

# Package ‘L1mstate’

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**Type** Package

**Title** L1-Regularized Multi-State Models

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**Description** Fitting the regularization path of the L1-regularized multi-state models since they can exploit sparsity structure of input. Different tuning regularization parameter methods are provided. The cumulative hazard rate estimation and the transition probability predictions can be made from the fitted models.

**License** GPL (>= 2)

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L1mstate-package	<i>L1-Regularized Multi-State Models</i>
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## Description

Fitting the regularization path of the L1-regularized multi-state models since they can exploit sparsity structure of input. Different tuning regularization parameter methods are provided. The cumulative hazard rate estimation and the transition probability predictions can be made from the fitted models.

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plot.probs.l1mstate	Plot the transition probabilities
probs.l1mstate	Compute subject-specific or overall transition probabilities

## Maintainer

Xuan Dang <xuandang11289@gmail.com>

## Author(s)

Xuan Dang

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coefl1mstate	<i>Obtain the coefficients</i>
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### Description

Obtain the optimal coefficients using the first cross-validation method or the penalized cross-validation method by setting `s = "lambda.min"` or `"lambda.pcv1"`. Can also obtain the coefficient at any lambda value within the range of sequence.

### Usage

```
coefl1mstate(object, s=c("lambda.pcv1", "lambda.min"))
```

### Arguments

<code>object</code>	fitted <code>cv.l1mstateR</code> object
<code>s</code>	lambda value (numeric type) or cross-validation methods (character type)

### Details

Return the coefficient values

### Value

Return the coefficient values.

### Author(s)

Xuan Dang <xuandang11289@gmail.com>

### Examples

```
set.seed(1001)
p <- 9
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)

beta12 <- c(-.65,-.65,-.65,0,0,-.65,-.65,0,0)
beta13 <- c(-.65,-.65,0,0,0,0,-.65,0,0)
beta23 <- c(0,-.65,-.65,0,0,-.65,-.65,0,-.65)

N <- 200
x <- mvrnorm(n=N, mu, sigma)
col_names <- c(sprintf("%d", seq(1,dim(x)[2])))
```

```

colnames(x) <- col_names
N12 <- N-2*N%/5
N13 <- N-N12
hx12 <- exp(x[1:N12,] %>% beta12)
ty12 <- rexp(N12,hx12)
hx13 <- exp(x[(N12+1):N,] %>% beta13)
ty13 <- rexp(N13,hx13)
hx23 <- exp(x[1:N12,] %>% beta23)
ty23 <- rexp(N12,hx23)
t2 <- rep(0,N)
t2[1:N12] <- ty12
t2[(N12+1):N] <- ty13
s2 <- rep(0,N)
s2[1:N12] <- rep(1,N12)
t3 <- rep(0,N)
t3[1:N12] <- ty12+ty23
t3[(N12+1):N] <- ty13
#censor case for state 3
tcens3 <- rbinom(n=N, prob = 0.3, size = 1)
s3 <- 1-tcens3

# pre-process data into long-format using msprep
dt <- data.frame(illt=t2, ills=s2, dt=t3, ds=s3, x)
tmat <- matrix(c(NA,NA,NA,1,NA,NA,2,3,NA), nrow = 3)
longdt <- msprep(time=c(NA,"illt","dt"),status=c(NA,"ills","ds"),
                 keep = col_names, data=dt,trans=tmat)

cv.l1fit <- cv.l1mstateR(longdt, nlambda = 100, nolds = 10)
coefl1mstate(cv.l1fit, s="lambda.min")

```

---

cumhaz.l1mstate

*Compute subject-specific transition hazard rates*


---

## Description

This function computes subject-specific cumulative transition hazard rates for each of the possible transitions in the multi-state model.

## Usage

```

cumhaz.l1mstate(object, longdt, newdata, cv.method = c("pcv1", "min"))

```

## Arguments

object	fitted cv.l1mstateR object.
longdt	long-format data input
newdata	a new data with the same long-format with the same covariate names as longdt.
cv.method	the cross-validation method used to select the optimal result.

