# Package 'seqHMM' 

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Title Mixture Hidden Markov Models for Social Sequence Data and Other Multivariate, Multichannel Categorical Time Series

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Description Designed for fitting hidden (latent) Markov models and mixture hidden Markov models for social sequence data and other categorical time series. Also some more restricted versions of these type of models are available: Markov models, mixture Markov models, and latent class models. The package supports models for one or multiple subjects with one or multiple parallel sequences (channels). External covariates can be added to explain cluster membership in mixture models. The package provides functions for evaluating and comparing models, as well as functions for visualizing of multichannel sequence data and hidden Markov models. Models are estimated using maximum likelihood via the EM algorithm and/or direct numerical maximization with analytical gradients. All main algorithms are written in C++ with support for parallel computation.
Documentation is available via several vignettes in this page, and the paper by Helske and Helske (2019, [doi:10.18637/jss.v088.i03](doi:10.18637/jss.v088.i03)).

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biofam3c Three-channel biofam data

## Description

Biofam data from the TraMineR package converted into three channels.

## Format

A list including three sequence data sets for 2000 individuals with 16 state variables, and a separate data frame with 1 id variable, 8 covariates, and 2 weights variables.

## Details

This data is constructed from the biofam data in the TraMineR package. Here the original state sequences are converted into three separate data sets: children, married, and left. These include the corresponding life states from age 15 to 30 : childless or (having) children; single, married, or divorced; and (living) with parents or left home.
Note that the divorced state does not give information on parenthood or residence, so a guess is made based on preceeding states.

The fourth data frame covariates is a collection of additional variables from the original data:

| idhous | id |
| :--- | :--- |
| sex | sex |
| birthyr | birth year |
| nat_1_02 | first nationality |
| plingu02 | language of questionnaire |
| p02r01 | religion |
| p02r04 | religious participation |
| cspfaj | father's social status |
| cspmoj | mother's social status |
| wp00tbgp | weights inflating to the Swiss population |
| wp00tbgs | weights respecting sample size |

The data is loaded by calling data(biofam3c). It was built using following code:

```
data("biofam" , package = "TraMineR")
biofam3c <- with(biofam, {
## Building one channel per type of event left, children or married
bf <- as.matrix(biofam[, 10:25])
children <- bf == 4 | bf == 5 | bf == 6
married <- bf == 2 | bf == 3 | bf == 6
left <- bf == 1 | bf == 3 | bf == 5 | bf == 6 | bf == 7
```

```
children[children == TRUE] <- "children"
children[children == FALSE] <- "childless"
# Divorced parents
div <- bf[(rowSums(bf == 7) > 0 & rowSums(bf == 5) > 0) |
    (rowSums(bf == 7) > 0 & rowSums(bf == 6) > 0),]
children[rownames(bf) %in% rownames(div) & bf == 7] <- "children"
married[married == TRUE] <- "married"
married[married == FALSE] <- "single"
married[bf == 7] <- "divorced"
left[left == TRUE] <- "left home"
left[left == FALSE] <- "with parents"
# Divorced living with parents (before divorce)
wp <- bf[(rowSums(bf == 7) > 0 & rowSums(bf == 2) > 0 &
    rowSums(bf == 3) == 0 & rowSums(bf == 5) == 0 &
    rowSums(bf == 6) == 0) |
    (rowSums(bf == 7) > 0 & rowSums(bf == 4) > 0 &
    rowSums(bf == 3) == 0 & rowSums(bf == 5) == 0 &
    rowSums(bf == 6) == 0), ]
left[rownames(bf) %in% rownames(wp) & bf == 7] <- "with parents"
list("children" = children, "married" = married, "left" = left,
    "covariates" = biofam[, c(1:9, 26:27)])
})
```


## Source

biofam data constructed from the Swiss Household Panel https://forscenter.ch/projects/ swiss-household-panel/

## References

Müller, N. S., M. Studer, G. Ritschard (2007). Classification de parcours de vie à l'aide de l'optimal matching. In XIVe Rencontre de la Société francophone de classification (SFC 2007), Paris, 5-7 septembre 2007, pp. 157-160.

## Description

Function build_hmm constructs a hidden Markov model object of class hmm.

## Usage

```
build_hmm(
    observations,
    n_states,
    transition_probs,
    emission_probs,
    initial_probs,
    state_names = NULL,
    channel_names = NULL,
)
```


## Arguments

```
observations An stslist object (see seqdef) containing the sequences, or a list of such
objects (one for each channel).
n_states A scalar giving the number of hidden states (not used if starting values for model
    parameters are given with initial_probs, transition_probs,or emission_probs).
transition_probs
    A matrix of transition probabilities.
emission_probs A matrix of emission probabilities or a list of such objects (one for each chan-
    nel). Emission probabilities should follow the ordering of the alphabet of obser-
    vations (alphabet(observations), returned as symbol_names).
initial_probs A vector of initial state probabilities.
state_names A list of optional labels for the hidden states. If NULL, the state names are taken
    from the row names of the transition matrix. If this is also NULL, numbered states
    are used.
channel_names A vector of optional names for the channels.
... Additional arguments to simulate_transition_probs.
```


## Details

The returned model contains some attributes such as nobs and $d f$, which define the number of observations in the model and the number of estimable model parameters, used in computing BIC. When computing nobs for a multichannel model with $C$ channels, each observed value in a single channel amounts to $1 / C$ observation, i.e. a fully observed time point for a single sequence amounts to one observation. For the degrees of freedom df, zero probabilities of the initial model are defined as structural zeroes.

## Value

Object of class hmm with the following elements:
observations State sequence object or a list of such objects containing the data.
transition_probs A matrix of transition probabilities.
emission_probs A matrix or a list of matrices of emission probabilities.
initial_probs A vector of initial probabilities.
state_names Names for hidden states.
symbol_names Names for observed states.
channel_names Names for channels of sequence data.
length_of_sequences (Maximum) length of sequences.
n _sequences Number of sequences.
n _symbols Number of observed states (in each channel).
n_states Number of hidden states.
n_channels Number of channels.

## See Also

fit_model for estimating model parameters; and plot. hmm for plotting hmm objects.

## Examples

```
# Single-channel data
data("mvad", package = "TraMineR")
mvad_alphabet <- c("employment", "FE", "HE", "joblessness", "school",
                "training")
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet, states = mvad_scodes,
                        labels = mvad_labels, xtstep = 6)
# Initializing an HMM with 4 hidden states, random starting values
init_hmm_mvad1 <- build_hmm(observations = mvad_seq, n_states = 4)
# Starting values for the emission matrix
emiss <- matrix(NA, nrow = 4, ncol = 6)
emiss[1,] <- seqstatf(mvad_seq[, 1:12])[, 2] + 1
emiss[2,] <- seqstatf(mvad_seq[, 13:24])[, 2] + 1
emiss[3,] <- seqstatf(mvad_seq[, 25:48])[, 2] + 1
emiss[4,] <- seqstatf(mvad_seq[, 49:70])[, 2] + 1
emiss <- emiss / rowSums(emiss)
# Starting values for the transition matrix
tr <- matrix(
    c(0.80, 0.10, 0.05, 0.05,
        0.05, 0.80, 0.10, 0.05,
        0.05, 0.05, 0.80, 0.10,
        0.05, 0.05, 0.10, 0.80),
    nrow=4, ncol=4, byrow=TRUE)
```

```
# Starting values for initial state probabilities
init <- c(0.3, 0.3, 0.2, 0.2)
# HMM with own starting values
init_hmm_mvad2 <- build_hmm(
    observations = mvad_seq, transition_probs = tr,
    emission_probs = emiss, initial_probs = init)
##########################################
# Multichannel data
# Three-state three-channel hidden Markov model
# See ?hmm_biofam for a five-state version
data("biofam3c")
# Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
# Define colors
attr(marr_seq, "cpal") <- c("violetred2", "darkgoldenrod2", "darkmagenta")
attr(child_seq, "cpal") <- c("darkseagreen1", "coral3")
attr(left_seq, "cpal") <- c("lightblue", "red3")
# Left-to-right HMM with 3 hidden states and random starting values
set.seed(1010)
init_hmm_bf1 <- build_hmm(
    observations = list(marr_seq, child_seq, left_seq),
    n_states = 3, left_right = TRUE, diag_c = 2)
# Starting values for emission matrices
emiss_marr <- matrix(NA, nrow = 3, ncol = 3)
emiss_marr[1,] <- seqstatf(marr_seq[, 1:5])[, 2] + 1
emiss_marr[2,] <- seqstatf(marr_seq[, 6:10])[, 2] + 1
emiss_marr[3,] <- seqstatf(marr_seq[, 11:16])[, 2] + 1
emiss_marr <- emiss_marr / rowSums(emiss_marr)
emiss_child <- matrix(NA, nrow = 3, ncol = 2)
emiss_child[1,] <- seqstatf(child_seq[, 1:5])[, 2] + 1
emiss_child[2,] <- seqstatf(child_seq[, 6:10])[, 2] + 1
emiss_child[3,] <- seqstatf(child_seq[, 11:16])[, 2] + 1
emiss_child <- emiss_child / rowSums(emiss_child)
emiss_left <- matrix(NA, nrow = 3, ncol = 2)
```

```
emiss_left[1,] <- seqstatf(left_seq[, 1:5])[, 2] + 1
emiss_left[2,] <- seqstatf(left_seq[, 6:10])[, 2] + 1
emiss_left[3,] <- seqstatf(left_seq[, 11:16])[, 2] + 1
emiss_left <- emiss_left / rowSums(emiss_left)
# Starting values for transition matrix
trans <- matrix(c(0.9, 0.07, 0.03,
            0, 0.9, 0.1,
            0, 0, 1),
        nrow = 3, ncol = 3, byrow = TRUE)
# Starting values for initial state probabilities
inits <- c(0.9, 0.09, 0.01)
# HMM with own starting values
init_hmm_bf2 <- build_hmm(
    observations = list(marr_seq, child_seq, left_seq),
    transition_probs = trans,
    emission_probs = list(emiss_marr, emiss_child, emiss_left),
    initial_probs = inits)
```

build_lcm

## Description

Function build_lcm is a shortcut for constructing a latent class model as a restricted case of an mhmm object.

## Usage

```
build_lcm(
    observations,
    n_clusters,
    emission_probs,
    formula,
    data,
    coefficients,
    cluster_names = NULL,
    channel_names = NULL
)
```


## Arguments

observations An stslist object (see seqdef) containing the sequences, or a list of such objects (one for each channel).
n_clusters A scalar giving the number of clusters/submodels (not used if starting values for model parameters are given with emission_probs).

| emission_probs | A matrix containing emission probabilities for each class by rows, or in case <br> of multichannel data a list of such matrices. Note that the matrices must have <br> dimensions kx s where k is the number of latent classes and s is the number <br> of unique symbols (observed states) in the data. Emission probabilities should <br> follow the ordering of the alphabet of observations (alphabet (observations), <br> returned as symbol_names). |
| :--- | :--- |
| formula | Covariates as an object of class formula, left side omitted. |
| data | An optional data frame, list or environment containing the variables in the model. <br> If not found in data, the variables are taken from environment (formula). |
| coefficients | An optional $k x l$ matrix of regression coefficients for time-constant covariates <br> for mixture probabilities, where $l$ is the number of clusters and $k$ is the number <br> of covariates. A logit-link is used for mixture probabilities. The first column is |
| set to zero. |  |$\quad$| A vector of optional names for the classes/clusters. |
| :--- |

## Value

Object of class mhmm with the following elements:
observations State sequence object or a list of such containing the data.
transition_probs A matrix of transition probabilities.
emission_probs A matrix or a list of matrices of emission probabilities.
initial_probs A vector of initial probabilities.
coefficients A matrix of parameter coefficients for covariates (covariates in rows, clusters in columns).
$X$ Covariate values for each subject.
cluster_names Names for clusters.
state_names Names for hidden states.
symbol_names Names for observed states.
channel_names Names for channels of sequence data
length_of_sequences (Maximum) length of sequences.
n _sequences Number of sequences.
n _symbols Number of observed states (in each channel).
n_states Number of hidden states.
n_channels Number of channels.
n_covariates Number of covariates.
n_clusters Number of clusters.

## See Also

fit_model for estimating model parameters; summary.mhmm for a summary of a mixture model; separate_mhmm for organizing an mhmm object into a list of separate hmm objects; and plot.mhmm for plotting mixture models.

## Examples

```
# Simulate observations from two classes
set.seed(123)
obs <- seqdef(rbind(
    matrix(sample(letters[1:3], 500, TRUE, prob = c(0.1, 0.6, 0.3)), 50, 10),
    matrix(sample(letters[1:3], 200, TRUE, prob = c(0.4, 0.4, 0.2)), 20, 10)))
# Initialize the model
set.seed(9087)
model <- build_lcm(obs, n_clusters = 2)
# Estimate model parameters
fit <- fit_model(model)
# How many of the observations were correctly classified:
sum(summary(fit$model)$most_probable_cluster == rep(c("Class 2", "Class 1"), times = c(500, 200)))
###############################################################
## Not run:
# LCM for longitudinal data
# Define sequence data
data("mvad", package = "TraMineR")
mvad_alphabet <- c("employment", "FE", "HE", "joblessness", "school",
    "training")
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet, states = mvad_scodes,
    labels = mvad_labels, xtstep = 6)
# Initialize the LCM with random starting values
set.seed(7654)
init_lcm_mvad1 <- build_lcm(observations = mvad_seq,
    n_clusters = 2, formula = ~male, data = mvad)
# Own starting values for emission probabilities
emiss <- rbind(rep(1/6, 6), rep(1/6, 6))
# LCM with own starting values
init_lcm_mvad2 <- build_lcm(observations = mvad_seq,
    emission_probs = emiss, formula = ~male, data = mvad)
# Estimate model parameters (EM algorithm with random restarts)
lcm_mvad <- fit_model(init_lcm_mvad1,
    control_em = list(restart = list(times = 5)))$model
# Plot the LCM
plot(lcm_mvad, interactive = FALSE, ncol = 2)
######################################################################
```

```
    # Binomial regression (comparison to glm)
    require("MASS")
    data("birthwt")
    model <- build_lcm(
        observations = seqdef(birthwt$low), emission_probs = diag(2),
        formula = ~age + lwt + smoke + ht, data = birthwt)
fit <- fit_model(model)
summary(fit$model)
summary(glm(low ~ age + lwt + smoke + ht, binomial, data = birthwt))
# Multinomial regression (comparison to multinom)
require("nnet")
set.seed(123)
n <- 100
x <- cbind(1, x1 = runif(n, 0, 1), x2 = runif(n, 0, 1))
coefs <- cbind(0,c(-2, 5, -2), c(0, -2, 2))
pr <- exp(X %*% coefs) + rnorm(n*3)
pr <- pr/rowSums(pr)
y <- apply(pr, 1, which.max)
table(y)
model <- build_lcm(
    observations = seqdef(y), emission_probs = diag(3),
    formula = ~x1 + x2, data = data.frame(X[, -1]))
fit <- fit_model(model)
summary(fit$model)
summary(multinom(y ~ x1 + x2, data = data.frame(X[,-1])))
## End(Not run)
```


## Description

Function build_mhmm constructs a mixture hidden Markov model object of class mhmm.

