraints can also be specified using the conrows argument as follows
conr <- matrix(0, 2, 18)
# parameters 4 and 8 have to be equal, otherwise stated, their difference should be zero,
# and similarly for parameters 6 & 10
conr[1, 4] <- 1
conr[1, 8] <- -1
conr[2, 6] <- 1
conr[2, 10] <- -1

# note here that we use the fitted model fmod2 as that has appropriate
# starting values
fmod3b <- fit(mod3, conrows=conr, fixed=!free) # using free defined above

## End(Not run)

data(balance)
# four binary items on the balance scale task
mod4 <- mix(list(d1~1, d2~1, d3~1, d4~1), data=balance, nstates=2,
family=list(multinomial("identity"), multinomial("identity"),
multinomial("identity"), multinomial("identity")))

set.seed(1)
fmod4 <- fit(mod4)

## Not run:
# add age as covariate on class membership by using the prior argument
mod5 <- mix(list(d1~1, d2~1, d3~1, d4~1), data=balance, nstates=2,
family=list(multinomial("identity"), multinomial("identity"),
multinomial("identity"), multinomial("identity")),
prior=~age, initdata=balance)

set.seed(1)
fmod5 <- fit(mod5)

# check the likelihood ratio; adding age significantly improves the goodness-of-fit
llratio(fmod5, fmod4)

## End(Not run)

---

**formatperc**

*Format percentage for level in printing confidence interval*

**Description**

See title.
Usage

\texttt{formatperc(x,digits)}

Arguments

\begin{itemize}
  \item \texttt{x} \hspace{1cm} \text{a vector of probabilities to be formatted as percentages.}
  \item \texttt{digits} \hspace{1cm} \text{the number of required digits in the percentages.}
\end{itemize}

Author(s)

Ingmar Visser

\begin{longtable}{ll}
\texttt{forwardbackward} & \textit{Forward and backward variables} \\
\end{longtable}

Description

Compute the forward and backward variables of a \texttt{depmix} object.

Usage

\begin{verbatim}
## S4 method for signature 'mix'
forwardbackward(object, return.all=TRUE, useC=TRUE, ...)
\end{verbatim}

Arguments

\begin{itemize}
  \item \texttt{object} \hspace{1cm} \text{A \texttt{depmix} object.}
  \item \texttt{return.all} \hspace{1cm} \text{If FALSE, only gamma and xi and the log likelihood are returned (which are the only variables needed in using EM).}
  \item \texttt{useC} \hspace{1cm} \text{Flag used to set whether the C-code is used to compute the forward, backward, gamma and xi variables or not; the R-code is basically obsolete (but retained for now for debugging purposes).}
  \item \ldots \hspace{1cm} \text{Not currently used.}
\end{itemize}

Value

\texttt{forwardbackward} returns a list of the following (the variables are named after the notation from Rabiner, 1989):

\begin{itemize}
  \item \texttt{alpha} \hspace{1cm} \text{The forward variables.}
  \item \texttt{beta} \hspace{1cm} \text{The backward variables.}
  \item \texttt{gamma} \hspace{1cm} \text{The smoothed state probabilities.}
  \item \texttt{xi} \hspace{1cm} \text{The smoothed transition probabilities.}
\end{itemize}
GLMresponse

Methods for creating depmix response models

Description

Create GLMresponse objects for depmix models using formulae and family objects.

Usage

GLMresponse(formula, data=NULL, family=gaussian(), pstart=NULL, fixed=NULL, prob=TRUE, ...)

## S4 method for signature 'response'
getdf(object)
GLMresponse

Arguments

- **formula**: A model formula.
- **data**: An optional data.frame to interpret the variables from the formula argument in.
- **family**: A family object;
- **pstart**: Starting values for the coefficients and other parameters, e.g. the standard deviation for the gaussian() family.
- **fixed**: Logical vector indicating which parameters are to be fixed.
- **prob**: Logical indicating whether the starting values for multinomial() family models are probabilities or logistic parameters (see details).
- **object**: Object of class response.
- **...**: Not used currently.

Details

GLMresponse is the default driver for specifying response distributions of depmix models. It uses the familiar formula interface from glm to specify how responses depend on covariates/predictors.

Currently available options for the family argument are binomial, gaussian, poisson, Gamma, and multinomial. Except for the latter option, the GLMresponse model is an interface to the glm functions of which the functionality is used: predict, fit and density functions.

The multinomial model takes as link functions mlogit, the default, and then uses functionality from the nnet package to fit multinomial logistic models; using mlogit as link allows only n=1 models to be specified, i.e. a single observation for each occasion; it also takes identity as a link function. The latter is typically faster and is hence preferred when no covariates are present.

See the responses help page for examples.

Value

GLMresponse returns an object of class GLMresponse which extends the response-class.

getdf returns the number of free parameters of a response model.

Author(s)

Ingmar Visser & Maarten Speekenbrink

See Also

makeDepmix has an example of specifying a model with a multivariate normal response and an example of how to add a user-defined response model, in particular an ex-gauss distribution used for the speed data.
llratio \hspace{1cm} *Log likelihood ratio test on two fitted models*

**Description**

Performs a log likelihood ratio test on two fitted depmix models.

**Usage**

\[ \text{llratio}(\text{basemodel, constrainedmodel, ...}) \]

**Arguments**

- `basemodel`: Fitted model with a `logLik` method.
- `constrainedmodel`: Fitted model with a `logLik` method.
- `...`: Not currently used.

**Details**

See the `fit` help page for an example.

**Value**

`llratio` returns an object of class `llratio` which has slots:

- `value`: Minus twice the loglikelihood difference.
- `df`: The degrees of freedom, i.e., the difference in number of freely estimated parameters between the models.

The print method shows the value, the degrees of freedom and the corresponding p-value under the chisquared distribution.

**Author(s)**

Ingmar Visser
Description

makeDepmix creates an object of class `depmix`. This function is meant for full control, e.g. specifying each response model and the transition and prior models 'by hand'. For the default easier specification of models, please see `depmix`. This function is meant for specifying one's own response models.