**Value**

time	A list of all time points for each of the transitions in the multi-state model.
baseHaz	A list of the estimated subject-specific baseline hazards for each of the transitions in the multi-state model.
var	A list of the variance of the estimated subject-specific hazards for each of the transitions in the multi-state model.
cumbaseHaz	A list of the estimated subject-specific cumulative hazards for each of the transitions in the multi-state model.
Haz	A list of the estimated subject-specific hazards for each of the transitions in the multi-state model.

**Author(s)**

Xuan Dang <xuandang11289@gmail.com>

**Examples**

```
#### EBMT data
data(EBMT)

x1 <- ifelse(EBMT$match=="gender mismatch", 1, 0)
x2 <- ifelse(EBMT$proph=="yes", 1, 0)
x3 <- ifelse(EBMT$year=="1990-1994", 1, 0)
x4 <- ifelse(EBMT$year=="1995-1998", 1, 0)
x5 <- ifelse(EBMT$agecl=="20-40", 1, 0)
x6 <- ifelse(EBMT$agecl==">40", 1, 0)
dt <- data.frame(EBMT[,1:11],x1,x2,x3,x4,x5,x6)
col_names <- c("x1", "x2", "x3", "x4", "x5", "x6")
tmat <- transMat(x = list(c(2,3,5,6),c(4,5,6),c(4,5,6),c(5,6),c()),c()),
  names = c("Tx", "Rec", "AE", "Rec+AE", "Rel", "Death"))
dat <- msprep(data = dt, trans = tmat, time = c(NA, "rec", "ae",
  "recae", "rel", "srv"), status = c(NA, "rec.s", "ae.s",
  "recae.s", "rel.s", "srv.s"), keep = col_names)
# select the first 450 lines for a sake of illustration
dat <- dat[1:450,]
out <- cv.l1mstateR(dat, nlambda = 100, nolds = 10)

#### Compute the cumulative hazard and the transition probabilities
## patient A: include the highest number of patients (287)
ptsA <- dat[which(dat$x1==0 & dat$x2==0 & dat$x3==1 & dat$x4==0 & dat$x5==1 & dat$x6==0),]
## observed transitions (ground truth)
# predicted time = 0
events(ptsA)
# use models to predict the transition probabilities
# L1MSTATE
ptA <- ptsA[which(ptsA$id == unique(ptsA$id)[1]),]
ptA <- ptA[which(ptA$trans %in% c(1:4)),c(4,9:14)]

cumhazA <- cumhaz.l1mstate(object=out, longdt=dat, newdata=ptA, cv.method="pcv1")
```

---

 cv.l1mstateR

*Cross-validation for l1mstateR*


---

**Description**

Does k-fold cross-validation for l1mstateR

**Usage**

```
cv.l1mstateR(longdt, lambda=NULL, nlambda=100, rlambda=NULL,
             nfolds=1, foldid=NULL, thresh=1e-7, maxit=1e+5)
```

**Arguments**

longdt	input in long-format structure
lambda	A user supplied lambda sequence. Typical usage is to have the program compute its own lambda sequence based on nlambda and rlambda. Supplying a value of lambda overrides this. <b>WARNING:</b> use with care. Avoid supplying a single value for lambda. Supply instead a decreasing sequence of lambda values. l1mstateR relies on its warm starts for speed, and its often faster to fit a whole path than compute a single fit.
nlambda	The number of lambda values- default is 100.
rlambda	Smallest value for lambda, as a fraction of the maximum lambda, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size relative to the number of covariates. If sample size > #covariates, the default is 0.001, close to zero. If sample size < #covariates, the default is 0.01.
nfolds	Number of folds - default is 10. Smallest value allowable is nfolds=3.
foldid	an optional vector of values between 1 and nfolds identifying what fold each observation is in.
thresh	Convergence threshold for one-step coordinate descent. Defaults value is 1E-7.
maxit	Maximum number of passes over the data for all lambda values; default is 1E+5.