## Usage

build_mhmm( observations, n_states, transition_probs, emission_probs, initial_probs,

```
    formula,
    data,
    coefficients,
    cluster_names = NULL,
    state_names = NULL,
    channel_names = NULL,
)
```


## Arguments

observations An stslist object (see seqdef) containing the sequences, or a list of such objects (one for each channel).
n_states A numerical vector giving the number of hidden states in each submodel (not used if starting values for model parameters are given with initial_probs, transition_probs, or emission_probs).
transition_probs
A list of matrices of transition probabilities for the submodel of each cluster.
emission_probs A list which contains matrices of emission probabilities or a list of such objects (one for each channel) for the submodel of each cluster. Note that the matrices must have dimensions $m x s$ where $m$ is the number of hidden states and $s$ is the number of unique symbols (observed states) in the data. Emission probabilities should follow the ordering of the alphabet of observations (alphabet(observations), returned as symbol_names).
initial_probs A list which contains vectors of initial state probabilities for the submodel of each cluster.
formula Covariates as an object of class formula, left side omitted.
data An optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment (formula).
coefficients An optional $k x l$ matrix of regression coefficients for time-constant covariates for mixture probabilities, where $l$ is the number of clusters and $k$ is the number of covariates. A logit-link is used for mixture probabilities. The first column is set to zero.
cluster_names A vector of optional names for the clusters.
state_names A list of optional labels for the hidden states. If NULL, the state names are taken as row names of transition matrices. If this is also NULL, numbered states are used.
channel_names A vector of optional names for the channels.
... Additional arguments to simulate_transition_probs.

## Details

The returned model contains some attributes such as nobs and df , which define the number of observations in the model and the number of estimable model parameters, used in computing BIC. When computing nobs for a multichannel model with $C$ channels, each observed value in a single channel amounts to $1 / C$ observation, i.e. a fully observed time point for a single sequence amounts
to one observation. For the degrees of freedom df , zero probabilities of the initial model are defined as structural zeroes.

## Value

Object of class mhmm with following elements:
observations State sequence object or a list of such containing the data.
transition_probs A matrix of transition probabilities.
emission_probs A matrix or a list of matrices of emission probabilities.
initial_probs A vector of initial probabilities.
coefficients A matrix of parameter coefficients for covariates (covariates in rows, clusters in columns).
X Covariate values for each subject.
cluster_names Names for clusters.
state_names Names for hidden states.
symbol_names Names for observed states.
channel_names Names for channels of sequence data
length_of_sequences (Maximum) length of sequences.
n _sequences Number of sequences.
n _symbols Number of observed states (in each channel).
n_states Number of hidden states.
n_channels Number of channels.
n_covariates Number of covariates.
n_clusters Number of clusters.

## References

Helske S. and Helske J. (2019). Mixture Hidden Markov Models for Sequence Data: The seqHMM Package in R, Journal of Statistical Software, 88(3), 1-32. doi:10.18637/jss.v088.i03

## See Also

fit_model for fitting mixture Hidden Markov models; summary .mhmm for a summary of a MHMM; separate_mhmm for reorganizing a MHMM into a list of separate hidden Markov models; and plot. mhmm for plotting mhmm objects.

## Examples

```
data("biofam3c")
## Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
```

```
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
## Choosing colors
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
## MHMM with random starting values, no covariates
set.seed(468)
init_mhmm_bf1 <- build_mhmm(
    observations = list(marr_seq, child_seq, left_seq),
    n_states = c(4, 4, 6),
    channel_names = c("Marriage", "Parenthood", "Residence"))
## Starting values for emission probabilities
# Cluster 1
B1_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
            0.8, 0.1, 0.1,
            0.3, 0.6, 0.1, # High probability for married
            0.3, 0.3, 0.4), # High probability for divorced
    nrow = 4, ncol = 3, byrow = TRUE)
B1_child <- matrix(
    c(0.9, 0.1, # High probability for childless
            0.9, 0.1,
            0.9, 0.1,
            0.9, 0.1),
    nrow = 4, ncol = 2, byrow = TRUE)
B1_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
            0.1, 0.9, # High probability for having left home
            0.1, 0.9,
            0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
# Cluster 2
B2_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
            0.8, 0.1, 0.1,
            0.1, 0.8, 0.1, # High probability for married
            0.7, 0.2, 0.1),
    nrow = 4, ncol = 3, byrow = TRUE)
B2_child <- matrix(
    c(0.9, 0.1, # High probability for childless
```

```
        0.9, 0.1,
        0.9, 0.1,
        0.1, 0.9)
    nrow = 4, ncol = 2, byrow = TRUE)
B2_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9,
        0.1, 0.9,
        0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
# Cluster 3
B3_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
        0.8, 0.1, 0.1,
        0.8, 0.1, 0.1,
        0.1, 0.8, 0.1, # High probability for married
        0.3, 0.4, 0.3,
        0.1, 0.1, 0.8), # High probability for divorced
    nrow = 6, ncol = 3, byrow = TRUE)
B3_child <- matrix(
    c(0.9, 0.1, # High probability for childless
        0.9, 0.1,
        0.5, 0.5,
        0.5, 0.5,
        0.5, 0.5,
        0.1, 0.9),
    nrow = 6, ncol = 2, byrow = TRUE)
B3_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9,
        0.5, 0.5,
        0.5, 0.5,
        0.1, 0.9,
        0.1, 0.9),
    nrow = 6, ncol = 2, byrow = TRUE)
# Starting values for transition matrices
A1 <- matrix(
    c(0.80, 0.16, 0.03, 0.01,
        0, 0.90, 0.07, 0.03,
        0, 0, 0.90, 0.10,
        0, 0, 0, 1)
    nrow = 4, ncol = 4, byrow = TRUE)
A2 <- matrix(
    c(0.80, 0.10, 0.05, 0.03, 0.01, 0.01,
        0, 0.70, 0.10, 0.10, 0.05, 0.05,
        0, 0, 0.85, 0.01, 0.10, 0.04,
```

```
        0, 0, 0, 0.90, 0.05, 0.05,
        0, 0, 0, 0, 0.90, 0.10,
        0, 0, 0, 0, 0, 1),
    nrow = 6, ncol = 6, byrow = TRUE)
# Starting values for initial state probabilities
initial_probs1 <- c(0.9, 0.07, 0.02, 0.01)
initial_probs2 <- c(0.9, 0.04, 0.03, 0.01, 0.01, 0.01)
# Birth cohort
biofam3c$covariates$cohort <- cut(biofam3c$covariates$birthyr, c(1908, 1935, 1945, 1957))
biofam3c$covariates$cohort <- factor(
    biofam3c$covariates$cohort, labels=c("1909-1935", "1936-1945", "1946-1957"))
## MHMM with own starting values and covariates
init_mhmm_bf2 <- build_mhmm(
    observations = list(marr_seq, child_seq, left_seq),
    initial_probs = list(initial_probs1, initial_probs1, initial_probs2),
    transition_probs = list(A1, A1, A2),
    emission_probs = list(list(B1_marr, B1_child, B1_left),
        list(B2_marr, B2_child, B2_left),
        list(B3_marr, B3_child, B3_left)),
    formula = ~sex + cohort, data = biofam3c$covariates,
    cluster_names = c("Cluster 1", "Cluster 2", "Cluster 3"),
    channel_names = c("Marriage", "Parenthood", "Residence"),
    state_names = list(paste("State", 1:4), paste("State", 1:4),
                paste("State", 1:6)))
```

    build_mm Build a Markov Model
    
## Description

Function build_mm builds and automatically estimates a Markov model. It is also a shortcut for constructing a Markov model as a restricted case of an hmm object.

## Usage

build_mm(observations)

## Arguments

observations An stslist object (see seqdef) containing the sequences.

## Details

Unlike the other build functions in seqHMM, the build_mm function automatically estimates the model parameters. In case of no missing values, initial and transition probabilities are directly estimated from the observed initial state probabilities and transition counts. In ase of missing values, the EM algorithm is run once.

## Value

Object of class hmm with following elements:
observations State sequence object or a list of such containing the data.
transition_probs A matrix of transition probabilities.
emission_probs A matrix or a list of matrices of emission probabilities.
initial_probs A vector of initial probabilities.
state_names Names for hidden states.
symbol_names Names for observed states.
channel_names Names for channels of sequence data.
length_of_sequences (Maximum) length of sequences.
n _sequences Number of sequences.
n _symbols Number of observed states (in each channel).
n_states Number of hidden states.
n_channels Number of channels.

## See Also

plot. hmm for plotting the model.

## Examples

```
# Construct sequence data
data("mvad", package = "TraMineR")
mvad_alphabet <-
    c("employment", "FE", "HE", "joblessness", "school", "training")
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet,
    states = mvad_scodes, labels = mvad_labels, xtstep = 6)
# Define a color palette for the sequence data
attr(mvad_seq, "cpal") <- colorpalette[[6]]
# Estimate the Markov model
mm_mvad <- build_mm(observations = mvad_seq)
```


## Description

Function build_mmm is a shortcut for constructing a mixture Markov model as a restricted case of an mhmm object.

## Usage

```
build_mmm(
    observations,
    n_clusters,
    transition_probs,
    initial_probs,
    formula,
    data,
    coefficients,
    cluster_names = NULL,
    )
```


## Arguments

observations An stslist object (see seqdef) containing the sequences.
n_clusters A scalar giving the number of clusters/submodels (not used if starting values for model parameters are given with initial_probs and transition_probs).
transition_probs
A list of matrices of transition probabilities for submodels of each cluster.
initial_probs A list which contains vectors of initial state probabilities for submodels of each cluster.
formula Covariates as an object of class formula, left side omitted.
data An optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment (formula).
coefficients An optional $k x l$ matrix of regression coefficients for time-constant covariates for mixture probabilities, where $l$ is the number of clusters and $k$ is the number of covariates. A logit-link is used for mixture probabilities. The first column is set to zero.
cluster_names A vector of optional names for the clusters.
...
Additional arguments to simulate_transition_probs.

## Value

Object of class mhmm with following elements:
observations State sequence object or a list of such containing the data.
transition_probs A matrix of transition probabilities.
emission_probs A matrix or a list of matrices of emission probabilities.
initial_probs A vector of initial probabilities.
coefficients A matrix of parameter coefficients for covariates (covariates in rows, clusters in columns).

X Covariate values for each subject.
cluster_names Names for clusters.
state_names Names for hidden states.
symbol_names Names for observed states.
channel_names Names for channels of sequence data
length_of_sequences (Maximum) length of sequences.
n _sequences Number of sequences.
n _symbols Number of observed states (in each channel).
n_states Number of hidden states.
n_channels Number of channels.
n_covariates Number of covariates.
n_clusters Number of clusters.

## See Also

fit_model for estimating model parameters; summary.mhmm for a summary of a mixture model; separate_mhmm for organizing an mhmm object into a list of separate hmm objects; and plot.mhmm for plotting mixture models.

## Examples

```
# Define sequence data
data("mvad", package = "TraMineR")
mvad_alphabet <- c("employment", "FE", "HE", "joblessness", "school",
    "training")
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet, states = mvad_scodes,
    labels = mvad_labels, xtstep = 6)
# Initialize the MMM
set.seed(123)
mmm_mvad <- build_mmm(observations = mvad_seq,
```

```
    n_clusters = 2,
    formula = ~male, data = mvad)
    ## Not run:
    # Estimate model parameters
    mmm_mvad <- fit_model(mmm_mvad)$model
    # Plot model (both clusters in the same plot)
    require(igraph)
    plot(mmm_mvad, interactive = FALSE,
    # Modify legend position and properties
    with.legend = "right", legend.prop = 0.3, cex.legend = 1.2,
    # Define vertex layout
    layout = layout_as_star,
    # Modify edge properties
    edge.label = NA, edge.arrow.size = 0.8, edge.curved = 0.2,
    # Modify vertex label positions (initial probabilities)
    vertex.label.pos = c("left", "right", "right", "left", "left", "right"))
    # Summary of the MMM
    summary (mmm_mvad)
    ## End(Not run)
```

    colorpalette
        Color palettes
    
## Description

A list containing ready defined color palettes with distinct colors using iWantHue. By default, seqHMM uses these palettes when assigning colors.

## Format

A list with 200 color palettes.

## Source

iWantHue web page https://medialab.github.io/iwanthue/

## See Also

plot_colors for visualization of color palettes. Implementations of iWantHue for R:

- https://github.com/hoesler/rwantshue
- https://github.com/johnbaums/hues


## Examples

```
data("colorpalette")
# Color palette with 9 colors
colorpalette[[9]]
# Color palette with 24 colors
colorpalette[[24]]
```

    estimate_coef Estimate Regression Coefficients of Mixture Hidden Markov Models
    
## Description

Function estimate_coef estimates the regression coefficients of mixture hidden Markov models and its restricted variants while keeping other parameters fixed.

## Usage

estimate_coef(model, threads = 1)

## Arguments

model An object of class hmm or mhmm.
threads Number of threads to use in parallel computing. The default is 1 .

fit_model $\quad$| Estimate Parameters of (Mixture) Hidden Markov Models and Their |
| :--- |
| Restricted Variants | Restricted Variants

## Description

Function fit_model estimates the parameters of mixture hidden Markov models and its restricted variants using maximimum likelihood. Initial values for estimation are taken from the corresponding components of the model with preservation of original zero probabilities.

## Usage

```
fit_model(
        model,
        em_step = TRUE,
    global_step = FALSE,
    local_step = FALSE,
    control_em = list(),
    control_global = list(),
    control_local = list(),
```

```
    lb,
    ub,
    threads = 1,
    log_space = FALSE,
    constraints = NULL,
    fixed_inits = NULL,
    fixed_emissions = NULL,
    fixed_transitions = NULL,
)
```


## Arguments

## model

An object of class hmm or mhmm.

```
em_step
```

Logical. Whether or not to use the EM algorithm at the start of the parameter estimation. The default is TRUE.
global_step Logical. Whether or not to use global optimization via nloptr (possibly after the EM step). The default is FALSE.
local_step Logical. Whether or not to use local optimization via nloptr (possibly after the EM and/or global steps). The default is FALSE.
control_em Optional list of control parameters for the EM algorithm. Possible arguments are
maxeval The maximum number of iterations, the default is 1000 . Note that iteration counter starts with -1 so with maxeval=1 you get already two iterations. This is for backward compatibility reasons.
print_level The level of printing. Possible values are 0 (prints nothing), 1 (prints information at the start and the end of the algorithm), 2 (prints at every iteration), and for mixture models 3 (print also during optimization of coefficients).
reltol Relative tolerance for convergence defined as $\left(\log L i k_{n} e w-\log L i k_{o} l d\right) /\left(a b s\left(\log L i k_{o} l d\right)+\right.$ $0.1)$. The default is $1 \mathrm{e}-10$.
restart A list containing options for possible EM restarts with the following components:
times Number of restarts of the EM algorithm using random initial values. The default is 0 , i.e. no restarts.
transition Logical. Should the original transition probabilities be varied? The default is TRUE.
emission Logical. Should the original emission probabilities be varied? The default is TRUE.
sd Standard deviation for rnorm used in randomization. The default is 0.25 .
maxeval Maximum number of iterations, the default is control_em\$maxeval
print_level Level of printing in restarted EM steps. The default is control_em\$print_level.
reltol Relative tolerance for convergence at restarted EM steps. The default is control_em\$reltol. If the relative change of the final model of the restart phase is larger than the tolerance for the original EM phase, the
final model is re-estimated with the original reltol and maxeval at the end of the EM step.
n_optimum Save the log-likelihood values of the n_optimum best models (from all estimated models including the the first EM run.). The default is min(times $+1,25$ ).
use_original If TRUE. Use the initial values of the input model as starting points for the permutations. Otherwise permute the results of the first EM run.
control_global Optional list of additional arguments for nloptr argument opts. The default values are
algorithm "NLOPT_GD_MLSL_LDS"
local_opts list(algorithm = "NLOPT_LD_LBFGS", ftol_rel = 1e-6, xtol_rel $=1 \mathrm{e}-4$ )
maxeval 10000 (maximum number of iterations in global optimization algorithm.)
maxtime 60 (maximum time for global optimization. Set to 0 for unlimited time.)
control_local Optional list of additional arguments for nloptr argument opts. The default values are
algorithm "NLOPT_LD_LBFGS"
ftol_rel 1e-10
xtol_rel 1e-8
maxeval 10000 (maximum number of iterations)
lb, ub Lower and upper bounds for parameters in Softmax parameterization. The default interval is $[\operatorname{pmin}(-25,2 *$ initialvalues $), \operatorname{pmax}(25,2 *$ initialvalues $)$ ], except for gamma coefficients, where the scale of covariates is taken into account. Note that it might still be a good idea to scale covariates around unit scale. Bounds are used only in the global optimization step.
threads Number of threads to use in parallel computing. The default is 1.
log_space Make computations using log-space instead of scaling for greater numerical stability at a cost of decreased computational performance. The default is FALSE.
constraints Integer vector defining equality constraints for emission distributions. Not supported for EM algorithm. See details.
fixed_inits Can be used to fix some of the probabilities to their initial values. Should have same structure as model\$initial_probs, where each element is either TRUE (fixed) or FALSE (to be estimated). Note that zero probabilities are always fixed to 0 . Not supported for EM algorithm. See details.
fixed_emissions
Can be used to fix some of the probabilities to their initial values. Should have same structure as model\$emission_probs, where each element is either TRUE (fixed) or FALSE (to be estimated). Note that zero probabilities are always fixed to 0 . Not supported for EM algorithm. See details.
fixed_transitions
Can be used to fix some of the probabilities to their initial values. Should have same structure as model\$transition_probs, where each element is either TRUE (fixed) or FALSE (to be estimated). Note that zero probabilities are always fixed to 0 . Not supported for EM algorithm. See details.