Usage

```r
makeDepmix(response, transition, prior, ntimes = NULL, homogeneous = TRUE, stationary = NULL, ...)

makeMix(response, prior, ...)
```

Arguments

- `response`: A two-dimensional list of response models. See 'Details'.
- `transition`: A list of transition models, each created by a call to `transInit`. The length of this list should be the number of states of the model.
- `prior`: The initial state probabilities model; created through a call to `transInit`.
- `ntimes`: A vector specifying the lengths of individual, i.e., independent, time series. If not specified, the responses are assumed to form a single time series.
- `homogeneous`: Logical indicating whether the transition models include time-varying covariates; used internally to determine the dimensions of certain arrays, notably `trDens`.
- `stationary`: This argument should no longer be used; if not NULL, the value of stationary is copied to the homogeneous argument, with a warning. In future versions this argument may be dropped or used for different purposes, i.e., for specifying models in which the initial state probabilities are constrained to be the stationary distribution of the transition matrix.
- `...`: Not used currently.

Details

The function `makeDepmix` creates an S4 object of class `depmix`, which needs to be fitted using `fit` to optimize the parameters. This function is provided to have full control, e.g., by specifying one's own response models with distributions that are not provided.

The response model(s) should be created by call(s) to `GLMresponse`, `MVNresponse` (see example below) or user-defined response models (see example below) that should extend the `response-class` and have the following methods: `dens`, `predict` and optionally `fit`. The `fit` function should have an
argument w, providing the weights. If the fit function is not provided, optimization should be done by using Rdonlp (use method="donlp" in calling fit on the depmix model, note that this is not the default). The first index of response models runs over the states of the model, and the second index over the responses to be modeled.

Value

See the depmix help page for the return value, a depmix object.

Author(s)

Ingmar Visser & Maarten Speekenbrink

See Also

fit, transInit, GLMresponse, depmix-methods for accessor functions to depmix objects.

Examples

# below example recreates the same model as on the fit help page in a roundabout way
# there we had:
# mod1 <- depmix(list(rt~1,corr~1),data=speed,transition=~Pacc,nstates=2,
#   family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
data(speed)

rModels <- list(
  list(   
    GLMresponse(formula=rt~1,data=speed,family=gaussian()),
    GLMresponse(formula=corr~1,data=speed,family=multinomial("identity"))
  ),
  list(   
    GLMresponse(formula=rt~1,data=speed,family=gaussian()),
    GLMresponse(formula=corr~1,data=speed,family=multinomial("identity"))
  )
)

transition <- list()
transition[[1]] <- transInit(~Pacc,nstates=2,data=speed)
transition[[2]] <- transInit(~Pacc,nstates=2,data=speed)
inMod <- transInit(~1,ns=2,data=data.frame(rep(1,3)),family=multinomial("identity"))
mod <- makeDepmix(response=rModels,transition=transition,prior=inMod,
  ntimes=c(168,134,137),homogeneous=FALSE)

set.seed(3)
fm1 <- fit(mod)
fml
summary(fm1)
# generate data from two different multivariate normal distributions
m1 <- c(0,1)
sd1 <- matrix(c(1,0.7,.7,1),2,2)
m2 <- c(1,0)
sd2 <- matrix(c(2,.1,.1,1),2,2)
set.seed(2)
y1 <- mvrnorm(50,m1,sd1)
y2 <- mvrnorm(50,m2,sd2)
# this creates data with a single change point
y <- rbind(y1,y2)

# now use makeDepmix to create a depmix model for this bivariate normal timeseries
rModels <- list()
rModels[[1]] <- list(MVNresponse(y~1))
rModels[[2]] <- list(MVNresponse(y~1))
trstart=c(0.9,0.1,0.1,0.9)
transition <- list()
transition[[1]] <- transInit(~1,nstates=2,data=data.frame(1),pstart=c(trstart[1:2]))
transition[[2]] <- transInit(~1,nstates=2,data=data.frame(1),pstart=c(trstart[3:4]))
instart=runif(2)
inMod <- transInit(~1,ns=2,ps=instart,data=data.frame(1))
mod <- makeDepmix(response=rModels,transition=transition,prior=inMod)
fm2 <- fit(mod,emc=em.control(random=FALSE))

# where is the switch point?
plot(as.ts(posterior(fm2, type="smoothing")[,1]))

# in below example we add the exgaus distribution as a response model and fit
# this instead of the gaussian distribution to the rt slot of the speed data
# most of the actual computations for the exgaus distribution is done by calling
# functions from the gamlss family of packages; see their help pages for
# interpretation of the mu, nu and sigma parameters that are fitted below

## Not run:
require(gamlss)
require(gamlss.dist)
data(speed)
rt <- speed$rt

# define a response class which only contains the standard slots, no additional slots
setClass("exgaus", contains="response")

# define a generic for the method defining the response class
setGeneric("exgaus", function(y, pstart = NULL, fixed = NULL, ...) standardGeneric("exgaus"))
# define the method that creates the response class

```r
setMethod("exgaus", 
  signature(y="ANY"),
  function(y,pstart=NULL,fixed=NULL, ...) {
    y <- matrix(y,length(y))
    x <- matrix(1)
    parameters <- list()
    npar <- 3
    if(is.null(fixed)) fixed <- as.logical(rep(0,npar))
    if(!is.null(pstart)) {
      if(length(pstart)!=npar) stop("length of 'pstart' must be ",npar)
      parameters$mu <- pstart[1]
      parameters$sigma <- log(pstart[2])
      parameters$nu <- log(pstart[3])
    }
    mod <- new("exgaus",parameters=parameters,fixed=fixed,x=x,y=y,npar=npar)
    mod
  })
```

```r
setMethod("show","exgaus", 
  function(object) {
    cat("Model of type exgaus (see ?gamlss for details) \n")
    cat("Parameters: \n")
    cat("mu: ", object@parameters$mu, "\n")
    cat("sigma: ", object@parameters$sigma, "\n")
    cat("nu: ", object@parameters$nu, "\n")
  })
```

```r
setMethod("dens","exgaus", 
  function(object,log=FALSE) {
    dexGAUS(object@y, mu = predict(object),
    sigma = exp(object@parameters$sigma), nu = exp(object@parameters$nu), log = log)
  })
```

```r
setMethod("getpars","response", 
  function(object,which="pars",...) {
    switch(which, 
      "pars" = {
        parameters <- numeric()
        parameters <- unlist(object@parameters)
        pars <- parameters
      },
      "fixed" = {
        pars <- object@fixed
      }
    }
    return(pars)
  })
```