**Value**

aBetaSTD	A list of coefficients in standardized form, each one corresponds to each lambda value.
aBeta0	A list of coefficients in original form, each one corresponds to each lambda value.
pBetaSTD	The coefficient in standardized form gives maximum log-likelihood value using the penalized cross-validation method.
pBeta0	The coefficient in original form gives maximum log-likelihood value using the penalized cross-validation method.

mBetaSTD	The coefficient in standardized form gives maximum log-likelihood value using the first cross-validation method.
mBeta0	The coefficient in original form gives maximum log-likelihood value using the first cross-validation method.
fit	A matrix includes lambda value, the mean cross-validation error.
fit	A matrix of lambda values and log-likelihood values
numcovs	Number of covariates
numtrans	Number of transitions

### Author(s)

Xuan Dang <xuandang11289@gmail.com>

### Examples

```

set.seed(1001)
p <- 9
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)

beta12 <- c(-.65,-.65,-.65,0,0,-.65,-.65,0,0)
beta13 <- c(-.65,-.65,0,0,0,0,-.65,0,0)
beta23 <- c(0,-.65,-.65,0,0,-.65,-.65,0,-.65)

N <- 200
x <- mvrnorm(n=N, mu, sigma)
col_names <- c(sprintf("X%d", seq(1,dim(x)[2])))
colnames(x) <- col_names
N12 <- N-2*N%/5
N13 <- N-N12
hx12 <- exp(x[1:N12,] %*% beta12)
ty12 <- rexp(N12,hx12)
hx13 <- exp(x[(N12+1):N,] %*% beta13)
ty13 <- rexp(N13,hx13)
hx23 <- exp(x[1:N12,] %*% beta23)
ty23 <- rexp(N12,hx23)
t2 <- rep(0,N)
t2[1:N12] <- ty12
t2[(N12+1):N] <- ty13
s2 <- rep(0,N)
s2[1:N12] <- rep(1,N12)
t3 <- rep(0,N)
t3[1:N12] <- ty12+ty23
t3[(N12+1):N] <- ty13
# censor case for state 3

```

```

tcens3 <- rbinom(n=N, prob = 0.3, size = 1)
s3 <- 1-tcens3

# pre-process data into long-format using msprep
dt <- data.frame(illt=t2, illt=s2, dt=t3, ds=s3, x)
tmat <- matrix(c(NA,NA,NA,1,NA,NA,2,3,NA), nrow = 3)
longdt <- msprep(time=c(NA,"illt","dt"),status=c(NA,"illt","ds"),
                 keep = col_names, data=dt,trans=tmat)

cv.l1fit <- cv.l1mstateR(longdt, nlambda = 100, nfolds = 10)

```

---

l1mstateR

*Fit multi-state models with lasso regularization*


---

## Description

Fit a multi-state models via penalized partial likelihood. The regularization path is computed for the lasso at a path of values for the regularization parameter lambda. Can deal with right-censoring and left-truncated data.

## Usage

```
l1mstateR(longdt, lambda=NULL, nlambda=100, rlambda=NULL, thresh=1e-7, maxit=1e+5)
```

## Arguments

longdt	input in long-format structure
lambda	A user supplied lambda sequence. Typical usage is to have the program compute its own lambda sequence based on nlambda and rlambda. Supplying a value of lambda overrides this. <b>WARNING:</b> use with care. Avoid supplying a single value for lambda. Supply instead a decreasing sequence of lambda values. l1mstateR relies on its warm starts for speed, and it's often faster to fit a whole path than compute a single fit.
nlambda	The number of lambda values- default is 100.
rlambda	Smallest value for lambda, as a fraction of the maximum lambda, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size relative to the number of covariates. If sample size > #covariates, the default is 0.001, close to zero. If sample size < #covariates, the default is 0.01.
thresh	Convergence threshold for one-step coordinate descent. Default value is 1E-7.
maxit	Maximum number of passes over the data for all lambda values; default is 1E+5.

## Details

Please note that input has to be long-format structure.