Additional arguments to nloptr.

## Details

The fitting function provides three estimation steps: 1) EM algorithm, 2) global optimization, and 3) local optimization. The user can call for one method or any combination of these steps, but should note that they are preformed in the above-mentioned order. The results from a former step are used as starting values in a latter, except for some of global optimization algorithms (such as MLSL and StoGO) which only use initial values for setting up the boundaries for the optimization.

It is possible to rerun the EM algorithm automatically using random starting values based on the first run of EM. Number of restarts is defined by the restart argument in control_em. As the EM algorithm is relatively fast, this method might be preferred option compared to the proper global optimization strategy of step 2.

The default global optimization method (triggered via global_step = TRUE) is the multilevel singlelinkage method (MLSL) with the LDS modification (NLOPT_GD_MLSL_LDS as algorithm in control_global), with L-BFGS as the local optimizer. The MLSL method draws random starting points and performs a local optimization from each. The LDS modification uses low-discrepancy sequences instead of pseudo-random numbers as starting points and should improve the convergence rate. In order to reduce the computation time spent on non-global optima, the convergence tolerance of the local optimizer is set relatively large. At step 3, a local optimization (L-BFGS by default) is run with a lower tolerance to find the optimum with high precision.

There are some theoretical guarantees that the MLSL method used as the default optimizer in step 2 shoud find all local optima in a finite number of local optimizations. Of course, it might not always succeed in a reasonable time. The EM algorithm can help in finding good boundaries for the search, especially with good starting values, but in some cases it can mislead. A good strategy is to try a couple of different fitting options with different combinations of the methods: e.g. all steps, only global and local steps, and a few evaluations of EM followed by global and local optimization.
By default, the estimation time is limited to 60 seconds in global optimization step, so it is advisable to change the default settings for the proper global optimization.
Any algorithm available in the nloptr function can be used for the global and local steps.
Equality constraints for emission distributions can be defined using the argument constraints. This should be a vector with length equal to the number of states, with numbers starting from 1 and increasing for each unique row of the emission probability matrix. For example in case of five states with emissions of first and third states being equal, constraints $=c(1,2,1,3,4)$. Similarly, some of the model parameters can be fixed to their initial values by using arguments fixed_inits, fixed_emissions, and fixed_transitions, where the structure of the arguments should be same as the corresponding model components, so that TRUE value means that the parameter should be fixed and FALSE otherwise (it is still treated as fixed if it is zero though). For both types of constrains, only numerical optimisation (local or global) is available, and currently the gradients are computed numerically (if needed) in these cases.

## Value

logLik Log-likelihood of the estimated model.
em_results Results after the EM step: log-likelihood (logLik), number of iterations (iterations), relative change in log-likelihoods between the last two iterations (change), and the log-likelihoods of the n_optimum best models after the EM step (best_opt_restart).
global_results Results after the global step.
local_results Results after the local step.
call The matched function call.

## References

Helske S. and Helske J. (2019). Mixture Hidden Markov Models for Sequence Data: The seqHMM Package in R, Journal of Statistical Software, 88(3), 1-32. doi:10.18637/jss.v088.i03

## See Also

build_hmm, build_mhmm, build_mm, build_mmm, and build_lcm for constructing different types of models; summary.mhmm for a summary of a MHMM; separate_mhmm for reorganizing a MHMM into a list of separate hidden Markov models; plot. hmm and plot.mhmm for plotting model objects; and ssplot and mssplot for plotting stacked sequence plots of hmm and mhmm objects.

## Examples

```
# Left-to-right HMM with equality constraint:
set.seed(1)
# Transition matrix
# Either stay or move to next state
A <- diag(c(0.9, 0.95, 0.95, 1))
A[1, 2] <- 0.1
A[2, 3] <- 0.05
A[3, 4] <- 0.05
# Emission matrix, rows 1 and 3 equal
B <- rbind(
    c(0.4,0.2,0.3,0.1),
        c(0.1,0.5,0.1,0.3),
        c(0.4,0.2,0.3,0.1),
        c(0, 0.2,0.4,0.4))
# Start from first state
init <- c(1,0,0,0)
# Simulate sequences
sim <- simulate_hmm(n_sequences = 100,
    sequence_length = 20, init, A, B)
# initial model, use true values as inits for faster estimation here
```

```
model <- build_hmm(sim$observations, init = init, trans = A, emiss = B)
# estimate the model subject to constraints:
# First and third row of emission matrix are equal (see details)
fit <- fit_model(model, constraints = c(1, 2, 1, 3),
    em_step = FALSE, local_step = TRUE)
fit$model
## Fix some emissions:
fixB <- matrix(FALSE, 4, 4)
fixB[2, 1] <- fixB[1, 3] <- TRUE # these are fixed to their initial values
fit <- fit_model(model, fixed_emissions = fixB,
    em_step = FALSE, local_step = TRUE)
fit$model$emission_probs
# Hidden Markov model for mvad data
data("mvad", package = "TraMineR")
mvad_alphabet <-
    c("employment", "FE", "HE", "joblessness", "school", "training")
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet,
    states = mvad_scodes, labels = mvad_labels, xtstep = 6)
attr(mvad_seq, "cpal") <- colorpalette[[6]]
# Starting values for the emission matrix
emiss <- matrix(
    c(0.05, 0.05, 0.05, 0.05, 0.75, 0.05, # SC
        0.05, 0.75, 0.05, 0.05, 0.05, 0.05, # FE
        0.05, 0.05, 0.05, 0.4, 0.05, 0.4, # JL, TR
        0.05, 0.05, 0.75, 0.05, 0.05, 0.05, # HE
        0.75, 0.05, 0.05, 0.05, 0.05, 0.05),# EM
    nrow = 5, ncol = 6, byrow = TRUE)
# Starting values for the transition matrix
trans <- matrix(0.025, 5, 5)
diag(trans) <- 0.9
# Starting values for initial state probabilities
initial_probs <- c(0.2, 0.2, 0.2, 0.2, 0.2)
# Building a hidden Markov model
init_hmm_mvad <- build_hmm(observations = mvad_seq,
    transition_probs = trans, emission_probs = emiss,
    initial_probs = initial_probs)
## Not run:
set.seed(21)
```

```
fit_hmm_mvad <- fit_model(init_hmm_mvad, control_em = list(restart = list(times = 50)))
hmm_mvad <- fit_hmm_mvad$model
## End(Not run)
# save time, load the previously estimated model
data("hmm_mvad")
# Markov model
# Note: build_mm estimates model parameters from observations,
# no need for estimating with fit_model
mm_mvad <- build_mm(observations = mvad_seq)
# Comparing likelihoods, MM fits better
logLik(hmm_mvad)
logLik(mm_mvad)
## Not run:
require("igraph") #for layout_in_circle
plot(mm_mvad, layout = layout_in_circle, legend.prop = 0.3,
    edge.curved = 0.3, edge.label = NA,
    vertex.label.pos = c(0, 0, pi, pi, pi, 0))
###############################################################
#' # Three-state three-channel hidden Markov model
# See ?hmm_biofam for five-state version
data("biofam3c")
# Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
# Define colors
attr(marr_seq, "cpal") <- c("violetred2", "darkgoldenrod2", "darkmagenta")
attr(child_seq, "cpal") <- c("darkseagreen1", "coral3")
attr(left_seq, "cpal") <- c("lightblue", "red3")
# Starting values for emission matrices
emiss_marr <- matrix(NA, nrow = 3, ncol = 3)
emiss_marr[1,] <- seqstatf(marr_seq[, 1:5])[, 2] + 1
emiss_marr[2,] <- seqstatf(marr_seq[, 6:10])[, 2] + 1
emiss_marr[3,] <- seqstatf(marr_seq[, 11:16])[, 2] + 1
emiss_marr <- emiss_marr / rowSums(emiss_marr)
```

```
emiss_child <- matrix(NA, nrow = 3, ncol = 2)
emiss_child[1,] <- seqstatf(child_seq[, 1:5])[, 2] + 1
emiss_child[2,] <- seqstatf(child_seq[, 6:10])[, 2] + 1
emiss_child[3,] <- seqstatf(child_seq[, 11:16])[, 2] + 1
emiss_child <- emiss_child / rowSums(emiss_child)
emiss_left <- matrix(NA, nrow = 3, ncol = 2)
emiss_left[1,] <- seqstatf(left_seq[, 1:5])[, 2] + 1
emiss_left[2,] <- seqstatf(left_seq[, 6:10])[, 2] + 1
emiss_left[3,] <- seqstatf(left_seq[, 11:16])[, 2] + 1
emiss_left <- emiss_left / rowSums(emiss_left)
# Starting values for transition matrix
trans <- matrix(c(0.9, 0.07, 0.03,
    0, 0.9, 0.1,
    0, 0, 1), nrow = 3, ncol = 3, byrow = TRUE)
# Starting values for initial state probabilities
inits <- c(0.9, 0.09, 0.01)
# Building hidden Markov model with initial parameter values
init_hmm_bf <- build_hmm(
    observations = list(marr_seq, child_seq, left_seq),
    transition_probs = trans,
    emission_probs = list(emiss_marr, emiss_child, emiss_left),
    initial_probs = inits)
# Fitting the model with different optimization schemes
# Only EM with default values
hmm_1 <- fit_model(init_hmm_bf)
hmm_1$logLik # -24179.1
# Only L-BFGS
hmm_2 <- fit_model(init_hmm_bf, em_step = FALSE, local_step = TRUE)
hmm_2$logLik # -22267.75
# Global optimization via MLSL_LDS with L-BFGS as local optimizer and final polisher
# This can be slow, use parallel computing by adjusting threads argument
# (here threads = 1 for portability issues)
hmm_3 <- fit_model(
    init_hmm_bf, em_step = FALSE, global_step = TRUE, local_step = TRUE,
    control_global = list(maxeval = 5000, maxtime = 0), threads = 1)
hmm_3$logLik # -21675.42
# EM with restarts, much faster than MLSL
set.seed(123)
hmm_4 <- fit_model(init_hmm_bf, control_em = list(restart = list(times = 5)))
hmm_4$logLik # -21675.4
# Global optimization via StoGO with L-BFGS as final polisher
# This can be slow, use parallel computing by adjusting threads argument
```

```
# (here threads = 1 for portability issues)
set.seed(123)
hmm_5 <- fit_model(
    init_hmm_bf, em_step = FALSE, global_step = TRUE, local_step = TRUE,
    lb = -50, ub = 50, control_global = list(algorithm = "NLOPT_GD_STOGO",
    maxeval = 2500, maxtime = 0), threads = 1)
hmm_5$logLik # -21675.4
###############################################################
# Mixture HMM
data("biofam3c")
## Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
## Choosing colors
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
## Starting values for emission probabilities
# Cluster 1
B1_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
        0.8, 0.1, 0.1,
        0.3, 0.6, 0.1, # High probability for married
        0.3, 0.3, 0.4), # High probability for divorced
    nrow = 4, ncol = 3, byrow = TRUE)
B1_child <- matrix(
    c(0.9, 0.1, # High probability for childless
        0.9, 0.1,
        0.9, 0.1,
        0.9, 0.1),
    nrow = 4, ncol = 2, byrow = TRUE)
B1_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9, # High probability for having left home
        0.1, 0.9,
        0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
# Cluster 2
B2_marr <- matrix(
```

```
    c(0.8, 0.1, 0.1, # High probability for single
        0.8, 0.1, 0.1,
        0.1, 0.8, 0.1, # High probability for married
        0.7, 0.2, 0.1),
    nrow = 4, ncol = 3, byrow = TRUE)
B2_child <- matrix(
    c(0.9, 0.1, # High probability for childless
        0.9, 0.1,
        0.9, 0.1,
        0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
B2_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9,
        0.1, 0.9,
        0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
# Cluster 3
B3_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
        0.8, 0.1, 0.1,
        0.8, 0.1, 0.1,
        0.1, 0.8, 0.1, # High probability for married
        0.3, 0.4, 0.3,
        0.1, 0.1, 0.8), # High probability for divorced
    nrow = 6, ncol = 3, byrow = TRUE)
B3_child <- matrix(
    c(0.9, 0.1, # High probability for childless
        0.9, 0.1,
        0.5, 0.5,
        0.5, 0.5,
        0.5, 0.5,
        0.1, 0.9),
    nrow = 6, ncol = 2, byrow = TRUE)
B3_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9,
        0.5, 0.5,
        0.5, 0.5,
        0.1, 0.9,
        0.1, 0.9)
    nrow = 6, ncol = 2, byrow = TRUE)
# Starting values for transition matrices
A1 <- matrix(
    c(0.80, 0.16, 0.03, 0.01,
        0, 0.90, 0.07, 0.03,
```

```
        0, 0, 0.90, 0.10,
        0, 0, 0, 1),
    nrow = 4, ncol = 4, byrow = TRUE)
A2 <- matrix(
    c(0.80, 0.10, 0.05, 0.03, 0.01, 0.01,
        0, 0.70, 0.10, 0.10, 0.05, 0.05,
        0, 0, 0.85, 0.01, 0.10, 0.04,
        0, 0, 0, 0.90, 0.05, 0.05,
        0, 0, 0, 0, 0.90, 0.10,
        0, 0, 0, 0, 0, 1),
    nrow = 6, ncol = 6, byrow = TRUE)
# Starting values for initial state probabilities
initial_probs1 <- c(0.9, 0.07, 0.02, 0.01)
initial_probs2 <- c(0.9, 0.04, 0.03, 0.01, 0.01, 0.01)
# Birth cohort
biofam3c$covariates$cohort <- cut(biofam3c$covariates$birthyr, c(1908, 1935, 1945, 1957))
biofam3c$covariates$cohort <- factor(
    biofam3c$covariates$cohort, labels=c("1909-1935", "1936-1945", "1946-1957"))
# Build mixture HMM
init_mhmm_bf <- build_mhmm(
    observations = list(marr_seq, child_seq, left_seq),
    initial_probs = list(initial_probs1, initial_probs1, initial_probs2),
    transition_probs = list(A1, A1, A2),
    emission_probs = list(list(B1_marr, B1_child, B1_left),
        list(B2_marr, B2_child, B2_left),
        list(B3_marr, B3_child, B3_left)),
    formula = ~sex + cohort, data = biofam3c$covariates,
    channel_names = c("Marriage", "Parenthood", "Residence"))
# Fitting the model with different settings
# Only EM with default values
mhmm_1 <- fit_model(init_mhmm_bf)
mhmm_1$logLik # -12713.08
# Only L-BFGS
mhmm_2 <- fit_model(init_mhmm_bf, em_step = FALSE, local_step = TRUE)
mhmm_2$logLik # -12966.51
# Use EM with multiple restarts
set.seed(123)
mhmm_3 <- fit_model(init_mhmm_bf, control_em = list(restart = list(times = 5, transition = FALSE)))
mhmm_3$logLik # -12713.08
## End(Not run)
```


## Description

The forward_backward function computes scaled forward and backward probabilities of a hidden Markov model.