setMethod("setpars","exgaus",
  function(object, values, which="pars", ...) {
    npar <- npar(object)
    if(length(values)!=npar) stop("length of 'values' must be",npar)
    # determine whether parameters or fixed constraints are being set
    nms <- names(object@parameters)
    switch(which,
      "pars"= {
        object@parameters$mu <- values[1]
        object@parameters$sigma <- values[2]
        object@parameters$nu <- values[3]
      },
      "fixed" = {
        object@fixed <- as.logical(values)
      }
    )
    names(object@parameters) <- nms
    return(object)
  })

setMethod("fit","exgaus",
  function(object,w) {
    if(missing(w)) w <- NULL
    y <- object@y
    fit <- gamlss(y~1,weights=w,family=exGAUS(),
      control=gamlss.control(n.cyc=100,trace=FALSE),
      mu.start=object@parameters$mu,
      sigma.start=exp(object@parameters$sigma),
      nu.start=exp(object@parameters$nu))
    pars <- c(fit$mu.coefficients,fit$sigma.coefficients,fit$nu.coefficients)
    object <- setpars(object,pars)
    object
  })

setMethod("predict","exgaus",
  function(object) {
    ret <- object@parameters$mu
    return(ret)
  })
)

rModels <- list(
  list( exgaus(rt,pstart=c(5,.1,.1)),
    GLMresponse(formula=corr~1, data=speed,
      family=multinomial("identity"), pstart=c(0.5,0.5))
  ),
  list( exgaus(rt,pstart=c(6,.1,.1)),
    GLMresponse(formula=corr~1, data=speed,
family=multinomial("identity"), pstart=c(.1,.9))
)
)

trstart=c(0.9,0.1,0.1,0.9)
transition <- list()
transition[[1]] <- transInit(~Pacc,nstates=2,data=speed,pstart=c(trstart[1:2],0,0))
transition[[2]] <- transInit(~Pacc,nstates=2,data=speed,pstart=c(trstart[3:4],0,0))

instart=c(0.5,0.5)
inMod <- transInit(~1,ns=2,ps=instart,family=multinomial("identity"), data=data.frame(rep(1,3))

mod <- makeDepmix(response=rModels,transition=transition,prior=inMod,ntimes=c(168,134,137), homogeneous=FALSE)

fm3 <- fit(mod,emc=em.control(rand=FALSE))
summary(fm3)

## End(Not run)

---

mix Mixture Model Specification

Description
mix creates an object of class mix, an (independent) mixture model (as a limit case of dependent mixture models in which all observed time series are of length 1), otherwise known latent class or mixture model. For a short description of the package see depmixS4.

Usage

mix(response, data=NULL, nstates, family=gaussian(), prior=~1, initaldata=NULL, respstart=NULL, instart=NULL,...)

Arguments

response The response to be modeled; either a formula or a list of formulae in the multivariate case; this interfaces to the glm distributions. See 'Details'.
data An optional data.frame to interpret the variables in the response and transition arguments.
nstates The number of states of the model.
family A family argument for the response. This must be a list of family's if the response is multivariate.
prior A one-sided formula specifying the density for the prior or initial state probabilities.
initdata  An optional data.frame to interpret the variables occurring in prior. The number of rows of this data.frame must be equal to the number of cases being modeled. See 'Details'.

respstart  Starting values for the parameters of the response models.

instart  Starting values for the parameters of the prior or initial state probability model.

...  Not used currently.

Details

The function `mix` creates an S4 object of class `mix`, which needs to be fitted using `fit` to optimize the parameters.

The response model(s) are by default created by call(s) to `GLMresponse` using the formula and the family arguments, the latter specifying the error distribution. See `GLMresponse` for possible values of the family argument for glm-type responses (i.e. a subset of the glm family options, and the multinomial). Alternative response distributions are specified by using the `makeDepmix` function. Its help page has examples of specifying a model with a multivariate normal response, as well as an example of adding a user-defined response model, in this case for the ex-gauss distribution.

If `response` is a list of formulae, the response’s are assumed to be independent conditional on the latent state.

The prior density is modeled as a multinomial logistic. This model is created by a call to `transInit`. Starting values may be provided by the respective arguments. The order in which parameters must be provided can be easily studied by using the `setpars` and `getpars` functions.

Linear constraints on parameters can be provided as argument to the `fit` function.

The print function prints the formulae for the response and prior models along with their parameter values.

Value

`mix` returns an object of class `mix` which has the following slots:

- **response**: A list of a list of response models; the first index runs over states; the second index runs over the independent responses in case a multivariate response is provided.
- **prior**: A multinomial logistic model for the initial state probabilities.
- **dens,init**: See `mix-class` help for details. For internal use.
- **ntimes**: A vector made by `rep(1,nrow(data))`; for internal use only.
- **nstates**: The number of states of the model.
- **nresp**: The number of independent responses.
- **npars**: The total number of parameters of the model. Note: this is not the degrees of freedom because there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior probabilities.

Author(s)

Ingmar Visser & Maarten Speekenbrink
References


See Also

`fit, transInit, GLMresponse, depmix-methods` for accessor functions to `depmix` objects.

Examples

```r
# four binary items on the balance scale task
data(balance)

# define a latent class model
instart=c(0.5,0.5)
set.seed(1)
respstart=runif(16)
# note that ntimes argument is used to make this a mixture model
mod <- mix(list(d1~1,d2~1,d3~1,d4~1), data=balance, nstates=2,
family=list(multinomial(),multinomial(),multinomial(),multinomial()),
respstart=respstart,instart=instart)
# to see the model
mod
```

mix-class

Class "mix"

Description

A `mix` model.