**Value**

aBetaSTD	A list of coefficients in standardized form, each one corresponds to each lambda value.
aBeta0	A list of coefficients in original form, each one corresponds to each lambda value.
fit	A matrix of lambda values and log-likelihood values
numcovs	Number of covariates
numtrans	Number of transitions

**Author(s)**

Xuan Dang <xuandang11289@gmail.com>

**Examples**

```

set.seed(1001)
p <- 9
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)

beta12 <- c(-.65,-.65,-.65,0,0,-.65,-.65,0,0)
beta13 <- c(-.65,-.65,0,0,0,0,-.65,0,0)
beta23 <- c(0,-.65,-.65,0,0,-.65,-.65,0,-.65)

N <- 200
x <- mvrnorm(n=N, mu, sigma)
col_names <- c(sprintf("X%d", seq(1,dim(x)[2])))
colnames(x) <- col_names
N12 <- N-2*N/%5
N13 <- N-N12
hx12 <- exp(x[1:N12,] %*% beta12)
ty12 <- rexp(N12,hx12)
hx13 <- exp(x[(N12+1):N,] %*% beta13)
ty13 <- rexp(N13,hx13)
hx23 <- exp(x[1:N12,] %*% beta23)
ty23 <- rexp(N12,hx23)
t2 <- rep(0,N)
t2[1:N12] <- ty12
t2[(N12+1):N] <- ty13
s2 <- rep(0,N)
s2[1:N12] <- rep(1,N12)
t3 <- rep(0,N)
t3[1:N12] <- ty12+ty23
t3[(N12+1):N] <- ty13
# censor case for state 3

```

```

tcens3 <- rbinom(n=N, prob = 0.3, size = 1)
s3 <- 1-tcens3

# pre-process data into long-format using msprep
dt <- data.frame(illt=t2, ills=s2, dt=t3, ds=s3, x)
tmat <- matrix(c(NA,NA,NA,1,NA,NA,2,3,NA), nrow = 3)
longdt <- msprep(time=c(NA,"illt","dt"),status=c(NA,"ills","ds"),
                 keep = col_names, data=dt,trans=tmat)

l1fit <- l1mstateR(longdt, nlambda = 100)

```

---

plot.cumhaz.l1mstate *Plot the estimated cumulative hazard rates of the multi-state model.*

---

### Description

Plot the estimated cumulative hazard rates of the multi-state model.

### Usage

```

## S3 method for class 'cumhaz.l1mstate'
plot(x,type=c("single","separate"),cols,
     xlab="Years since transplant",ylab="Cumulative hazard",
     ylim,lwd=3,lty,legend,legend.pos,bty="o",...)

```

### Arguments

x	fitted cumhaz.l1mstate object
type	One of "single" or "separate"; in case of "single", all estimated cumulative hazards are drawn in a single plot, in case of "separate", separate plots are shown for the estimated cumulative hazards.
cols	A vector specifying colors for the different transitions
xlab	A title for the x-axis; default is "Years since transplant"
ylab	A title for the y-axis; default is "Cumulative hazard"
ylim	The y limits of the plot
lwd	The line width; default is 3
lty	The line type
legend	The transition numbers; if missing, these will be taken from the transition matrix contained in cumhaz.l1mstate object.
legend.pos	The position of the legend; default is "topleft"
bty	The box type of the legend
...	Further arguments to plot

### Details

A plot is produced, and nothing is returned.