## Usage

forward_backward(model, forward_only = FALSE, log_space = FALSE, threads = 1)

## Arguments

| model | Object of class hmm or mhmm. |
| :--- | :--- |
| forward_only | If TRUE, only forward probabilities are computed. The default is FALSE. |
| log_space | Compute forward and backward probabilities in logarithmic scale instead of <br> scaling. The default is FALSE. |
| threads | Number of threads used in parallel computing. The default is 1. |

## Value

List with components
forward_probs If log_space = FALSE, scaled forward probabilities, i.e. probability of state given observations up to that time point. If log_space $=$ TRUE, logarithms of non-scaled forward probabilities.
backward_probs Scaled backward probabilities (log_space = FALSE), or logarithms of non-scaled backward probabilities(log_space = TRUE).
scaling_factors
Sum of non-scaled forward probabilities at each time point. Only computed if log_space $=$ FALSE.

In case of multiple observations, these are computed independently for each sequence.

## Examples

```
# Load a pre-defined MHMM
data("mhmm_biofam")
# Compute forward and backward probabilities
fb <- forward_backward(mhmm_biofam)
# The most probable hidden state at time t
# given the observations up to time t for the first subject:
apply(fb$forward_probs[, , 1], 2, which.max)
```


## Description

Function gridplot plots multiple ssp objects to a grid.

```
Usage
    gridplot(
        x,
        nrow = NA,
        ncol = NA,
        byrow = FALSE,
        with.legend = "auto",
        legend.pos = "auto",
        legend.pos2 = "center",
        title.legend = "auto",
        ncol.legend = "auto",
        with.missing.legend = "auto",
        row.prop = "auto",
        col.prop = "auto",
        cex.legend = 1
    )
```


## Arguments

X
nrow, ncol
byrow
with.legend
legend.pos
Defines the positions of the legend boxes relative to the whole plot. Either one of "bottom" (equivalent to "auto") or "right", or a numerical vector of grid cells (by order) to print the legends to (the cells must be in one row/column).
legend. pos2 Defines the positions of the legend boxes relative to the cell(s). One of "bottomright", "bottom", "bottomleft", "left", "topleft", "top" (the default), "topright", "right" and "center".
title.legend The titles for the legend boxes. The default "auto" takes the titles from the channel labels provided by the first object in $x$. NA prints no title.
ncol.legend (A vector of) the number of columns for the legend(s). The default "auto" creates one column for each legend.

```
with.missing.legend
                    If set to "auto" (the default), a legend for the missing state is added automat-
                        ically if one or more of the sequences in data contain missing states. With the
                        value TRUE a legend for the missing state is added in any case; equivalently
                FALSE omits the legend for the missing state.
row.prop Sets the proportions of the row heights of the grid. The default value is "auto"
                for even row heights. Takes a vector of values from 0 to 1, with values summing
                to 1.
col.prop Sets the proportion of the column heights of the grid. The default value is
        "auto" for even column widths. Takes a vector of values from 0 to 1, with
        values summing to 1.
cex.legend Expansion factor for setting the size of the font for the labels in the legend. The
    default value is 1. Values lesser than 1 will reduce the size of the font, values
    greater than 1 will increase the size.
```


## See Also

ssp for defining the plot before using gridplot, and plot.ssp for plotting only one ssp object.

## Examples

```
## Not run:
data("biofam3c")
# Creating sequence objects
child_seq <- seqdef(biofam3c$children, start = 15)
marr_seq <- seqdef(biofam3c$married, start = 15)
left_seq <- seqdef(biofam3c$left, start = 15)
## Choosing colors
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
# Preparing plot for state distribution plots of observations for women
ssp_f <- ssp(
    list(child_seq[biofam3c$covariates$sex == "woman",],
            marr_seq[biofam3c$covariates$sex == "woman",],
            left_seq[biofam3c$covariates$sex == "woman",]),
    type = "d", plots = "obs", title = "Women",
    ylab = c("Children", "Married", "Left home"))
# Preparing plot for state distribution plots of observations for men
# (Updating the previous plot, only arguments that change values)
ssp_m <- update(ssp_f, title = "Men",
    x = list(child_seq[biofam3c$covariates$sex == "man",],
            marr_seq[biofam3c$covariates$sex == "man",],
            left_seq[biofam3c$covariates$sex == "man",]))
# Plotting state distribution plots of observations for women and men in two columns
```

```
gridplot(list(ssp_f, ssp_m), ncol = 2, with.legend = FALSE)
# Preparing plots for women's state distributions
ssp_f2 <- ssp(
    list(marr_seq[biofam3c$covariates$sex == "woman",],
            child_seq[biofam3c$covariates$sex == "woman",],
            left_seq[biofam3c$covariates$sex == "woman",]),
        type = "d", border = NA, with.legend = FALSE,
        title = "State distributions for women", title.n = FALSE, xtlab = 15:30,
        ylab.pos = c(1, 2, 1), ylab = c("Married", "Children", "Left home"))
# The same plot with sequences instead of state distributions
ssp_f3 <- update(
    ssp_f2, type = "I", sortv = "mds.obs", title = "Sequences for women")
# State distributions with men's data
ssp_m2 <- update(
    ssp_f2, title = "State distributions for men",
    x = list(marr_seq[biofam3c$covariates$sex == "man",],
            child_seq[biofam3c$covariates$sex == "man",],
            left_seq[biofam3c$covariates$sex == "man",]))
# Men's sequences
ssp_m3 <- update(
    ssp_m2, type = "I", sortv = "mds.obs", title = "Sequences for men")
# Plotting state distributions and index plots of observations
# for women and men in two columns (+ one column for legends)
gridplot(
    list(ssp_f2, ssp_f3, ssp_m2, ssp_m3), ncol = 3, byrow = TRUE,
    with.legend = "combined", legend.pos = "right", col.prop = c(0.35, 0.35, 0.3))
# The same with different positioning and fixed cells for legends
gridplot(
    list(ssp_f2, ssp_f3, ssp_m2, ssp_m3), ncol = 2, nrow = 3, byrow = TRUE,
    # defining the legend positions by the cell numbers
    legend.pos = 3:4)
## End(Not run)
```


## Description

Function hidden_paths computes the most probable path of hidden states of a (mixture) hidden Markov model given the observed sequences.

## Usage

hidden_paths(model, respect_void = TRUE)

## Arguments

$$
\begin{array}{ll}
\text { model } & \text { A hidden Markov model of class hmm or a mixture HMM of class mhmm. } \\
\text { respect_void } & \begin{array}{l}
\text { If TRUE (default), states at the time points corresponding to TraMineR's void in } \\
\text { the observed sequences are set to void in the hidden state sequences as well. }
\end{array}
\end{array}
$$

## Value

The most probable paths of hidden states as an stslist object (see seqdef). The log-probability is included as an attribute log_prob.

## See Also

hmm_biofam for information on the model used in the example; and seqIplot, ssplot, or mssplot for plotting hidden paths.

## Examples

```
# Load a pre-defined HMM
data("hmm_biofam")
# Compute the most probable hidden state paths given the data and the model
mpp <- hidden_paths(hmm_biofam)
# Plot hidden paths for the first 100 individuals
ssplot(mpp, type = "I", tlim = 1:100)
# Because the model structure is so sparse that the posterior probabilities are
# mostly peaked to single state at each time point, the joint probability of
# observations and most probable paths of hidden states is almost identical to
# log-likelihood:
sum(attr(mpp, "log_prob"))
logLik(hmm_biofam)
```

hmm_biofam

Hidden Markov model for the biofam data

## Description

A five-state hidden Markov model (HMM) fitted for the biofam data.

## Format

A hidden Markov model of class hmm; a left-to-right model with four hidden states.

## Details

The model is loaded by calling data(hmm_biofam). It was created with the following code:

```
data("biofam3c")
# Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
## Choosing colors
attr(marr_seq, "cpal") <- c("violetred2", "darkgoldenrod2", "darkmagenta")
attr(child_seq, "cpal") <- c("darkseagreen1", "coral3")
attr(left_seq, "cpal") <- c("lightblue", "red3")
init <- c(0.9, 0.05, 0.02, 0.02, 0.01)
# Starting values for transition matrix
trans <- matrix(
        c(0.8, 0.10, 0.05, 0.03, 0.02,
            0, 0.9, 0.05, 0.03, 0.02,
            0, 0, 0.9, 0.07, 0.03,
            0, 0, 0, 0.9, 0.1,
            0, 0, 0, 0, 1),
        nrow = 5, ncol = 5, byrow = TRUE)
# Starting values for emission matrices
emiss_marr <- matrix(
        c(0.9, 0.05, 0.05, # High probability for single
            0.9, 0.05, 0.05,
            0.05, 0.9, 0.05, # High probability for married
            0.05, 0.9, 0.05,
            0.3, 0.3, 0.4), # mixed group
        nrow = 5, ncol = 3, byrow = TRUE)
emiss_child <- matrix(
    c(0.9, 0.1, # High probability for childless
            0.9, 0.1,
            0.1, 0.9,
            0.1, 0.9,
            0.5, 0.5),
        nrow = 5, ncol = 2, byrow = TRUE)
emiss_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
```

```
        0.1, 0.9,
        0.1, 0.9,
        0.1, 0.9,
        0.5, 0.5),
    nrow = 5, ncol = 2, byrow = TRUE)
initmod <- build_hmm(
    observations = list(marr_seq, child_seq, left_seq),
    initial_probs = init, transition_probs = trans,
    emission_probs = list(emiss_marr, emiss_child,
        emiss_left),
    channel_names = c("Marriage", "Parenthood", "Residence"))
fit_biofam <- fit_model(initmod, em = FALSE, local = TRUE)
hmm_biofam <- fit_biofam$model
```


## See Also

Examples of building and fitting HMMs in build_hmm and fit_model; and biofam for the original data and biofam3c for the three-channel version used in this model.

## Examples

\# Plotting the model
plot(hmm_biofam)
hmm_mvad
Hidden Markov model for the mvad data

## Description

A hidden Markov model (MMM) fitted for the mvad data.

## Format

A hidden Markov model of class hmm; unrestricted model with six hidden states.

## Details

Model was created with the following code:
data("mvad", package = "TraMineR")
mvad_alphabet <-

```
    c("employment", "FE", "HE", "joblessness", "school", "training")
```

```
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet,
    states = mvad_scodes, labels = mvad_labels, xtstep = 6)
attr(mvad_seq, "cpal") <- colorpalette[[6]]
# Starting values for the emission matrix
emiss <- matrix(
    c(0.05, 0.05, 0.05, 0.05, 0.75, 0.05, # SC
            0.05, 0.75, 0.05, 0.05, 0.05, 0.05, # FE
            0.05, 0.05, 0.05, 0.4, 0.05, 0.4, # JL, TR
            0.05, 0.05, 0.75, 0.05, 0.05, 0.05, # HE
            0.75, 0.05, 0.05, 0.05, 0.05, 0.05),# EM
    nrow = 5, ncol = 6, byrow = TRUE)
# Starting values for the transition matrix
trans <- matrix(0.025, 5, 5)
diag(trans) <- 0.9
# Starting values for initial state probabilities
initial_probs <- c(0.2, 0.2, 0.2, 0.2, 0.2)
# Building a hidden Markov model
init_hmm_mvad <- build_hmm(observations = mvad_seq,
    transition_probs = trans, emission_probs = emiss,
    initial_probs = initial_probs)
set.seed(21)
fit_hmm_mvad <- fit_model(init_hmm_mvad, control_em = list(restart = list(times = 100)))
hmm_mvad <- fit_hmm_mvad$model
```


## See Also

Examples of building and fitting HMMs in build_hmm and fit_model; and mvad for more information on the data.

## Examples

data("hmm_mvad")
\# Plotting the model
plot(hmm_mvad)

## Description

Function logLik. hmm computes the log-likelihood value of a hidden Markov model.

## Usage

\#\# S3 method for class 'hmm'
$\operatorname{logLik}($ object, partials $=$ FALSE, threads $=1$, log_space $=$ FALSE, $\ldots$ )

## Arguments

| object | A hidden Markov model of class hmm. |
| :--- | :--- |
| partials | Return a vector containing the individual contributions of each sequence to the <br> total log-likelihood. The default is FALSE, which returns the sum of all log- <br> likelihood components. |
| threads | Number of threads to use in parallel computing. The default is 1. |
| log_space | Make computations using log-space instead of scaling for greater numerical sta- <br> bility at the cost of decreased computational performance. The default is TRUE. |
| $\ldots$ | Ignored. |

## Value

Log-likelihood of the hidden Markov model. This is an object of class logLik with attributes nobs and $d f$ inherited from the model object.

## See Also

build_hmm and fit_model for building and fitting Hidden Markov models.
logLik.mhmm Log-likelihood of the Mixture Hidden Markov Model

## Description

Function logLik.mhmm computes the log-likelihood value of a mixture hidden Markov model.

## Usage

\#\# S3 method for class 'mhmm'
$\operatorname{logLik}($ object, partials $=$ FALSE, threads $=1$, log_space $=$ FALSE,.. )

## Arguments

object A mixture hidden Markov model of class mhmm.
partials Return a vector containing the individual contributions of each sequence to the total log-likelihood. The default is FALSE, which returns the sum of all loglikelihood components.
threads Number of threads to use in parallel computing. The default is 1 .
log_space Make computations using log-space instead of scaling for greater numerical stability at the cost of decreased computational performance. The default is TRUE.
... Ignored.

## Value

Log-likelihood of the mixture hidden Markov model. This is an object of class logLik with attributes nobs and df inherited from the model object.

## See Also

build_mhmm and fit_model for building and fitting mixture Hidden Markov models.

mc_to_sc | Transform a Multichannel Hidden Markov Model into a Single Chan- |
| :--- | :--- |
| nel Representation |

## Description

Transforms data and parameters of a multichannel model into a single channel model. Observed states (symbols) are combined and parameters multiplied across channels.

## Usage

mc_to_sc(model, combine_missing = TRUE, all_combinations = FALSE)

## Arguments

model An object of class hmm or mhmm.
combine_missing
Controls whether combined states of observations at time $t$ are coded missing (coded with $*$ in stslists) if one or more of the channels include missing information at time $t$. Defaults to TRUE. FALSE keeps missing states as they are, producing more states in data; e.g. single/childless/* where the observation in channel 3 is missing.
all_combinations
Controls whether all possible combinations of observed states are included in the single channel representation or only combinations that are found in the data. Defaults to FALSE, i.e. only actual observations are included.

## Details

Note that in case of no missing observations, the log-likelihood of the original and transformed models are identical but the AIC and BIC can be different as the model attribute df is recomputed based on the single channel representation.

## See Also

build_hmm and fit_model for building and fitting Hidden Markov models; and hmm_biofam for information on the model used in the example.