Objects from the Class

Objects can be created by calls to `mix`.

Slots

response: List of list of response objects.
prior: `transInit` object; model for the prior probabilities, also unconditional probabilities
dens: Array of dimension `sum(ntimes)*nresp*nstates` providing the densities of the observed responses for each state.
init: Array of dimension `length(ntimes)*nstates` with the current predictions for the initial state probabilities.
nstates: The number of states (classes) of the model.
nresp: The number of independent responses.
ntimes: A vector of 1's for each case; for internal use.
mix.fitted-class

npars: The total number of parameters of the model. This is not the degrees of freedom, ie there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior.

Accessor Functions

The following functions should be used for accessing the corresponding slots:

npar: The number of parameters of the model.
nresp: The number of responses.
nstates: The number of states.
ntimes: The vector of independent time series lengths.

Author(s)

Ingmar Visser & Maarten Speekenbrink

Examples

showClass("mix")

mix.fitted-class  Class "mix.fitted" (and "mix.fitted.classLik")

Description

A fitted mix model.

Slots

A mix.fitted object is a mix object with three additional slots, here is the complete list:

response: List of list of response objects.
prior: transInit object.
dens: Array of dimension sum(ntimes)*nresp*nstates providing the densities of the observed responses for each state.
init: Array of dimension length(ntimes)*nstates with the current predictions for the initial state probabilities.
ntimes: A vector containing the lengths of independent time series; if data is provided, sum(ntimes) must be equal to nrow(data).
nstates: The number of states of the model.
nresp: The number of independent responses.
npars: The total number of parameters of the model. This is not the degrees of freedom, ie there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior.
message: This provides some information on convergence, either from the EM algorithm or from Rdonlp2.
conMat: The linear constraint matrix, which has zero rows if there were no constraints.
lin.lower The lower bounds on the linear constraints.
lin.upper The upper bounds on the linear constraints.
posterior: Posterior (Viterbi) state sequence.

Details
The print function shows some convergence information, and the summary method shows the parameter estimates.

Extends
Class "mix" directly. mix.fitted.classLik is similar to mix.fitted, the only difference being that the model is fitted by maximising the classification likelihood.

Author(s)
Ingmar Visser & Maarten Speekenbrink

Examples
showClass("mix.fitted")
nresp: The number of independent responses.
npars: The total number of parameters of the model. This is not the degrees of freedom, ie there are redundancies in the parameters, in particular in the multinomial models for the transitions and prior.
states: A matrix with the true states/classes.

Accessor Functions
The following functions should be used for accessing the corresponding slots:

npar: The number of parameters of the model.
nresp: The number of responses.
nstates: The number of states.
ntimes: The vector of independent time series lengths.

Author(s)
Maarten Speekenbrink & Ingmar Visser

Description
Fit a model using multiple sets of starting values.

Usage

## S4 method for signature 'mix'
multistart(object, nstart=10, initIters=10, ...)

Arguments

object: An object of class mix or depmix.
nstart: The number of sets of starting values that are used.
initIters: The number of EM iterations that each set of starting values is run.
... Not used currently.

Details
Starting values in the EM algorithm are generated by randomly assigning posterior state probabilities for each observation using a Dirichlet distribution. This is done nstart times. The EM algorithm is run initIters times for each set of starting values. The then best fitting model is then optimized until convergence. A warning is provided about the number of sets of starting values that are infeasible, e.g. due to non-finite log likelihood, if that number is larger than zero. Note that the number of iterations reported in the final fitted model does not include the initial number of iterations that EM was run for.
posterior

Value

A fitted (dep)mix object.

Author(s)

Ingmar Visser & Maarten Speekenbrink

Examples

data(speed)

# this example is from ?fit with fit now replaced by multistart and the
# set.seed statement is left out
mod1 <- depmix(list(rt~1,corr~1),data=speed,transition=Pacc,nstates=2,
    family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
set.seed(3)
  fmod1 <- fit(mod1)
  fmod2 <- multistart(mod1)
  fmod1
  fmod2

posterior

Posterior state/class probabilities and classification

Description

Return posterior state classifications and/or probabilities for a fitted (dep-)mix object. In the case
of a latent class or mixture model, states refer to the classes/mixture components.

There are different ways to define posterior state probabilities and the resulting classifications. The
‘type’ argument can be used to specify the desired definition. The default is currently set to ‘viterbi’. Other options are ‘global’ and ‘local’ for state classification, and ‘filtering’ and ‘smoothing’ for state probabilities. See Details for more information.

Usage

## S4 method for signature 'depmix'
posterior(object, type = c("viterbi", "global", "local", "filtering", "smoothing"))
## S4 method for signature 'depmix.fitted'
posterior(object, type = c("viterbi", "global", "local", "filtering", "smoothing"))
## S4 method for signature 'mix'
posterior(object, type = c("viterbi", "global", "local", "filtering", "smoothing"))
## S4 method for signature 'mix.fitted'
posterior(object, type = c("viterbi", "global", "local", "filtering", "smoothing"))
Arguments

object A (fitted)(dep-)mix object.

type character, partial matching allowed. The type of classification or posterior probability desired.

Details

After fitting a mix or depmix model, one is often interested in determining the most probable mixture components or hidden states at each time-point $t$. This is also called decoding the hidden states from the observed data. There are at least two general ways to consider state classification: ‘global’ decoding means determining the most likely state sequence, whilst ‘local’ decoding means determining the most likely state at each time point whilst not explicitly considering the identity of the hidden states at other time points. For mixture models, both forms of decoding are identical.