**Value**

No return value

**Author(s)**

Xuan Dang <xuandang11289@gmail.com>

**Examples**

```
#### EBMT data
data(ebmt4)

x1 <- ifelse(ebmt4$match=="gender mismatch", 1, 0)
x2 <- ifelse(ebmt4$proph=="yes", 1, 0)
x3 <- ifelse(ebmt4$year=="1990-1994", 1, 0)
x4 <- ifelse(ebmt4$year=="1995-1998", 1, 0)
x5 <- ifelse(ebmt4$agecl=="20-40", 1, 0)
x6 <- ifelse(ebmt4$agecl==">40", 1, 0)
dt <- data.frame(ebmt4[,1:11],x1,x2,x3,x4,x5,x6)
col_names <- c("x1", "x2", "x3", "x4", "x5", "x6")
tmat <- transMat(x = list(c(2,3,5,6),c(4,5,6),c(4,5,6),c(5,6),c(),c()),
  names = c("Tx", "Rec", "AE", "Rec+AE", "Rel", "Death"))
dat <- msprep(data = dt, trans = tmat, time = c(NA, "rec", "ae",
  "recae","rel", "srv"),status = c(NA, "rec.s", "ae.s",
  "recae.s","rel.s", "srv.s"), keep = col_names)
# select the first 450 lines for a sake of illustration
dat <- dat[1:450,]
out <- cv.l1mstateR(dat, nlambda = 100, nfolds = 10)

### Compute the cumulative hazard and the transition probabilities
## patient A: include the highest number of patients (287)
ptsA <- dat[which(dat$x1==0 & dat$x2==0 & dat$x3==1 & dat$x4==0 & dat$x5==1 & dat$x6==0),]
## observed transitions (ground truth)
# predicted time = 0
events(ptsA)
# use models to predict the transition probabilities
# L1MSTATE
ptA <- ptsA[which(ptsA$id == unique(ptsA$id)[1]),]
ptA <- ptA[which(ptA$trans %in% c(1:4)),c(4,9:14)]

cumhazA <- cumhaz.l1mstate(object = out, longdt = dat, newdata = ptA, cv.method = "pcv1")
plot.cumhaz.l1mstate(cumhazA, main = "L1MSTATE", type = "single", lwd=2, las=1,
ylim = c(0,1))
legend("topleft", inset=.01, legend=c("1", "2", "3", "4"), col=c("black", "red", "green",
"blue"), lty=1, cex=0.8, title="Transitions", text.font=2, bg='white')
```

---

plot.l1mstateCoef

*Plots the coefficient paths of transitions produced by l1mstateR or cv.l1mstateR*

---

**Description**

Plots the coefficient values as a function of the lambda values used.

**Usage**

```
## S3 method for class 'l1mstateCoef'
plot(x, trans = NULL,...)
```

**Arguments**

x	fitted l1mstateR or cv.l1mstateR object.
trans	choose the transition you want to plot the coefficient. It can be a single transition or a set of several transitions.
...	further arguments to plot

**Details**

A plot is produced, and nothing is returned.

**Value**

No return value.

**Author(s)**

Xuan Dang <xuandang11289@gmail.com>

**Examples**

```
set.seed(1001)
p <- 9
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)

beta12 <- c(-.65,-.65,-.65,0,0,-.65,-.65,0,0)
beta13 <- c(-.65,-.65,0,0,0,0,-.65,0,0)
beta23 <- c(0,-.65,-.65,0,0,-.65,-.65,0,-.65)

N <- 200
x <- mvrnorm(n=N, mu, sigma)
col_names <- c(sprintf("X%d", seq(1,dim(x)[2])))
colnames(x) <- col_names
N12 <- N-2*N%/5
N13 <- N-N12
hx12 <- exp(x[1:N12,] %*% beta12)
```

```

ty12 <- rexp(N12,hx12)
hx13 <- exp(x[(N12+1):N,] %*% beta13)
ty13 <- rexp(N13,hx13)
hx23 <- exp(x[1:N12,] %*% beta23)
ty23 <- rexp(N12,hx23)
t2 <- rep(0,N)
t2[1:N12] <- ty12
t2[(N12+1):N] <- ty13
s2 <- rep(0,N)
s2[1:N12] <- rep(1,N12)
t3 <- rep(0,N)
t3[1:N12] <- ty12+ty23
t3[(N12+1):N] <- ty13
#censor case for state 3
tcens3 <- rbinom(n=N, prob = 0.3, size = 1)
s3 <- 1-tcens3

# pre-process data into long-format using msprep
dt <- data.frame(illt=t2, ills=s2, dt=t3, ds=s3, x)
tmat <- matrix(c(NA,NA,NA,1,NA,NA,2,3,NA), nrow = 3)
longdt <- msprep(time=c(NA,"illt","dt"),status=c(NA,"ills","ds"),
                 keep = col_names, data=dt,trans=tmat)

l1fit <- l1mstateR(longdt, nlambda = 100)
plot.l1mstateCoef(l1fit, trans=1)

```

---

plot.l1mstateCV

*plot the cross-validation curve produced by cv.l1mstateR*


---

### Description

Plots the cross-validation curve, and upper and lower standard deviation curves, as a function of the lambda values used.