## Examples

```
# Loading a hidden Markov model of the biofam data (hmm object)
data("hmm_biofam")
# Convert the multichannel model to a single-channel model
sc <- mc_to_sc(hmm_biofam)
# Likelihoods of the single-channel and the multichannel model are the same
# (Might not be true if there are missing observations)
logLik(sc)
logLik(hmm_biofam)
```

mc_to_sc_data $\quad$| Merge Multiple Sequence Objects into One (from Multichannel to Sin- |
| :--- |
| gle Channel Data) |

## Description

Function mc_to_sc_data combines observed states of multiple sequence objects into one, time point by time point.

## Usage

mc_to_sc_data(data, combine_missing = TRUE, all_combinations = FALSE)

## Arguments

data A list of state sequence objects (stslists) created with the seqdef function.
combine_missing
Controls whether combined states of observations at time $t$ are coded missing (coded with * in stslists) if one or more of the channels include missing information at time t . Defaults to TRUE. FALSE keeps missing states as they are, producing more states in data; e.g. single/childless/* where the observation in channel 3 is missing.
all_combinations
Controls whether all possible combinations of observed states are included in the single channel representation or only combinations that are found in the data. Defaults to FALSE, i.e. only actual observations are included.

## See Also

mc_to_sc for transforming multichannel hmm or mhmm objects into single-channel representations; ssplot for plotting multiple sequence data sets in the same plot; and seqdef for creating state sequence objects.

## Examples

```
# Load three-channel sequence data
data("biofam3c")
# Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
# Define colors
attr(marr_seq, "cpal") <- c("violetred2", "darkgoldenrod2", "darkmagenta")
attr(child_seq, "cpal") <- c("darkseagreen1", "coral3")
attr(left_seq, "cpal") <- c("lightblue", "red3")
# Converting multichannel data to single-channel data
sc_data <- mc_to_sc_data(list(marr_seq, child_seq, left_seq))
# 10 combined states
alphabet(sc_data)
# Colors for combined states
attr(sc_data, "cpal") <- colorpalette[[14]][1:10]
# Plotting sequences for the first 10 subjects
ssplot(list("Marriage" = marr_seq, "Parenthood" = child_seq,
    "Residence" = left_seq, "Combined" = sc_data), type = "I",
    tlim = 1:10)
# Including all combinations (whether or not available in data)
sc_data_all <- mc_to_sc_data(list(marr_seq, child_seq, left_seq),
    all_combinations = TRUE)
# 12 combined states, 2 with no observations in data
seqstatf(sc_data_all)
```


## Description

A mixture hidden Markov model (MHMM) fitted for the biofam data.

## Format

A mixture hidden Markov model of class mhmm: three clusters with left-to-right models including 4, 4, and 6 hidden states. Two covariates, sex and cohort, explaining the cluster membership.

## Details

The model was created with the following code:

```
data("biofam3c")
## Building sequence objects
marr_seq <- seqdef(biofam3c$married, start = 15,
    alphabet = c("single", "married", "divorced"))
child_seq <- seqdef(biofam3c$children, start = 15,
    alphabet = c("childless", "children"))
left_seq <- seqdef(biofam3c$left, start = 15,
    alphabet = c("with parents", "left home"))
## Choosing colors
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
## Starting values for emission probabilities
# Cluster 1
B1_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
            0.8, 0.1, 0.1,
            0.3, 0.6, 0.1, # High probability for married
            0.3, 0.3, 0.4), # High probability for divorced
    nrow = 4, ncol = 3, byrow = TRUE)
B1_child <- matrix(
    c(0.9, 0.1, # High probability for childless
            0.9, 0.1,
            0.9, 0.1,
            0.9, 0.1),
    nrow = 4, ncol = 2, byrow = TRUE)
B1_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9, # High probability for having left home
        0.1, 0.9,
        0.1, 0.9),
```

```
    nrow = 4, ncol = 2, byrow = TRUE)
# Cluster 2
B2_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
        0.8, 0.1, 0.1,
        0.1, 0.8, 0.1, # High probability for married
        0.7, 0.2, 0.1),
    nrow = 4, ncol = 3, byrow = TRUE)
B2_child <- matrix(
    c(0.9, 0.1, # High probability for childless
        0.9, 0.1,
        0.9, 0.1,
        0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
B2_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9,
        0.1, 0.9,
        0.1, 0.9),
    nrow = 4, ncol = 2, byrow = TRUE)
# Cluster 3
B3_marr <- matrix(
    c(0.8, 0.1, 0.1, # High probability for single
        0.8, 0.1, 0.1,
        0.8, 0.1, 0.1,
        0.1, 0.8, 0.1, # High probability for married
        0.3, 0.4, 0.3,
        0.1, 0.1, 0.8), # High probability for divorced
    nrow = 6, ncol = 3, byrow = TRUE)
B3_child <- matrix(
    c(0.9, 0.1, # High probability for childless
        0.9, 0.1,
        0.5, 0.5,
        0.5, 0.5,
        0.5, 0.5,
        0.1, 0.9),
    nrow = 6, ncol = 2, byrow = TRUE)
B3_left <- matrix(
    c(0.9, 0.1, # High probability for living with parents
        0.1, 0.9,
```

```
        0.5, 0.5,
        0.5, 0.5,
        0.1, 0.9,
        0.1, 0.9),
    nrow = 6, ncol = 2, byrow = TRUE)
# Starting values for transition matrices
A1 <- matrix(
    c(0.80, 0.16, 0.03, 0.01,
            0, 0.90, 0.07, 0.03,
            0, 0, 0.90, 0.10,
            0, 0, 0, 1),
    nrow = 4, ncol = 4, byrow = TRUE)
A2 <- matrix(
    c(0.80, 0.10, 0.05, 0.03, 0.01, 0.01,
        0, 0.70, 0.10, 0.10, 0.05, 0.05,
        0, 0, 0.85, 0.01, 0.10, 0.04,
        0, 0, 0, 0.90, 0.05, 0.05,
        0, 0, 0, 0, 0.90, 0.10,
        0, 0, 0, 0, 0, 1),
    nrow = 6, ncol = 6, byrow = TRUE)
# Starting values for initial state probabilities
initial_probs1 <- c(0.9, 0.07, 0.02, 0.01)
initial_probs2 <- c(0.9, 0.04, 0.03, 0.01, 0.01, 0.01)
# Birth cohort
biofam3c$covariates$cohort <- factor(cut(biofam3c$covariates$birthyr,
    c(1908, 1935, 1945, 1957)), labels = c("1909-1935", "1936-1945", "1946-1957"))
# Build mixture HMM
init_mhmm_bf <- build_mhmm(
    observations = list(marr_seq, child_seq, left_seq),
    initial_probs = list(initial_probs1, initial_probs1, initial_probs2),
    transition_probs = list(A1, A1, A2),
    emission_probs = list(list(B1_marr, B1_child, B1_left),
            list(B2_marr, B2_child, B2_left),
            list(B3_marr, B3_child, B3_left)),
    formula = ~sex + cohort, data = biofam3c$covariates,
    channel_names = c("Marriage", "Parenthood", "Residence"))
# Fitting the model
mhmm_biofam <- fit_model(init_mhmm_bf)$model
```


## See Also

Examples of building and fitting MHMMs in build_mhmm and fit_model; and biofam for the original data and biofam3c for the three-channel version used in this model.

## Examples

```
data("mhmm_biofam")
# use conditional_se = FALSE for more accurate standard errors
# (these are considerebly slower to compute)
summary(mhmm_biofam$model)
if (interactive()) {
    # Plotting the model for each cluster (change with Enter)
    plot(mhmm_biofam)
}
```

```
mhmm_mvad Mixture hidden Markov model for the mvad data
```


## Description

A mixture hidden Markov model (MHMM) fitted for the mvad data.

## Format

A mixture hidden Markov model of class mhmm: two clusters including 3 and 4 hidden states. No covariates.

## Details

The model is loaded by calling data(mhmm_mvad). It was created with the following code:

```
data("mvad", package = "TraMineR")
mvad_alphabet <-
    c("employment", "FE", "HE", "joblessness", "school", "training")
mvad_labels <- c("employment", "further education", "higher education",
    "joblessness", "school", "training")
mvad_scodes <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad_seq <- seqdef(mvad, 17:86, alphabet = mvad_alphabet,
    states = mvad_scodes, labels = mvad_labels, xtstep = 6)
attr(mvad_seq, "cpal") <- colorpalette[[6]]
# Starting values for the emission matrices
emiss_1 <- matrix(
    c(0.01, 0.01, 0.01, 0.01, 0.01, 0.95,
            0.95, 0.01, 0.01, 0.01, 0.01, 0.01,
            0.01, 0.01, 0.01, 0.95, 0.01, 0.01),
    nrow = 3, ncol = 6, byrow = TRUE)
```

```
emiss_2 <- matrix(
    c(0.01, 0.01, 0.01, 0.06, 0.90, 0.01,
        0.01, 0.95, 0.01, 0.01, 0.01, 0.01,
        0.01, 0.01, 0.95, 0.01, 0.01, 0.01,
        0.95, 0.01, 0.01, 0.01, 0.01, 0.01),
    nrow = 4, ncol = 6, byrow = TRUE)
# Starting values for the transition matrix
trans_1 <- matrix(
    c(0.95, 0.03, 0.02,
        0.01, 0.98, 0.01,
        0.01, 0.01, 0.98),
    nrow = 3, ncol = 3, byrow = TRUE)
trans_2 <- matrix(
    c(0.97, 0.01, 0.01, 0.01,
        0.01, 0.97, 0.01, 0.01,
        0.01, 0.01, 0.97, 0.01,
        0.01, 0.01, 0.01, 0.97),
    nrow = 4, ncol = 4, byrow = TRUE)
# Starting values for initial state probabilities
initial_probs_1 <- c(0.5, 0.25, 0.25)
initial_probs_2 <- c(0.4, 0.4, 0.1, 0.1)
# Building a hidden Markov model with starting values
init_mhmm_mvad <- build_mhmm(observations = mvad_seq,
    transition_probs = list(trans_1, trans_2),
    emission_probs = list(emiss_1, emiss_2),
    initial_probs = list(initial_probs_1, initial_probs_2))
# Fit the model
set.seed(123)
mhmm_mvad <- fit_model(init_mhmm_mvad, control_em = list(restart = list(times = 25)))$model
```


## See Also

Examples of building and fitting MHMMs in build_mhmm and fit_model; and mvad for more information on the data.

## Examples

```
data("mhmm_mvad")
summary (mhmm_mvad)
if (interactive()) {
```

```
    # Plotting the model for each cluster (change with Enter)
    plot(mhmm_mvad)
}
```

mssplot

Interactive Stacked Plots of Multichannel Sequences and/or Most Probable Paths for Mixture Hidden Markov Models

## Description

Function mssplot plots stacked sequence plots of observation sequences and/or most probable hidden state paths for each model of the mhmm object (model chosen according to the most probable path).

## Usage

```
mssplot(
    x,
    ask = FALSE,
    which.plots = NULL,
    hidden. paths = NULL,
    plots = "obs",
    type = "d",
    tlim \(=0\),
    sortv = NULL,
    sort. channel = 1 ,
    dist.method = "OM",
    with.missing = FALSE,
    missing.color = NULL,
    title = NA,
    title.n = TRUE,
    cex.title = 1,
    title.pos = 1,
    with.legend = "auto",
    ncol.legend = "auto",
    with.missing.legend = "auto",
    legend.prop \(=0.3\),
    cex.legend \(=1\),
    hidden.states.colors = "auto",
    hidden.states.labels = "auto",
    xaxis = TRUE,
    xlab = NA,
    xtlab = NULL,
    xlab.pos = 1 ,
    ylab = "auto",
    hidden.states.title = "Hidden states",
```

```
    yaxis = FALSE,
    ylab.pos = "auto",
    cex.lab = 1,
    cex.axis = 1,
    respect_void = TRUE,
)
```


## Arguments

$\begin{array}{ll}\text { x } & \text { Mixture hidden Markov model object of class mhmm. } \\ \text { ask } & \text { If TRUE and which.plots is NULL, plot. mhmm operates in interactive mode, } \\ \text { via menu. Defaults to FALSE. }\end{array}$ which.plots $\left.\begin{array}{l}\text { The number(s) of the requested model(s) as an integer vector. The default NULL } \\ \text { produces all plots. }\end{array}\right\}$

| title | A vector of main titles for the graphics. The default is NA: if title. $n=T R U E$, the name of the cluster and the number of subjects is plotted. FALSE prints no titles, even when title. $n=$ TRUE. |
| :---: | :---: |
| title.n | Controls whether the number of subjects is printed in the main titles of the plots. The default is TRUE: n is plotted if title is anything but FALSE. |
| cex.title | Expansion factor for setting the size of the font for the main titles. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size. |
| title.pos | Controls the position of the main titles of the plots. The default value is 1 . Values greater than 1 will place the title higher. |
| with.legend | Defines if and where the legend for the states is plotted. The default value "auto" (equivalent to TRUE and "right") creates separate legends for each requested plot and positiones them on the right-hand side of the plot. Other possible values are "bottom", "right.combined", and "bottom.combined", of which the last two create a combined legend in the selected position. FALSE prints no legend. |
| ncol.legend | (A vector of) the number of columns for the legend(s). The default "auto" creates one column for each legend. |
| with.missing.legend |  |
|  | If set to "auto" (the default), a legend for the missing state is added automatically if one or more of the sequences in the data/channel contains missing states and type = "I". If type $=" d "$ missing states are omitted from the legends unless with.missing $=$ TRUE. With the value TRUE a legend for the missing state is added in any case; equivalently FALSE omits the legend for the missing state. |
| legend.prop | Sets the proportion of the graphic area used for plotting the legend when with. legend is not FALSE. The default value is 0.3 . Takes values from 0 to 1 . |
| cex.legend | Expansion factor for setting the size of the font for the labels in the legend. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size. |
| hidden.states.colors |  |
|  | A vector of colors assigned to hidden states. The default value "auto" uses the colors assigned to the stslist object (created with seqdef) if hidden. paths is given; otherwise colors from colorpalette are automatically used. |
| hidden.states.labels |  |
|  | Labels for the hidden states. The default value "auto" uses the names provided in $x \$$ state_names if $x$ is an hmm object; otherwise the number of the hidden state. |
| xaxis | Controls whether an $x$-axis is plotted below the plot at the bottom. The default value is TRUE. |
| xlab | An optional label for the x -axis. If set to NA, no label is drawn. |
| xtlab | Optional labels for the $x$-axis tick labels. If unspecified, the column names of the seqdata sequence object are used (see seqdef). |
| xlab.pos | Controls the position of the x -axis label. The default value is 1 . Values greater than 1 will place the label further away from the plot. |


| ylab | Labels for the channels shown as labels for $y$-axes. A vector of names for each channel (observations). The default value "auto" uses the names provided in $x \$$ channel_names if $x$ is an hmm object; otherwise the names of the list in $x$ if given, or the number of the channel if names are not given. FALSE prints no labels. |
| :---: | :---: |
| hidden.states.title |  |
|  | Optional label for the hidden state plot (in the $y$-axis). The default is "Hidden states". |
| yaxis | Controls whether or not to plot the y-axis. The default is FALSE. |
| ylab.pos | Controls the position of the y axis labels (labels for channels and/or hidden states). Either "auto" or a numerical vector indicating how far away from the plots the titles are positioned. The default value "auto" positions all titles on line 1 . Shorter vectors are recycled. |
| cex.lab | Expansion factor for setting the size of the font for the axis labels. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size. |
| cex.axis | Expansion factor for setting the size of the font for the $x$-axis tick labels. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size. |
| respect_void | If TRUE (default), states at the time points corresponding to TraMineR's void in the observed sequences are set to void in the hidden state sequences as well. |
|  | Other arguments to be passed on to seqplot. |

## See Also

build_mhmm and fit_model for building and fitting mixture hidden Markov models, hidden_paths for computing the most probable paths (Viterbi paths) of hidden states, plot . mhmm for plotting mhmm objects as directed graphs, and colorpalette for default colors.