Global decoding is based on the conditional probability $p(S_1, \ldots, S_T \mid Y_1, \ldots, Y_T)$, and consists of determining, at each time point $t = 1, \ldots, T$:

$$s^*_t = \arg \max_{i=1}^N p(S_1 = s_1, \ldots, S_{t-1} = s_{t-1}, S_t = i, S_{t+1} = s_{t+1}, \ldots, S_T = s_T \mid Y_1, \ldots, Y_T)$$

where $N$ is the total number of states. These probabilities and the resulting classifications, are computed through the viterbi algorithm. Setting type = ’viterbi' returns a data.frame with the Viterbi-decoded global state sequence in the first column, and the normalized ''delta'' probabilities in the remaining columns. These ''delta'' probabilities are defined as the joint probability of the most likely state sequence ending in state $i$ at time $t$, and all the observations up to time $t$. The normalization of these joint probabilities is done on a time-point basis (i.e., dividing the delta probability by the sum of the delta probabilities for that time point for all possible states $j$ (including state $i$)). These probabilities are not straightforward to interpret. Setting type = ''global'' returns just a vector with the Viterbi-decoded global state sequence.

Local decoding is based on the smoothing probabilities $p(S_t \mid Y_1, \ldots, Y_T)$, which are the ''gamma'' probabilities computed with the forwardbackward algorithm. Local decoding then consists of determining, at each time point $t = 1, \ldots, T$:

$$s^*_t = \arg \max_{i=1}^N p(S_t = i \mid Y_1, \ldots, Y_T)$$

where $N$ is the total number of states. Setting type = ''local'' returns a vector with the local decoded states. Setting type = ''smoothing'' returns the smoothing probabilities which underlie this classification. When considering the posterior probability of each state, the values returned by type = ''smoothing'' are most likely what is wanted by the user.

The option type = ''filtering'' returns a matrix with the so-called filtering probabilities, defined as $p(S_t \mid Y_1, \ldots, Y_t)$, i.e. the probability of a hidden state at time $t$ considering the observations up to and including time $t$.

See the fit help page for an example.

Value

The return value of posterior depends on the value of the type argument:
type = 'viterbi'
Returns a data.frame with nstates(object) + 1 columns; the first column contains the states decoded through the Viterbi algorithm, the remaining columns contain the (normalized) delta probabilities.

type = 'global'
Returns a vector which contains the states decoded through the Viterbi algorithm.

type = 'local'
Returns a vector which contains the states decoded as the maximum of the smoothing probabilities.

type = 'filtering'
Returns a matrix which contains the posterior probabilities of each state, conditional upon the responses observed thus far.

type = 'smoothing'
Returns a matrix which contains the posterior probabilities of each state, conditional upon all the responses observed.

See Details for more information.

Note
The initial version of this function was a simple wrapper to return the value of the posterior slot in a mix-fitted or depmix-fitted object. The value of this slot is set by a call of the viterbi method. For backwards compatibility, the default value of the type argument is set to "viterbi", which returns the same. As the "delta" probabilities returned as part of this may be misinterpreted, and may not be the desired posterior probabilities, the updated version of this method now allows for other return values, and the type = "viterbi" option should be considered deprecated.

Author(s)
Maarten Speekenbrink & Ingmar Visser

References

Examples

data(speed)

# 2-state model on rt and corr from speed data set
# with Pacc as covariate on the transition matrix
# ntimes is used to specify the lengths of 3 separate series
mod <- depmix(list(rt~1,corr~1),data=speed,transition=~Pacc,nstates=2,
family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))
fmod <- fit(mod)

# Global decoding:
pst_global <- posterior(fmod, type = "global")
response-class

Class "response"

Description

A generic response model for depmix models.

Arguments

object An object of class "response".

Details

This class offers a framework from which to build specific response models such as glm based responses or multinomial responses. For extensibility, objects with class response should have at least methods: dens to return the density of responses, and getpars and setpars methods to get and set parameters.

The constr slot is used for information on constraints that are inherently part of a model; the only such constraints which are currently used are 1) the sum constraint in multinomial models with identity link, and 2) a lower bound of zero of sd parameters in gaussian distributions. Such constraints are only used in fitting models with general optimization routines such as Rsolnp; the EM algorithm automagically respects the sum constraint.

lin: Derivative of linear constraint.
linup: Upper bounds for linear constraints.
lnlow: Lower bounds for linear constraints.
parup: Upper bounds on parameters.
parlow: Lower bounds on parameters.
Slots

parameters: A (named) list of parameters.
fixed: A logical vector indicating which parameters are fixed.
y: A matrix with the actual response; possibly multivariate.
x: A design matrix; possibly only an intercept term.
npar: The number of parameters.
constr: Information on constraints.

Accessor Functions

The following functions should be used for accessing the corresponding slots:

npar: The number of parameters of the model.
getdf: The number of non-fixed parameters.

Author(s)

Ingmar Visser & Maarten Speekenbrink

response-classes

Class "GLMresponse" and class "transInit"

Description

Specific instances of response models for depmix models.

Details

The GLMresponse-class offers an interface to the glm functions that are subsequently used in fitting the depmix model of which the response is a part.

The transInit is an extension of response that is used to model the transition matrix and the initial state probabilities by the use of a multinomial logistic model, the difference being that in fact the response is missing as the transitions between states are not observed. This class has its own fit function which is an interface to the multinom function in nnet.

Slots

Both GLMresponse and transInit contain the response-class. In addition to the slots of that class, these classes have the following slots:

formula: A formula that specifies the model.
family: A family object specifying the link function. See the GLMresponse help page for possible options.
Accessor Functions

The following functions should be used for accessing the corresponding slots:

- npar: The number of parameters of the model.
- getdf: The number of non-fixed parameters.

Author(s)

Ingmar Visser & Maarten Speekenbrink

Response models currently implemented in depmix.

Description

Depmix contains a number of default response models. We provide a brief description of these here.

BINOMresponse

BINOMresponse is a binomial response model. It derives from the basic GLMresponse class.

y: The dependent variable can be either a binary vector, a factor, or a 2-column matrix, with successes and misses.

x: The design matrix.

Parameters: A named list with a single element “coefficients”, which contains the GLM coefficients.