### Usage

```
## S3 method for class 'l1mstateCV'
plot(x, ...)
```

### Arguments

x	fitted cv.l1mstateR object
...	Further arguments to plot

### Details

A plot is produced, and nothing is returned.

**Value**

No return value.

**Author(s)**

Xuan Dang <xuandang11289@gmail.com>

**Examples**

```

set.seed(1001)
p <- 9
times <- 1:p
rho <- 0.5
H <- abs(outer(times, times, "-"))
C <- 1 * rho^H
C[cbind(1:p, 1:p)] <- C[cbind(1:p, 1:p)]
sigma <- matrix(C,p,p)
mu <- rep(0,p)

beta12 <- c(-.65,-.65,-.65,0,0,-.65,-.65,0,0)
beta13 <- c(-.65,-.65,0,0,0,0,-.65,0,0)
beta23 <- c(0,-.65,-.65,0,0,-.65,-.65,0,-.65)

N <- 200
x <- mvrnorm(n=N, mu, sigma)
col_names <- c(sprintf("%d", seq(1,dim(x)[2])))
colnames(x) <- col_names
N12 <- N-2*N%/5
N13 <- N-N12
hx12 <- exp(x[1:N12,] %*% beta12)
ty12 <- rexp(N12,hx12)
hx13 <- exp(x[(N12+1):N,] %*% beta13)
ty13 <- rexp(N13,hx13)
hx23 <- exp(x[1:N12,] %*% beta23)
ty23 <- rexp(N12,hx23)
t2 <- rep(0,N)
t2[1:N12] <- ty12
t2[(N12+1):N] <- ty13
s2 <- rep(0,N)
s2[1:N12] <- rep(1,N12)
t3 <- rep(0,N)
t3[1:N12] <- ty12+ty23
t3[(N12+1):N] <- ty13
#censor case for state 3
tcens3 <- rbinom(n=N, prob = 0.3, size = 1)
s3 <- 1-tcens3

# pre-process data into long-format using msprep
dt <- data.frame(illt=t2, ills=s2, dt=t3, ds=s3, x)
tmat <- matrix(c(NA,NA,NA,1,NA,NA,2,3,NA), nrow = 3)
longdt <- msprep(time=c(NA,"illt","dt"),status=c(NA,"ills","ds"),
  keep = col_names, data=dt,trans=tmat)

```

```
cv.l1fit <- cv.l1mstateR(longdt, nlambda = 100, nfolds = 10)
plot.l1mstateCV(cv.l1fit)
```

---

plot.probs.l1mstate     *Plot the transition probabilities*

---

## Description

Plot the transition probabilities produced by probs.l1mstate

## Usage

```
## S3 method for class 'probs.l1mstate'
plot(x, from, type=c("stacked", "filled", "single", "separate"),
     ord, cols, xlab="Years since transplant", ylab="Probability",
     xlim, ylim, lwd, lty, cex, legend, legend.pos, bty="o", ...)
```

## Arguments

x	object produced from probs.l1mstate
from	the starting state from which the probabilities are used to plot
type	with "stacked" type, the transition probabilities are stacked and the distance between two adjacent curves indicates the probability; with "filled" type, it is the same but the space between adjacent curves are filled; with "single" type, the probabilities are shown as different curves in a single plot; with "separate" type, separate plots are shown for the estimated transition probabilities
ord	A vector indicates the order of plotting in case "stacked" or "filled"
cols	A vector specifying colors for the different transitions
xlab	A title for the x-axis; default is "Years since transplant"
ylab	A title for the y-axis; default is "Probability"
xlim	The x limits of the plot; default is range of time
ylim	The y limits of the plot
lwd	The line width; default is 3
lty	The line type
cex	Character size
legend	The transition numbers; if missing, these will be taken from the transition matrix contained in cumhaz.l1mstate object.
legend.pos	The position of the legend; default is "topleft"
bty	The box type of the legend
...	Further arguments to plot

**Details**

A plot is produced, and nothing is returned.