## Examples

```
# Loading mixture hidden Markov model (mhmm object)
# of the biofam data
data("mhmm_biofam")
# Plotting the first cluster only
mssplot(mhmm_biofam, which.plots = 1)
if (interactive()) {
    # Interactive plot
    mssplot(mhmm_biofam)
}
```

plot.hmm Plot hidden Markov models

## Description

Function plot.hmm plots a directed graph with pie charts of emission probabilities as vertices/nodes.

```
Usage
    ## S3 method for class 'hmm'
    plot(
        x,
        layout = "horizontal",
        pie = TRUE,
        vertex.size = 40,
        vertex.label = "initial.probs",
        vertex.label.dist = "auto",
        vertex.label.pos = "bottom",
        vertex.label.family = "sans",
        loops = FALSE,
        edge.curved = TRUE,
        edge.label = "auto",
        edge.width = "auto",
        cex.edge.width = 1,
        edge.arrow.size = 1.5,
        edge.label.family = "sans",
        label.signif = 2,
        label.scientific = FALSE,
        label.max.length = 6,
        trim = 1e-15,
        combine.slices = 0.05,
        combined.slice.color = "white",
        combined.slice.label = "others",
        with.legend = "bottom",
        ltext = NULL,
        legend.prop = 0.5,
        cex.legend = 1,
        ncol.legend = "auto",
        cpal = "auto",
        cpal.legend = "auto",
        legend.order = TRUE,
        main = NULL,
        withlegend,
    )
```


## Arguments

x
A hidden Markov model object of class hmm created with build_hmm (or build_mm). Multichannel hmm objects are automatically transformed into single-channel objects. See function mc_to_sc for more information on the transformation.
layout specifies the layout of vertices (nodes). Accepts a numerical matrix, a layout_ function (without quotation marks), or either of the predefined options "horizontal" (the default) and "vertical". Options "horizontal" and "vertical" position vertices at the same horizontal or vertical line. A two-column numerical matrix can be used to give $x$ and $y$ coordinates of the vertices. The layout_ functions available in the igraph package offer other automatic layouts for graphs.
pie Are vertices plotted as pie charts of emission probabilities? Defaults to TRUE.
vertex.size Size of vertices, given as a scalar or numerical vector. The default value is 40 .
vertex.label Labels for vertices. Possible options include "initial. probs", "names", NA, and a character or numerical vector. The default "initial.probs" prints the initial probabilities of the model and "names" prints the names of the hidden states as labels. NA prints no labels.
vertex.label.dist
Distance of the label of the vertex from its center. The default value "auto" places the label outside the vertex.
vertex.label.pos
Positions of vertex labels, relative to the center of the vertex. A scalar or numerical vector giving position(s) as radians or one of "bottom" (pi/2 as radians), "top" (-pi/2), "left" (pi), or "right" (0).
vertex.label.family, edge.label.family
Font family to be used for vertex/edge labels. See argument family in par for more information.
loops Defines whether transitions back to same states are plotted.
edge. curved Defines whether to plot curved edges (arcs, arrows) between vertices. A logical or numerical vector or scalar. Numerical values specify curvatures of edges. The default value TRUE gives curvature of 0.5 to all edges. See igraph.plotting for more information.
edge. label Labels for edges. Possible options include "auto", NA, and a character or numerical vector. The default "auto" prints transition probabilities as edge labels. NA prints no labels.
edge.width Width(s) for edges. The default "auto" determines widths according to transition probabilities between hidden states. Other possibilities are a scalar or a numerical vector of widths.
cex.edge.width An expansion factor for edge widths. Defaults to 1 .
edge.arrow.size
Size of the arrow in edges (constant). Defaults to 1.5 .
label.signif Rounds labels of model parameters to specified number of significant digits, 2 by default. Ignored for user-given labels.
label.scientific
Defines if scientific notation should be used to describe small numbers. Defaults to FALSE, e.g. 0.0001 instead of 1e-04. Ignored for user-given labels.
label.max.length
Maximum number of digits in labels of model parameters. Ignored for usergiven labels.
trim Scalar between 0 and 1 giving the highest probability of transitions that are plotted as edges, defaults to $1 \mathrm{e}-15$.
combine.slices Scalar between 0 and 1 giving the highest probability of emission probabilities that are combined into one state. The dafault value is 0.05 .
combined.slice.color
Color of the combined slice that includes the smallest emission probabilities (only if argument "combine.slices" is greater than 0 ). The default color is white.
combined.slice.label
The label for combined states (when argument "combine.slices" is greater than 0) to appear in the legend.
with.legend Defines if and where the legend of state colors is plotted. Possible values include "bottom" (the default), "top", "left", and "right". FALSE omits the legend.
ltext Optional description of (combined) observed states to appear in the legend. A vector of character strings. See seqplot for more information.
legend.prop Proportion used for plotting the legend. A scalar between 0 and 1 , defaults to 0.5.
cex.legend Expansion factor for setting the size of the font for labels in the legend. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size.
ncol.legend The number of columns for the legend. The default value "auto" sets the number of columns automatically.
cpal Optional color palette for (combinations of) observed states. The default value "auto" uses automatic color palette. Otherwise a vector of length $\times \$ n^{\prime}$ symbols is given, i.e. the argument requires a color specified for all (combinations of) observed states even if they are not plotted (if the probability is less than combine.slices).
cpal.legend Optional color palette for the legend, only considered when legend.order is FALSE. Should match ltext.
legend. order Whether to use the default order in the legend, i.e., order by appearance (first by hidden state, then by emission probability). TRUE by default.
main Main title for the plot. Omitted by default.
withlegend Deprecated. Use with. legend instead.
Other parameters passed on to plot.igraph such as vertex.color, vertex.label.cex, or edge.lty.

## See Also

build_hmm and fit_model for building and fitting Hidden Markov models, mc_to_sc for transforming multistate hmm objects into single-channel objects, hmm_biofam and hmm_mvad for information on the models used in the examples, and plot.igraph for the general plotting function of directed graphs.

## Examples

```
# Multichannel data, left-to-right model
# Loading a HMM of the biofam data
data("hmm_biofam")
# Plotting hmm object
plot(hmm_biofam)
# Plotting HMM with
plot(hmm_biofam,
    # varying curvature of edges
    edge.curved = c(0, -0.7, 0.6, 0.7, 0, -0.7, 0),
    # legend with two columns and less space
    ncol.legend = 2, legend.prop = 0.4,
    # new label for combined slice
    combined.slice.label = "States with probability < 0.05")
# Plotting HMM with given coordinates
plot(hmm_biofam,
    # layout given in 2x5 matrix
    # x coordinates in the first column
    # y coordinates in the second column
    layout = matrix(c(1, 3, 3, 5, 3,
                        0, 0, 1, 0, -1), ncol = 2),
    # larger vertices
    vertex.size = 50,
    # straight edges
    edge.curved = FALSE,
    # thinner edges and arrows
    cex.edge.width = 0.5, edge.arrow.size = 1,
    # varying positions for vertex labels (initial probabilities)
    vertex.label.pos = c(pi, pi/2, -pi/2, 0, pi/2),
    # different legend properties
    with.legend = "top", legend.prop = 0.3, cex.legend = 1.1,
    # Fix axes to the right scale
    xlim = c(0.5, 5.5), ylim = c(-1.5, 1.5), rescale = FALSE,
    # all states (not combining states with small probabilities)
    combine.slices = 0,
    # legend with two columns
    ncol.legend = 2)
# Plotting HMM with own color palette
plot(hmm_biofam, cpal = 1:10,
    # States with emission probability less than 0.2 removed
    combine.slices = 0.2,
    # legend with two columns
    ncol.legend = 2)
# Plotting HMM without pie graph and with a layout function
require("igraph")
# Setting the seed for a random layout
```

```
set.seed(1234)
plot(hmm_biofam,
    # Without pie graph
    pie = FALSE,
    # Using an automatic layout function from igraph
    layout = layout_nicely,
    vertex.size = 30,
    # Straight edges and probabilities of moving to the same state
    edge.curved = FALSE, loops = TRUE,
    # Labels with three significant digits
    label.signif = 3,
    # Fixed edge width
    edge.width = 1,
    # Remove edges with probability less than 0.01
    trim = 0.01,
    # Hidden state names as vertex labels
    vertex.label = "names",
    # Labels insidde vertices
    vertex.label.dist = 0,
    # Fix x-axis (more space on the right-hand side)
    xlim = c(-1, 1.3))
# Single-channel data, unrestricted model
# Loading a hidden Markov model of the mvad data (hmm object)
data("hmm_mvad")
# Plotting the HMM
plot(hmm_mvad)
# Checking the order of observed states (needed for the next call)
require(TraMineR)
alphabet(hmm_mvad$observations)
# Plotting the HMM with own legend (note: observation "none" nonexistent in the observations)
plot(hmm_mvad,
    # Override the default order in the legend
    legend.order = FALSE,
    # Colours in the pies (ordered by the alphabet of observations)
    cpal = c("purple", "pink", "brown", "lightblue", "orange", "green"),
    # Colours in the legend (matching to ltext)
    cpal.legend = c("orange", "pink", "brown", "green", "lightblue", "purple", "gray"),
    # Labels in the legend (matching to cpal.legend)
    ltext = c("school", "further educ", "higher educ", "training", "jobless", "employed", "none"))
require("igraph")
plot(hmm_mvad,
    # Layout in circle (layout function from igraph)
    layout = layout_in_circle,
    # Less curved edges with smaller arrows, no labels
    edge.curved = 0.2, edge.arrow.size = 0.9, edge.label = NA,
    # Positioning vertex labels (initial probabilities)
```

```
vertex.label.pos = c("right", "right", "left", "left", "right"),
# Less space for the legend
legend.prop = 0.3)
```

plot.mhmm Interactive Plotting for Mixed Hidden Markov Model (mhmm)

## Description

Function plot.mhmm plots a directed graph of the parameters of each model with pie charts of emission probabilities as vertices/nodes.

## Usage

```
## S3 method for class 'mhmm'
plot(
    x,
    interactive = TRUE,
    ask = FALSE,
    which.plots = NULL,
    nrow = NA,
    ncol = NA,
    byrow = FALSE,
    row.prop = "auto",
    col.prop = "auto",
    layout = "horizontal",
    pie = TRUE,
    vertex.size = 40,
    vertex.label = "initial.probs",
    vertex.label.dist = "auto",
    vertex.label.pos = "bottom",
    vertex.label.family = "sans",
    loops = FALSE,
    edge.curved = TRUE,
    edge.label = "auto",
    edge.width = "auto",
    cex.edge.width = 1,
    edge.arrow.size = 1.5,
    edge.label.family = "sans",
    label.signif = 2,
    label.scientific = FALSE,
    label.max.length = 6,
    trim = 1e-15,
    combine.slices = 0.05,
    combined.slice.color = "white",
    combined.slice.label = "others",
    with.legend = "bottom",
```

```
    ltext = NULL,
    legend.prop = 0.5,
    cex.legend = 1,
    ncol.legend = "auto",
    cpal = "auto",
    main = "auto",
    withlegend,
)
```


## Arguments

$x \quad$ A hidden Markov model object of class mhmm created with build_mhmm (or build_mmm or build_lcm). Multichannel mhmm objects are automatically transformed into single-channel objects. See function mc_to_sc for more information on the transformation.
interactive Whether to plot each cluster in succession or in a grid. Defaults to TRUE, i.e. clusters are plotted one after another.
ask If TRUE and which.plots is NULL, plot.mhmm operates in interactive mode, via menu. Defaults to FALSE. Ignored if interactive = FALSE.
which.plots The number(s) of the requested cluster(s) as an integer vector. The default NULL produces all plots.
nrow, ncol Optional arguments to arrange plots in a grid. Ignored if interactive $=$ TRUE
byrow Controls the order of plotting in a grid. Defaults to FALSE, i.e. plots are arranged column-wise. Ignored if interactive = TRUE.
row.prop Sets the proportions of the row heights of the grid. The default value is "auto" for even row heights. Takes a vector of values from 0 to 1 , with values summing to 1. Ignored if interactive = TRUE .
col.prop Sets the proportion of the column heights of the grid. The default value is "auto" for even column widths. Takes a vector of values from 0 to 1 , with values summing to 1 . Ignored if interactive $=$ TRUE.
layout specifies the layout of vertices (nodes). Accepts a numerical matrix, a layout_ function (without quotation marks), or either of the predefined options "horizontal" (the default) and "vertical". Options "horizontal" and "vertical" position vertices at the same horizontal or vertical line. A two-column numerical matrix can be used to give x and y coordinates of the vertices. The layout_ functions available in the igraph package offer other automatic layouts for graphs.
pie Are vertices plotted as pie charts of emission probabilities? Defaults to TRUE.
vertex.size Size of vertices, given as a scalar or numerical vector. The default value is 40 .
vertex.label Labels for vertices. Possible options include "initial.probs", "names", NA, and a character or numerical vector. The default "initial.probs" prints the initial probabilities of the model and "names" prints the names of the hidden states as labels. NA prints no labels.

vertex.label. | dist |
| :--- |
|  |
| Distance of the label of the vertex from its center. The default value "auto" |
| places the label outside the vertex. |

vertex.label. pos
Positions of vertex labels, relative to the center of the vertex. A scalar or numer-
ical vector giving position(s) as radians or one of "bottom" (pi/2 as radians),
"top" (-pi/2), "left" (pi), or "right" (0).

| ltext | Optional description of (combined) observed states to appear in the legend. A <br> vector of character strings. See seqplot for more information. |
| :--- | :--- |
| legend.prop | Proportion used for plotting the legend. A scalar between 0 and 1, defaults to <br> 0.5. |
| cex.legend | Expansion factor for setting the size of the font for labels in the legend. The <br> default value is 1. Values lesser than 1 will reduce the size of the font, values <br> greater than 1 will increase the size. |
| ncol.legend | The number of columns for the legend. The default value "auto" sets the num- <br> ber of columns automatically. |
| cpal | Optional color palette for (combinations of) observed states. The default value <br> "auto" uses automatic color palette. Otherwise a vector of length x\$n_symbols <br> is given, i.e. the argument requires a color specified for all (combinations of) ob- <br> served states even if they are not plotted (if the probability is less than combine.slices). |
| main | Optional main titles for plots. The default "auto" uses cluster_names as titles, |
| withlegend | NULL prints no titles. |
| Deprecated. Use with. legend instead. |  |

## References

Helske S. and Helske J. (2019). Mixture Hidden Markov Models for Sequence Data: The seqHMM Package in R, Journal of Statistical Software, 88(3), 1-32. doi:10.18637/jss.v088.i03

## See Also

build_mhmm and fit_model for building and fitting mixture hidden Markov models; plot.igraph for plotting directed graphs; and mhmm_biofam and mhmm_mvad for the models used in examples.

## Examples

```
# Loading mixture hidden Markov model (mhmm object)
# of the biofam data
data("mhmm_biofam")
# Plotting only the first cluster
plot(mhmm_biofam, which.plots = 1)
if (interactive()) {
    # Plotting each cluster (change with Enter)
    plot(mhmm_biofam)
    # Choosing the cluster (one at a time)
    plot(mhmm_biofam, ask = TRUE)
    # Loading MHMM of the mvad data
    data("mhmm_mvad")
    # Plotting models in the same graph (in a grid)
```

```
    # Note: the plotting window must be high enough!
    set.seed(123)
    plot(mhmm_mvad, interactive = FALSE,
        # automatic layout, legend on the right-hand side
        layout = layout_nicely, with.legend = "right",
        # Smaller and less curved edges
        edge.curved = 0.2, cex.edge.width = 0.5, edge.arrow.size = 0.7,
        vertex.label.pos = -4 * pi / 5, vertex.label.dist = 5)
```

\}
plot.ssp Stack Multichannel Sequence Plots and/or Most Probable Paths Plots from Hidden Markov Models

## Description

Function plot.ssp plots stacked sequence plots from ssp objects defined with ssp.