GAMMAresponse

GAMMAresponse is a model for a Gamma distributed response. It extends the basic GLMresponse class directly.

y: The dependent variable.

x: The design matrix.

Parameters: A named list with a single element “coefficients”, which contains the GLM coefficients.

MULTINOMresponse

MULTINOMresponse is a model for a multinomial distributed response. It extends the basic GLMresponse class, although the functionality is somewhat different from other models that do so.

y: The dependent variable. This is a binary matrix with N rows and Y columns, where Y is the total number of categories.

x: The design matrix.

Parameters: A named list with a single element “coefficients”, which is an ncol(x) by ncol(y) matrix which contains the GLM coefficients.
MVNresponse

MVNresponse is a model for a multivariate normal distributed response. See code makeDepmix for an example of how to use this and other non-glm like distributions.

- **y**: The dependent variable. This is a matrix.
- **x**: The design matrix.
- **Parameters**: A named list with elements “coefficients”, which contains the GLM coefficients, and “Sigma”, which contains the covariance matrix.

NORMresponse

NORMresponse is a model for a normal (Gaussian) distributed response. It extends the basic GLMresponse class directly.

- **y**: The dependent variable.
- **x**: The design matrix.
- **Parameters**: A named list with elements “coefficients”, which contains the GLM coefficients, and “sd”, which contains the standard deviation.

POISSONresponse

POISSONresponse is a model for a Poisson distributed response. It extends the basic GLMresponse class directly.

- **y**: The dependent variable.
- **x**: The design matrix.
- **Parameters**: A named list with a single element “coefficients”, which contains the GLM coefficients.

Author(s)

Maarten Speekenbrink & Ingmar Visser

Examples

```r
# binomial response model
x <- rnorm(1000)
p <- plogis(x)
ss <- rbinom(1000,1,p)
mod <- GLMresponse(cbind(ss,1-ss)~x,family=binomial())
fit(mod)
glm(cbind(ss,1-ss)~x, family=binomial)

# gamma response model
x=runif(1000,1,5)
res <- rgamma(1000,x)
## note that gamma needs proper starting values which are not
## provided by depmixS4 (even with them, this may produce warnings)
```
mod <- GLMresponse(res~x,family=Gamma(),pst=c(0.8,1/0.8))
fit(mod)
glm(res~x,family=Gamma)

# multinomial response model
x <- sample(0:1,1000,rep=TRUE)
mod <- GLMresponse(sample(1:3,1000,rep=TRUE)*x,family=multinomial(),pstart=c(0.33,0.33,0.33,0,0,1))
mod@y <- simulate(mod)
fit(mod)
colSums(mod@y[which(x==0),])/length(which(x==0))
colSums(mod@y[which(x==1),])/length(which(x==1))
# note that the response is treated as factor here, internal representation is in 
# dummy coded format:
head(mod@y)
# similar to the binomial model, data may also be entered in multi-column format 
# where the n for each row can be different
dt <- data.frame(y1=c(0,1,1,2,4,5),y2=c(1,0,1,0,1,0),y3=c(4,4,3,2,1,1))
m2 <- mix(cbind(y1,y2,y3)~1,data=dt,ns=2,family=multinomial("identity"))
fm2 <- fit(m2)
summary(fm2)

# multivariate normal response model
mn <- c(1,2,3)
sig <- matrix(c(1,.5,0,.5,1,0,0,0,2),3,3)
y <- mvrnorm(1000,mn,sig)
mod <- MVNresponse(y~1)
fit(mod)
colMeans(y)
var(y)

# normal (gaussian) response model
y <- rnorm(1000)
mod <- GLMresponse(y~1)
fm <- fit(mod)
cat("Test gaussian fit: ", all.equal(getpars(fm),c(mean(y),sd(y)),check.attributes=FALSE))

# poisson response model
x <- abs(rnorm(1000,2))
res <- rpois(1000,x)
mod <- GLMresponse(res~x,family=poisson())
fit(mod)
glm(res~x, family=poisson)

# this creates data with a single change point with Poisson distributed data
set.seed(3)
y1 <- rpois(50,1)
y2 <- rpois(50,2)
ydf <- data.frame(y=c(y1,y2))

# fit models with 1 to 3 states
m1 <- depmix(y~1,ns=1,family=poisson(),data=ydf)
fm1 <- fit(m1)
m2 <- depmix(y~1,ns=2,family=poisson(),data=ydf)
fm2 <- fit(m2)
m3 <- depmix(y~1,ns=3,family=poisson(),data=ydf)
fm3 <- fit(m3,em=em.control(maxit=500))

# plot the BICs to select the proper model
plot(1:3,c(BIC(fm1),BIC(fm2),BIC(fm3)),ty="b")

### simulate

Methods to simulate from (de-)mix models

Description

Random draws from (de-)mix objects.

Usage

```
## S4 method for signature 'depmix'
simulate(object, nsim=1, seed=NULL, ...)

## S4 method for signature 'mix'
simulate(object, nsim=1, seed=NULL, ...)

## S4 method for signature 'response'
simulate(object, nsim=1, seed=NULL, times, ...)

## S4 method for signature 'GLMresponse'
simulate(object, nsim=1, seed=NULL, times, ...)

## S4 method for signature 'transInit'
simulate(object, nsim=1, seed=NULL, times, is.prior=FALSE, ...)
```

Arguments

- **object**: Object to generate random draws. An object of class `mix`, `depmix`, `response` or `transInit`.
- **nsim**: The number of draws (one draw simulates a data set of the size that is defined by `ntimes`); defaults to 1.
- **seed**: Set the seed.
- **times**: (optional) An indicator vector indicating for which times in the complete series to generate the data. For internal use.
- **is.prior**: For `transInit` objects, indicates whether it is a prior (init) model, or not (i.e., it is a transition model).
- **...**: Not used currently.
Details

For a `depmix` model, simulate generates `nsim` random state sequences, each of the same length as the observation sequence in the `depmix` model (i.e., `sum(ntimes(object))`). The state sequences are then used to generate `nsim` observation sequences of the same length.

For a `mix` model, simulate generates `nsim` random class assignments for each case. Those assignments are then used to generate observation/response values from the appropriate distributions.

Setting the `times` option selects the time points in the total state/observation sequence (i.e., counting is continued over `ntimes`). Direct calls of simulate with the `times` option are not recommended.

Value

For a `depmix` object, a new object of class `depmix.sim`.

For a `transInit` object, a state sequence.

For a `response` object, an observation sequence.

Author(s)

Maarten Speekenbrink & Ingmar Visser

Examples

```r
y <- rnorm(1000)
respst <- c(0,1,2,1)
trst <- c(0.9,0.1,0.1,0.9)

df <- data.frame(y=y)

mod <- depmix(y~1,data=df,respst=respst,trst=trst,inst=c(0.5,0.5),nti=1000,nst=2)

mod <- simulate(mod)
```

sp500

*Standard & Poor’s 500 index*

Description

This data set consists of (monthly) values of the S&P 500 stock exchange index. The variable of interest is the logarithm of the return values, i.e., the logarithm of the ratio of indices, in this case the closing index is used.

Usage

`data(speed)`
Format

A data frame with 744 observations and 6 variables.

Open   Index at the start of trading.
High   Highest index.
Low    Lowest index.
Close  Index at the close of trading.
Volume The volume of trading.
logret The log return of the closing index.

Source

Yahoo Data.

Examples

data(sp500)

# the data can be made with the following code (eg to include a longer or
# shorter time span)

## Not run:
require(TTR)

# load SP500 returns
Sys.setenv(tz="UTC")

sp500 <- getYahooData('^GSPC', start=19500101, end=20120228, freq='daily')
ep <- endpoints(sp500, on="months", k=1)
sp500 <- sp500[ep[2:(length(ep)-1)]]
sp500$ret <- log(sp500$Close) - lag(log(sp500$Close))
sp500 <- na.exclude(sp500)

## End(Not run)
Description

This data set is a bivariate series of response times and accuracy scores of a single participant switching between slow/accurate responding and fast guessing on a lexical decision task. The slow and accurate responding, and the fast guessing can be modelled using two states, with a switching regime between them. The dataset further contains a third variable called Pacc, representing the relative pay-off for accurate responding, which is on a scale of zero to one. The value of Pacc was varied during the experiment to induce the switching. This data set is from participant A in experiment 1a from Dutilh et al (2011).

Usage

data(speed)

Format

A data frame with 439 observations on the following 4 variables.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>rt</td>
<td>a numeric vector of response times (log ms)</td>
</tr>
<tr>
<td>corr</td>
<td>a numeric vector of accuracy scores (0/1)</td>
</tr>
<tr>
<td>Pacc</td>
<td>a numeric vector of the pay-off for accuracy</td>
</tr>
<tr>
<td>prev</td>
<td>a numeric vector of accuracy scores (0/1) on the previous trial</td>
</tr>
</tbody>
</table>

Source


Corresponding author: g.dutilh@uva.nl

Examples

data(speed)
## maybe str(speed) ; plot(speed) ...

```
stationary

  Compute the stationary distribution of a transition probability matrix.
```

Description

See title.

Usage

stationary(tpm)
Arguments

- `tpm`: a transition probability matrix.

Value

A vector with the stationary distribution such that $p = tpm * p$.

Author(s)

Ingmar Visser

---

transInit

*Methods for creating depmix transition and initial probability models*

Description

Create `transInit` objects for `depmix` models using formulae and family objects.

Usage

```r
transInit(formula, nstates, data=NULL, family=multinomial(),
pstart=NULL, fixed=NULL, prob=TRUE, ...)
```

```r
## S4 method for signature 'transInit'
getdf(object)
```

Arguments

- `formula`: A model formula.
- `data`: An optional data.frame to interpret the variables from the formula argument in.
- `family`: A family object; see details.
- `pstart`: Starting values for the coefficients.
- `fixed`: Logical vector indicating which parameters are to be fixed.
- `prob`: Logical indicating whether the starting values for multinomial() family models are probabilities or logistic parameters (see details).
- `nstates`: The number of states of the model.
- `object`: An object of class `transInit`.
- `...`: Not used currently.
Details

The transInit model provides functionality for the multinomial probabilities of the transitions between states, as well as for the prior or initial probabilities. These probabilities may depend on (time-varying) covariates. The model can be used with link functions mlogit and identity; the latter is the default when no covariates are. With the mlogit link function, the transition probabilities are modeled as baseline logistic multinomials (see Agresti, 2002, p. 272 ff.).

Start values for the parameters may be provided using the pstart argument; these can be provided as probabilities, the default option, or as baseline logistic parameters, use the prob argument to specify the chosen option. The default baseline category is set to 1, which can be modified through calling, say, family=multinomial(base=2).

Note that the transInit model extends the response-class, but that it actually lacks a response, i.e. the y-slot is empty, at the time of construction, as the transitions are not observed.

getdf returns the number of free parameters of a transInit model.

Value

transInit return objects of class transInit; this class extends the response-class.

Author(s)

Ingmar Visser & Maarten Speekenbrink

References


---

### Parameter standard errors

These functions provide standard errors for parameters of (dep-)mix models.

#### Usage

```r
## S4 method for signature 'mix'
vcov(object, fixed=NULL, equal=NULL, conrows=NULL, conrows.upper=NULL, conrows.lower=NULL, tolerance=1e-6,
method="finiteDifferences", ...)

## S4 method for signature 'mix'
standardError(object, fixed=NULL, equal=NULL, conrows=NULL, conrows.upper=NULL, conrows.lower=NULL, tolerance=1e-6,
method="finiteDifferences", ...)
```
## S4 method for signature 'mix'
confint(object, level=0.95, fixed=NULL, equal=NULL,
conrows=NULL, conrows.upper=NULL, conrows.lower=NULL, tolerance=1e-6,
method="finiteDifferences", ...)