## Usage

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


## Arguments

x
An ssp object.
... Ignored.

## References

Helske S. and Helske J. (2019). Mixture Hidden Markov Models for Sequence Data: The seqHMM Package in R, Journal of Statistical Software, 88(3), 1-32. doi:10.18637/jss.v088.i03

## See Also

ssp for more examples and information on defining the plot before using plot. ssp; ssplot for straight plotting of ssp objects; and gridplot for plotting multiple ssp objects.

## Examples

```
data("biofam3c")
## Building sequence objects
child_seq <- seqdef(biofam3c$children, start = 15)
marr_seq <- seqdef(biofam3c$married, start = 15)
left_seq <- seqdef(biofam3c$left, start = 15)
```

```
## Choosing colors
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
# Plotting state distribution plots of observations
ssp1 <- ssp(list(child_seq, marr_seq, left_seq))
plot(ssp1)
```

    plot_colors Plot Colorpalettes
    
## Description

Function plot_colors plots colors and their labels for easy visualization of a colorpalette.

## Usage

plot_colors(x, labels = NULL)

## Arguments

x
labels

A vector of colors.
A vector of labels for colors. If omitted, given color names are used.

## See Also

See e.g. the colorpalette data and RColorBrewer package for ready-made color palettes.

## Examples

```
plot_colors(colorpalette[[5]], labels = c("one", "two", "three", "four", "five"))
plot_colors(colorpalette[[10]])
plot_colors(1:7)
plot_colors(c("yellow", "orange", "red", "purple", "blue", "green"))
plot_colors(rainbow(15))
```


## Description

Function posterior_probs computes the posterior probabilities of hidden states of a (mixture) hidden Markov model.

## Usage

posterior_probs(model, log_space = FALSE)

## Arguments

model A (mixture) hidden Markov model of class hmm or mhmm.
log_space Compute posterior probabilities in logarithmic scale. The default is FALSE.

## Value

Posterior probabilities. In case of multiple observations, these are computed independentlsy for each sequence.

## Examples

\# Load a pre-defined MHMM
data("mhmm_biofam")
\# Compute posterior probabilities
pb <- posterior_probs(mhmm_biofam)
\# Locally most probable states for the first subject:
pb[, , 1]
print.hmm
Print Method for a Hidden Markov Model

## Description

Prints the parameters of a (mixture) hidden Markov model.

## Usage

```
## S3 method for class 'hmm'
print(x, digits = 3, ...)
    ## S3 method for class 'mhmm'
    print(x, digits = 3, ...)
    ## S3 method for class 'summary.mhmm'
    print(x, digits = 3, ...)
```


## Arguments

$x \quad$ Hidden Markov model of class hmm or mhmm.
digits Minimum number of significant digits to print.
... Further arguments to print. default.

## See Also

build_hmm and fit_model for building and fitting hidden Markov models.

## Description

The separate_mhmm function reorganizes the parameters of a mhmm object into a list where each list component is an object of class hmm consisting of the parameters of the corresponding cluster.

## Usage

separate_mhmm(model)

## Arguments

model Mixture hidden Markov model of class mhmm.

## Value

List with components of class hmm.

## See Also

build_mhmm and fit_model for building and fitting MHMMs; and mhmm_biofam for more information on the model used in examples.

## Examples

\# Loading mixture hidden Markov model (mhmm object)
\# of the biofam data
data("mhmm_biofam")
\# Separate models for clusters
sep_hmm <- separate_mhmm(mhmm_biofam)
\# Plotting the model for the first cluster
plot(sep_hmm[[1]])
seqdef Imported Functions from TraMineR

## Description

Imported functions for convinience. For details, see the corresponding help pages of seqstatf, alphabet and seqdef.
seqHMM The seqHMM package

## Description

The seqHMM package is designed for fitting hidden (or latent) Markov models (HMMs) and mixture hidden Markov models (MHMMs) for social sequence data and other categorical time series. The package supports models for one or multiple subjects with one or multiple interdependent sequences (channels). External covariates can be added to explain cluster membership in mixture models. The package provides functions for evaluating and comparing models, as well as functions for easy plotting of multichannel sequences and hidden Markov models. Common restricted versions of (M)HMMs are also supported, namely Markov models, mixture Markov models, and latent class models.

## Details

Maximum likelihood estimation via the EM algorithm and direct numerical maximization with analytical gradients is supported. All main algorithms are written in C++. Parallel computation is implemented via OpenMP.

## References

Helske S. and Helske J. (2019). Mixture Hidden Markov Models for Sequence Data: The seqHMM Package in R, Journal of Statistical Software, 88(3), 1-32. doi:10.18637/jss.v088.i03
seqHMM-deprecated Deprecated function(s) in the seqHMM package

## Description

These functions are provided for compatibility with older version of the seqHMM package. They will be eventually completely removed.

## Usage

```
fit_hmm(
    model,
    em_step = TRUE,
    global_step = FALSE,
    local_step = FALSE,
    control_em = list(),
    control_global = list(),
    control_local = list(),
    lb,
    ub,
    threads = 1,
    log_space = FALSE,
)
    fit_mhmm(
    model,
    em_step = TRUE,
    global_step = FALSE,
    local_step = FALSE,
    control_em = list(),
    control_global = list(),
    control_local = list(),
    lb,
    ub,
    threads = 1,
    log_space = FALSE,
    ...
    )
    trim_hmm(
    model,
    maxit = 0,
    return_loglik = FALSE,
    zerotol = 1e-08,
    verbose = TRUE,
```

)

## Arguments

model
em_step
global_step
local_step
control_em

An object of class hmm or mhmm.
Logical. Whether or not to use the EM algorithm at the start of the parameter estimation. The default is TRUE.

Logical. Whether or not to use global optimization via nloptr (possibly after the EM step). The default is FALSE.
Logical. Whether or not to use local optimization via nloptr (possibly after the EM and/or global steps). The default is FALSE.
Optional list of control parameters for the EM algorithm. Possible arguments are
maxeval The maximum number of iterations, the default is 1000 . Note that iteration counter starts with -1 so with maxeval $=1$ you get already two iterations. This is for backward compatibility reasons.
print_level The level of printing. Possible values are 0 (prints nothing), 1 (prints information at the start and the end of the algorithm), 2 (prints at every iteration), and for mixture models 3 (print also during optimization of coefficients).
reltol Relative tolerance for convergence defined as $\left(\log L i k_{n} e w-\log L i k_{o} l d\right) /\left(a b s\left(\log L i k_{o} l d\right)+\right.$ $0.1)$. The default is $1 \mathrm{e}-10$.
restart A list containing options for possible EM restarts with the following components:
times Number of restarts of the EM algorithm using random initial values. The default is 0 , i.e. no restarts.
transition Logical. Should the original transition probabilities be varied? The default is TRUE.
emission Logical. Should the original emission probabilities be varied? The default is TRUE.
sd Standard deviation for rnorm used in randomization. The default is 0.25 .
maxeval Maximum number of iterations, the default is control_em\$maxeval
print_level Level of printing in restarted EM steps. The default is control_em\$print_level.
reltol Relative tolerance for convergence at restarted EM steps. The default is control_em\$reltol. If the relative change of the final model of the restart phase is larger than the tolerance for the original EM phase, the final model is re-estimated with the original reltol and maxeval at the end of the EM step.
n_optimum Save the log-likelihood values of the n_optimum best models (from all estimated models including the the first EM run.). The default is min(times $+1,25$ ).
use_original If TRUE. Use the initial values of the input model as starting points for the permutations. Otherwise permute the results of the first EM run.
control_global Optional list of additional arguments for nloptr argument opts. The default values are
algorithm "NLOPT_GD_MLSL_LDS"
local_opts list(algorithm = "NLOPT_LD_LBFGS", ftol_rel = 1e-6, xtol_rel $=1 \mathrm{e}-4)$
maxeval 10000 (maximum number of iterations in global optimization algorithm.)
maxtime 60 (maximum time for global optimization. Set to 0 for unlimited time.)
control_local Optional list of additional arguments for nloptr argument opts. The default values are
algorithm "NLOPT_LD_LBFGS"
ftol_rel $1 \mathrm{e}-10$
xtol_rel 1e-8
maxeval 10000 (maximum number of iterations)
lb Lower and upper bounds for parameters in Softmax parameterization. The default interval is $[\operatorname{pmin}(-25,2 *$ initialvalues $), \operatorname{pmax}(25,2 *$ initialvalues $)$ ], except for gamma coefficients, where the scale of covariates is taken into account. Note that it might still be a good idea to scale covariates around unit scale. Bounds are used only in the global optimization step.
ub Lower and upper bounds for parameters in Softmax parameterization. The default interval is $[\operatorname{pmin}(-25,2 *$ initialvalues $), \operatorname{pmax}(25,2 *$ initialvalues $)]$, except for gamma coefficients, where the scale of covariates is taken into account. Note that it might still be a good idea to scale covariates around unit scale. Bounds are used only in the global optimization step.
threads Number of threads to use in parallel computing. The default is 1.
log_space Make computations using log-space instead of scaling for greater numerical stability at a cost of decreased computational performance. The default is FALSE.
... Additional arguments to nloptr.
maxit Number of iterations. After zeroing small values, the model is refitted, and this is repeated until there is nothing to trim or maxit iterations are done.
return_loglik Return the log-likelihood of the trimmed model together with the model object. The default is FALSE.
zerotol Values smaller than this are trimmed to zero.
verbose Print results of trimming. The default is TRUE.
simulate_hmm Simulate hidden Markov models

## Description

Simulate sequences of observed and hidden states given parameters of a hidden Markov model.

## Usage

```
simulate_hmm(
        n_sequences,
        initial_probs,
        transition_probs,
        emission_probs,
        sequence_length
)
```


## Arguments

n_sequences Number of simulations.
initial_probs A vector of initial state probabilities.
transition_probs
A matrix of transition probabilities.
emission_probs A matrix of emission probabilities or a list of such objects (one for each channel).
sequence_length
Length for simulated sequences.

## Value

A list of state sequence objects of class stslist.

## See Also

build_hmm and fit_model for building and fitting hidden Markov models; ssplot for plotting multiple sequence data sets; seqdef for more information on state sequence objects; and simulate_mhmm for simulating mixture hidden Markov models.

## Examples

```
# Parameters for the HMM
emission_probs <- matrix(c(0.5, 0.2, 0.5, 0.8), 2, 2)
transition_probs <- matrix(c(5/6, 1/6, 1/6, 5/6), 2, 2)
initial_probs <- c(1, 0)
# Setting the seed for simulation
set.seed(1)
# Simulating sequences
sim <- simulate_hmm(
    n_sequences = 10, initial_probs = initial_probs,
    transition_probs = transition_probs,
    emission_probs = emission_probs,
    sequence_length = 20)
ssplot(sim, sortv = "mds.obs", type = "I")
```

```
simulate_initial_probs
```


## Description

These are helper functions for quick construction of initial values for various model building functions. Mostly useful for global optimization algorithms which do not depend on initial values.

## Usage

```
    simulate_initial_probs(n_states, n_clusters = 1)
```

    simulate_transition_probs(
        n_states,
        n_clusters = 1,
        left_right = FALSE,
        diag_c = 0
    )
    simulate_emission_probs(n_states, n_symbols, \(\mathrm{n}_{-}\)clusters \(=1\) )
    
## Arguments

n_states Number of states in each cluster.
n_clusters Number of clusters.
left_right Constrain the transition probabilities to upper triangular. Default is FALSE.
diag_c A constant value to be added to diagonal of transition matrices before scaling.
n_symbols Number of distinct symbols in each channel.

## See Also

build_hmm, build_mhmm, build_mm, build_mmm, and build_lcm for constructing different types of models.

```
simulate_mhmm
```


## Description

Simulate sequences of observed and hidden states given the parameters of a mixture hidden Markov model.

## Usage

```
simulate_mhmm(
        n_sequences,
        initial_probs,
        transition_probs,
        emission_probs,
        sequence_length,
        formula,
        data,
        coefficients
    )
```


## Arguments

n_sequences The number of simulations.
initial_probs A list containing vectors of initial state probabilities for the submodel of each cluster.
transition_probs
A list of matrices of transition probabilities for the submodel of each cluster.
emission_probs A list which contains matrices of emission probabilities or a list of such objects (one for each channel) for the submodel of each cluster. Note that the matrices must have dimensions $s x m$ where $s$ is the number of hidden states and $m$ is the number of unique symbols (observed states) in the data.

```
sequence_length
```

The length of the simulated sequences.
formula Covariates as an object of class formula, left side omitted.
data An optional data frame, a list or an environment containing the variables in the model. If not found in data, the variables are taken from environment (formula).
coefficients An optional $k x l$ matrix of regression coefficients for time-constant covariates for mixture probabilities, where $l$ is the number of clusters and $k$ is the number of covariates. A logit-link is used for mixture probabilities. The first column is set to zero.

## Value

A list of state sequence objects of class stslist.

## See Also

build_mhmm and fit_model for building and fitting mixture hidden Markov models; ssplot for plotting multiple sequence data sets; seqdef for more information on state sequence objects; and simulate_hmm for simulating hidden Markov models.

## Examples

```
emission_probs_1 <- matrix(c(0.75, 0.05, 0.25, 0.95), 2, 2)
emission_probs_2 <- matrix(c(0.1, 0.8, 0.9, 0.2), 2, 2)
```

```
colnames(emission_probs_1) <- colnames(emission_probs_2) <-
    c("heads", "tails")
transition_probs_1 <- matrix(c(9, 0.1, 1, 9.9) / 10, 2, 2)
transition_probs_2 <- matrix(c(35, 1, 1, 35) / 36, 2, 2)
rownames(emission_probs_1) <- rownames(transition_probs_1) <-
    colnames(transition_probs_1) <- c("coin 1", "coin 2")
rownames(emission_probs_2) <- rownames(transition_probs_2) <-
    colnames(transition_probs_2) <- c("coin 3", "coin 4")
initial_probs_1 <- c(1, 0)
initial_probs_2 <- c(1, 0)
n <- 30
set.seed(123)
covariate_1 <- runif(n)
covariate_2 <- sample(c("A", "B"), size = n, replace = TRUE,
    prob = c(0.3, 0.7))
dataf <- data.frame(covariate_1, covariate_2)
coefs <- cbind(cluster_1 = c(0, 0, 0), cluster_2 = c(-1.5, 3, -0.7))
rownames(coefs) <- c("(Intercept)", "covariate_1", "covariate_2B")
sim <- simulate_mhmm(
    n = n, initial_probs = list(initial_probs_1, initial_probs_2),
    transition_probs = list(transition_probs_1, transition_probs_2),
    emission_probs = list(emission_probs_1, emission_probs_2),
    sequence_length = 20, formula = ~covariate_1 + covariate_2,
    data = dataf, coefficients = coefs)
ssplot(sim$observations, hidden.paths = sim$states, plots = "both",
    sortv = "from.start", sort.channel = 0, type = "I")
hmm <- build_mhmm(sim$observations,
    initial_probs = list(initial_probs_1, initial_probs_2),
    transition_probs = list(transition_probs_1, transition_probs_2),
    emission_probs = list(emission_probs_1, emission_probs_2),
    formula = ~covariate_1 + covariate_2,
    data = dataf)
fit <- fit_model(hmm)
fit$model
paths <- hidden_paths(fit$model)
ssplot(list(estimates = paths, true = sim$states), sortv = "from.start",
    sort.channel = 2, ylab = c("estimated paths", "true (simulated)"),
    type = "I")
```

Define Arguments for Plotting Multichannel Sequences and/or Most Probable Paths from Hidden Markov Models

## Description

Function ssp defines the arguments for plotting with plot.ssp or gridplot.