## S4 method for signature 'mix'
hessian(object, tolerance=1e-6,
method="finiteDifferences", ...)

### Arguments

- **object**: A (dep-)mix object; see `depmix` for details.
- **fixed, equal**: These arguments are used to specify constraints on a model; see usage details here: `fit`.
- **conrows**: These arguments are used to specify constraints on a model; see usage details here: `fit`.
- **conrows.upper**: These arguments are used to specify constraints on a model; see usage details here: `fit`.
- **conrows.lower**: These arguments are used to specify constraints on a model; see usage details here: `fit`.
- **tolerance**: Threshold used for testing whether parameters are estimated on the boundary of the parameter space; if so, they are ignored in these functions.
- **method**: The method used for computing the Hessian matrix of the parameters; currently only a finite differences method (using `fdHess` from package `nlme`) is implemented and hence used by default.
- **level**: The desired significance level for the confidence intervals.
- **...**: Further arguments passed to other methods; currently not in use.

### Details

`vcov` computes the variance-covariance matrix of a (dep-)mix object, either fitted or not. It does so by first constructing a Hessian matrix through the use of `hessian` and then transforming this as described in Visser et al (2000), taking into account the linear constraints that are part of the model. Currently, `hessian` has a single method using finite differences to arrive at an approximation of the second order derivative matrix of the parameters.

`confint` and `standardError` use `vcov` to compute confidence intervals (the confidence level can be set through an argument) and standard errors respectively. The latter are computed first by using `sqrt(diag(vcov))` and the confidence intervals are computed through the normal approximation.

If and when these methods are applied to fitted models, the linear constraint matrix is obtained from the `mix.fitted` or `depmix.fitted` slot `lincon` (supplemented with additional constraints if those are provided through the `equal` and other arguments to these functions).

All four functions exclude parameters that are estimated on or near (this can be controlled using the `tolerance` argument) their boundary values. Setting this argument to zero can result in error as the `fdHess` function requires an environment around the parameter estimate that provides proper log-likelihood values, which parameter on or over their boundary values are not guaranteed to provided. Fixed parameters are similarly ignored in these four functions.
vcov

Value

vcov returns a named list with elements vcov, elements, and lincon. standardError returns a data.frame with columns par, elements, and se. confint returns a data.frame with columns par, elements, and two columns for the lower and upper bounds of the confidence intervals (with the column names indicating the level of the interval.)

vcov : The variance-covariance matrix of the parameters.

elements : Vector of length npar(object) indicating which elements of the parameter vector are included in computing the hessian, the variance-covariance matrix, the standard errors and/or the confidence intervals.

inc : 'inc'luded parameter.

fix : 'fix'ed parameter.

bnd : parameter estimated on the boundary.

par : The values of the parameters.

se : The values of the standard errors of the parameters.

lower/upper : The lower and upper bounds of the confidence intervals; column names indicate the as in 0.5+/-level/2, using the level argument.

Note

Note that the quality of the resulting standard errors is similar to those reported in Visser et al (2000) for both bootstrap and the profile likelihood methods. In Visser et al (2000), the finite differences standard errors were somewhat less precise as they relied on a very parsimonious but indeed less precise method for computing the finite differences approximation (computation time was a much scarcer resource at the time then it is now).

Author(s)

Ingmar Visser

References


Examples

data(speed)

# 2-state model on rt and corr from speed data set
# with Pacc as covariate on the transition matrix
# ntimes is used to specify the lengths of 3 separate series
mod1 <- depmix(list(rt~1,corr~1),data=speed,transition=~Pacc,nstates=2,
family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))

# fit the model
set.seed(3)
fmod1 <- fit(mod1)
vcov(fmod1)$vcov # $
standardError(fmod1)
confint(fmod1)

---

**viterbi**

**Viterbi algorithm for decoding the most likely state sequence**

**Description**

Apply the Viterbi algorithm to compute the maximum a posteriori state sequence for a mix or depmix object.

**Usage**

viterbi(object, na.allow=TRUE)

**Arguments**

- **object** A mix or depmix object.
- **na.allow** logical. If TRUE, the density of missing responses is set to 1, similar as in the forwardbackward algorithm. If FALSE, missing values have NA as density values, and will result in an error.

**Details**

The Viterbi algorithm is used for global decoding of the hidden state sequence. Global decoding is based on the conditional probability $p(S_1, \ldots, S_T \mid Y_1, \ldots, Y_T)$, and consists of determining, at each time point $t = 1, \ldots, T$:

$$s^*_t = \arg\max_{i=1}^N p(S_1 = s^*_1, \ldots, S_t = i, S_{t-1} = s^*_{t-1}, S_{t+1} = s^*_{t+1}, \ldots, S_T = s^*_T \mid Y_1, \ldots, Y_T)$$

where $N$ is the total number of states.

The Viterbi algorithm is a dynamic programming algorithm that relies on “delta” probabilities (see Rabiner, 1989), which are defined as the joint probability of the most likely state sequence ending in state $i$ at time $t$, and all the observations up to time $t$. The implementation here normalizes these probabilities on a time-point basis, dividing the delta probability by the sum of the delta probabilities for that time point for all possible states $j$ (including state $i$). The normalized delta probabilities for each state are returned in columns 2: (nstates(object) + 1), whilst column 1 contains the indices of the maximum a posteriori states.
viterbi

Value

viterbi returns a data.frame with in the first column the maximum a posteriori state sequence. This is a vector with integers corresponding to the index of the most likely hidden states. The remaining columns contain the normalized "delta" probabilities (see Details).

Author(s)

Maarten Speekenbrink

References


Examples

data(speed)

# 2-state model on rt and corr from speed data set
# with Pacc as covariate on the transition matrix
# ntimes is used to specify the lengths of 3 separate series
mod <- depmix(list(rt~1,corr~1),data=speed,transition=~Pacc,nstates=2,
   family=list(gaussian(),multinomial("identity")),ntimes=c(168,134,137))

fmod <- fit(mod)

# result of viterbi is stored in a depmix-fitted object in slot "posterior"
identical(viterbi(fmod),fmod@posterior)
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