## Usage

```
ssp(
    x,
    hidden.paths = NULL,
    plots = "obs",
    type = "d",
    tlim = 0,
    sortv = NULL,
    sort.channel = 1,
    dist.method = "OM",
    with.missing = FALSE,
    missing.color = NULL,
    title = NA,
    title.n = TRUE,
    cex.title = 1,
    title.pos = 1,
    with.legend = "auto",
    ncol.legend = "auto",
    with.missing.legend = "auto",
    legend.prop = 0.3,
    cex.legend = 1,
    hidden.states.colors = "auto",
    hidden.states.labels = "auto",
    xaxis = TRUE,
    xlab = NA,
    xtlab = NULL,
    xlab.pos = 1,
    ylab = "auto",
    hidden.states.title = "Hidden states",
    yaxis = FALSE,
    ylab.pos = "auto",
    cex.lab = 1,
    cex.axis = 1,
    withlegend,
    respect_void = TRUE,
)
```


## Arguments

| x | Either a hidden Markov model object of class hmm or a state sequence object <br> of class stslist (created with the seqdef) function) or a list of state sequence <br> objects. |
| :--- | :--- |
| hidden.paths | Output from hidden_paths function. Optional, if x is a hmm object or if type = <br> "obs". <br> plots <br> What to plot. One of "obs" for observations (the default), "hidden.paths" for <br> most probable paths of hidden states, or "both" for observations and hidden |
| paths together. |  |
| type | The type of the plot. Available types are "I" for sequence index plots and "d" |
| for state distribution plots (the default). See seqplot for details. |  |
| tlim | Indexes of the subjects to be plotted (the default is 0, i.e. all subjects are plotted). |
| For example, tlim = 1:10 plots the first ten subjects in data. |  |

with.legend | Defines if and where the legend for the states is plotted. The default value |
| :--- |
| "auto" (equivalent to TRUE and "right") creates separate legends for each |
| requested plot and positiones them on the right-hand side of the plot. Other |
| possible values are "bottom", "right. combined", and "bottom. combined", |
| of which the last two create a combined legend in the selected position. FALSE |
| prints no legend. |
| (A vector of) the number of columns for the legend(s). The default "auto" |
| determines number of columns depending on the position of the legend. |

ncol.legend
with.missing.legend
If set to "auto" (the default), a legend for the missing state is added automati-
cally if one or more of the sequences in the data/channel contains missing states
and type = "I". If type = "d" missing states are omitted from the legends un-
less with.missing = TRUE. With the value TRUE a legend for the missing state
is added in any case; equivalently FALSE omits the legend for the missing state.
plots the titles are positioned. The default value "auto" positions all titles on line 1. Shorter vectors are recycled.
cex.lab Expansion factor for setting the size of the font for the axis labels. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size.
cex.axis Expansion factor for setting the size of the font for the $x$-axis tick labels. The default value is 1 . Values lesser than 1 will reduce the size of the font, values greater than 1 will increase the size.
withlegend Deprecated. Use with.legend instead.
respect_void If TRUE (default), states at the time points corresponding to TraMineR's void in the observed sequences are set to void in the hidden state sequences as well.
... Other arguments to be passed on to seqplot.

## Value

Object of class ssp.

## See Also

plot.ssp for plotting objects created with the ssp function; gridplot for plotting multiple ssp objects; build_hmm and fit_model for building and fitting hidden Markov models; hidden_paths for computing the most probable paths of hidden states; and biofam 3 c and hmm_biofam for information on the data and model used in the example.

## Examples

```
data("biofam3c")
## Building sequence objects
child_seq <- seqdef(biofam3c$children, start = 15)
marr_seq <- seqdef(biofam3c$married, start = 15)
left_seq <- seqdef(biofam3c$left, start = 15)
## Choosing colors
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
# Defining the plot for state distribution plots of observations
ssp1 <- ssp(list("Parenthood" = child_seq, "Marriage" = marr_seq,
    "Residence" = left_seq))
# Plotting ssp1
plot(ssp1)
## Not run:
# Defining the plot for sequence index plots of observations
ssp2 <- ssp(
    list(child_seq, marr_seq, left_seq), type = "I", plots = "obs",
    # Sorting subjects according to the beginning of the 2nd channel (marr_seq)
```

```
    sortv = "from.start", sort.channel = 2,
    # Controlling the size, positions, and names for channel labels
    ylab.pos = c(1, 2, 1), cex.lab = 1, ylab = c("Children", "Married", "Residence"),
    # Plotting without legend
    with.legend = FALSE)
plot(ssp2)
# Plotting hidden Markov models
# Loading data
data("hmm_biofam")
# Plotting observations and most probable hidden states paths
ssp3 <- ssp(
    hmm_biofam, type = "I", plots = "both",
    # Sorting according to multidimensional scaling of hidden states paths
    sortv = "mds.hidden",
    # Controlling title
    title = "Biofam", cex.title = 1.5,
    # Labels for x axis and tick marks
    xtlab = 15:30, xlab = "Age")
plot(ssp3)
# Computing the most probable paths of hidden states
hid <- hidden_paths(hmm_biofam)
# Giving names for hidden states
library(TraMineR)
alphabet(hid) <- paste("Hidden state", 1:5)
# Plotting observations and hidden state paths
ssp4 <- ssp(
    hmm_biofam, type = "I", plots = "hidden.paths",
    # Sequence object of most probable paths
    hidden.paths = hid,
    # Sorting according to the end of hidden state paths
    sortv = "from.end", sort.channel = 0,
    # Contolling legend position, type, and proportion
    with.legend = "bottom.combined", legend.prop = 0.15,
    # Plotting without title and y label
    title = FALSE, ylab = FALSE)
plot(ssp4)
## End(Not run)
```


## Description

Function ssplot plots stacked sequence plots of sequence object created with the seqdef function or observations and/or most probable paths of hmm objects.

## Usage

```
ssplot(
    x,
    hidden.paths = NULL,
    plots = "obs",
    type = "d",
    tlim = 0,
    sortv = NULL,
    sort.channel = 1,
    dist.method = "OM",
    with.missing = FALSE,
    missing.color = NULL,
    title = NA,
    title.n = TRUE,
    cex.title = 1,
    title.pos = 1,
    with.legend = "auto",
    ncol.legend = "auto",
    with.missing.legend = "auto",
    legend.prop = 0.3,
    cex.legend = 1,
    hidden.states.colors = "auto",
    hidden.states.labels = "auto",
    xaxis = TRUE,
    xlab = NA,
    xtlab = NULL,
    xlab.pos = 1,
    ylab = "auto",
    hidden.states.title = "Hidden states",
    yaxis = FALSE,
    ylab.pos = "auto",
    cex.lab = 1,
    cex.axis = 1,
    respect_void = TRUE,
)
```


## Arguments

x
Either a hidden Markov model object of class hmm or a state sequence object of class stslist (created with the seqdef) function) or a list of state sequence objects.

| hidden.paths | Output from hidden_paths function. Optional, if x is a hmm object or if type = <br> "obs". <br> plots <br>  <br>  <br>  <br>  <br>  <br>  <br> What to plot. One of "obs" for observations (the default), "hidden. paths" for <br> maths together. |
| :--- | :--- |
| type | The type of the plot. Available types are "I" for sequence index plots and "d" |
| for state distribution plots (the default). See seqplot for details. |  |

ncol.legend | (A vector of) the number of columns for the legend(s). The default "auto" |
| :--- |
| determines number of columns depending on the position of the legend. |
| with.missing. legend |
| If set to "auto" (the default), a legend for the missing state is added automati- |
| cally if one or more of the sequences in the data/channel contains missing states |
| and type = "I". If type = "d" missing states are omitted from the legends un- |
| less with.missing = TRUE. With the value TRUE a legend for the missing state |
| is added in any case; equivalently FALSE omits the legend for the missing state. |
| Sets the proportion of the graphic area used for plotting the legend when with.legend |
| is not FALSE. The default value is 0.3. Takes values from 0 to 1. |

Expansion factor for setting the size of the font for the labels in the legend. The
default value is 1. Values lesser than 1 will reduce the size of the font, values
greater than 1 will increase the size.

| cex.axis | Expansion factor for setting the size of the font for the x-axis tick labels. The <br> default value is 1 . Values lesser than 1 will reduce the size of the font, values <br> greater than 1 will increase the size. |
| :--- | :--- |
| respect_void | If TRUE (default), states at the time points corresponding to TraMineR's void in <br> the observed sequences are set to void in the hidden state sequences as well. |
| $\ldots$. | Other arguments to be passed on to seqplot. |

## See Also

ssp for creating ssp objects and plot. ssp and gridplot for plotting these; build_hmm and fit_model for building and fitting hidden Markov models; hidden_paths for computing the most probable paths of hidden states; and biofam3c hmm_biofam for information on the data and model used in the example.

## Examples

```
data("biofam3c")
# Creating sequence objects
child_seq <- seqdef(biofam3c$children, start = 15)
marr_seq <- seqdef(biofam3c$married, start = 15)
left_seq <- seqdef(biofam3c$left, start = 15)
## Choosing colors
attr(child_seq, "cpal") <- c("#66C2A5", "#FC8D62")
attr(marr_seq, "cpal") <- c("#AB82FF", "#E6AB02", "#E7298A")
attr(left_seq, "cpal") <- c("#A6CEE3", "#E31A1C")
# Plotting state distribution plots of observations
ssplot(list("Children" = child_seq, "Marriage" = marr_seq,
"Residence" = left_seq))
## Not run:
# Plotting sequence index plots of observations
ssplot(
    list(child_seq, marr_seq, left_seq), type = "I",
    # Sorting subjects according to the beginning of the 2nd channel (marr_seq)
    sortv = "from.start", sort.channel = 2,
    # Controlling the size, positions, and names for channel labels
    ylab.pos = c(1, 2, 1), cex.lab = 1, ylab = c("Children", "Married", "Residence"),
    # Plotting without legend
    with.legend = FALSE)
# Plotting hidden Markov models
# Loading a ready-made HMM for the biofam data
data("hmm_biofam")
# Plotting observations and hidden states paths
ssplot(
```

```
    hmm_biofam, type = "I", plots = "both",
    # Sorting according to multidimensional scaling of hidden states paths
    sortv = "mds.hidden",
    ylab = c("Children", "Married", "Left home"),
    # Controlling title
    title = "Biofam", cex.title = 1.5,
    # Labels for x axis and tick marks
    xtlab = 15:30, xlab = "Age")
    # Computing the most probable paths of hidden states
    hidden.paths <- hidden_paths(hmm_biofam)
    hidden.paths_seq <- seqdef(hidden.paths, labels = paste("Hidden state", 1:5))
    # Plotting observations and hidden state paths
    ssplot(
    hmm_biofam, type = "I", plots = "hidden.paths",
    # Sequence object of most probable paths
    hidden.paths = hidden.paths_seq,
    # Sorting according to the end of hidden state paths
    sortv = "from.end", sort.channel = 0,
    # Contolling legend position, type, and proportion
    with.legend = "bottom", legend.prop = 0.15,
    # Plotting without title and y label
    title = FALSE, ylab = FALSE)
    ## End(Not run)
```

summary.mhmm Summary method for mixture hidden Markov models

## Description

Function summary.mhmm gives a summary of a mixture hidden Markov model.

## Usage

```
    ## S3 method for class 'mhmm'
    summary(
        object,
    parameters = FALSE,
    conditional_se = TRUE,
    log_space = FALSE,
    )
```


## Arguments

object Mixture hidden Markov model of class mhmm.

```
parameters Whether or not to return transition, emission, and initial probabilities. FALSE by
    default.
conditional_se Return conditional standard errors of coefficients. See vcov.mhmm for details.
                    TRUE by default.
log_space Make computations using log-space instead of scaling for greater numerical sta-
    bility at cost of decreased computational performance. Default is FALSE.
... Further arguments to vcov.mhmm.
```


## Details

The summary.mhmm function computes features from a mixture hidden Markov model and stores them as a list. A print method prints summaries of these: log-likelihood and BIC, coefficients and standard errors of covariates, means of prior cluster probabilities, and information on most probable clusters.

## Value

transition_probs Transition probabilities. Only returned if parameters = TRUE.
emission_probs Emission probabilities. Only returned if parameters = TRUE.
initial_probs Initial state probabilities. Only returned if parameters = TRUE.
logLik Log-likelihood.
BIC Bayesian information criterion.
most_probable_cluster The most probable cluster according to posterior probabilities.
coefficients Coefficients of covariates.
vcov Variance-covariance matrix of coefficients.
prior_cluster_probabilities Prior cluster probabilities (mixing proportions) given the covariates.
posterior_cluster_probabilities Posterior cluster membership probabilities.
classification_table Cluster probabilities (columns) by the most probable cluster (rows).

## See Also

build_mhmm and fit_model for building and fitting mixture hidden Markov models; and mhmm_biofam for information on the model used in examples.

## Examples

```
# Loading mixture hidden Markov model (mhmm object)
# of the biofam data
data("mhmm_biofam")
# Model summary
summary(mhmm_biofam)
```


## Description

Function trim_model tries to set small insignificant probabilities to zero without decreasing the likelihood.

## Usage

```
trim_model(
    model,
    maxit = 0,
    return_loglik = FALSE,
    zerotol = 1e-08,
    verbose = TRUE,
    )
```


## Arguments

| model | Model of class hmm or mhmm for which trimming is performed. |
| :--- | :--- |
| maxit | Number of iterations. After zeroing small values, the model is refitted, and this <br> is repeated until there is nothing to trim or maxit iterations are done. |
| return_loglik | Return the log-likelihood of the trimmed model together with the model object. <br> The default is FALSE. |
| zerotol | Values smaller than this are trimmed to zero. |
| verbose | Print results of trimming. The default is TRUE. |
| $\ldots$ | Further parameters passed on to fit_model. |

## See Also

build_hmm and fit_model for building and fitting hidden Markov models; and hmm_biofam for information on the model used in the example.

## Examples

```
data("hmm_biofam")
# Testing if changing parameter values smaller than 1e-03 to zero
# leads to improved log-likelihood.
hmm_trim <- trim_model(hmm_biofam, zerotol = 1e-03, maxit = 10)
```

| vcov.mhmm | Variance-Covariance Matrix for Coefficients of Covariates of Mixture |
| :--- | :--- |
|  | Hidden Markov Model |

## Description

Returns the asymptotic covariances matrix of maximum likelihood estimates of the coefficients corresponding to the explanatory variables of the model.

## Usage

\#\# S3 method for class 'mhmm'
vcov(object, conditional $=$ TRUE, threads $=1$, log_space $=$ FALSE, ...)

## Arguments

| object <br> conditional | Object of class mhmm. <br> If TRUE (default), the standard errors are computed conditional on other model <br> parameters. See details. |
| :--- | :--- |
| threads | Number of threads to use in parallel computing. Default is 1. |
| log_space | Make computations using log-space instead of scaling for greater numerical sta- <br> bility at cost of decreased computational performance. Default is FALSE. |
| $\ldots$ | Additional arguments to function jacobian of numDeriv package. |

## Details

The conditional standard errors are computed using analytical formulas by assuming that the coefficient estimates are not correlated with other model parameter estimates (or that the other parameters are assumed to be fixed). This often underestimates the true standard errors, but is substantially faster approach for preliminary analysis. The non-conditional standard errors are based on the numerical approximation of the full Hessian of the coefficients and the model parameters corresponding to nonzero probabilities. Computing the non-conditional standard errors can be slow for large models as the Jacobian of analytical gradients is computed using finite difference approximation.

## Value

Matrix containing the variance-covariance matrix of coefficients.

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