**Value**

No return value.

**Author(s)**

Xuan Dang <xuandang11289@gmail.com>

**Examples**

```
#### EBMT data
data(ebmt4)

x1 <- ifelse(ebmt4$match=="gender mismatch", 1, 0)
x2 <- ifelse(ebmt4$proph=="yes", 1, 0)
x3 <- ifelse(ebmt4$year=="1990-1994", 1, 0)
x4 <- ifelse(ebmt4$year=="1995-1998", 1, 0)
x5 <- ifelse(ebmt4$agec1=="20-40", 1, 0)
x6 <- ifelse(ebmt4$agec1==">40", 1, 0)
dt <- data.frame(ebmt4[,1:11],x1,x2,x3,x4,x5,x6)
col_names <- c("x1", "x2", "x3", "x4", "x5", "x6")
tmat <- transMat(x = list(c(2,3,5,6),c(4,5,6),c(4,5,6),c(5,6),c(),c()),
                 names = c("Tx", "Rec", "AE", "Rec+AE", "Rel", "Death"))
dat <- msprep(data = dt, trans = tmat, time = c(NA, "rec", "ae",
        "recae", "rel", "srv"), status = c(NA, "rec.s", "ae.s",
        "recae.s", "rel.s", "srv.s"), keep = col_names)
# select the first 450 lines for a sake of illustration
dat <- dat[1:450,]
out <- cv.l1mstateR(dat, nlambda = 100, nfolds = 10)

### Compute the cumulative hazard and the transition probabilities
## patient A: include the highest number of patients (287)
ptsA <- dat[which(dat$x1==0 & dat$x2==0 & dat$x3==1 & dat$x4==0 & dat$x5==1 & dat$x6==0),]
## observed transitions (ground truth)
# predicted time = 0
events(ptsA)
# use models to predict the transition probabilities
# L1MSTATE
ptA <- ptsA[which(ptsA$id == unique(ptsA$id)[1]),]
ptA <- ptA[which(ptA$trans %in% c(1:4)),c(4,9:14)]

cumhazA <- cumhaz.l1mstate(object = out, longdt = dat, newdata = ptA, cv.method = "pcv1")
probA_0 <- probs.l1mstate(cumhazA, longdt = longdt, tmat, predt = 0, direction = "forward")
statecols <- heat_hcl(6, c = c(90, 10), l = c(20, 80), power = c(1/5, 2))[c(6, 5, 3, 4, 2, 1)]
ord <- c(1, 2, 3, 4, 5, 6)
plot.probs.l1mstate(probA_0, main = "L1MSTATE", from = 1, ord = ord,
                   las=1, cex=0, type = "filled", col = statecols[ord])
```

---

probs.l1mstate                      *Compute subject-specific or overall transition probabilities*

---

### Description

Compute subject-specific or overall transition probabilities.

### Usage

```
probs.l1mstate(object, longdt, tmat, predt, direction=c("forward", "fixedhorizon"))
```

### Arguments

object	fitted cumhaz.l1mstate object
longdt	long-format data input
tmat	a transition matrix for multi-state model
predt	a prediction time
direction	indicates whether prediction is forward or for a fixed horizon

### Value

A list with each element `[[s]]` containing a data frame with the estimated transition probabilities from state `s`. It also includes transition and `tmat` information for plotting purpose

### Author(s)

Xuan Dang <xuandang11289@gmail.com>

### Examples

```
#### EBMT data
data(ebmt4)

x1 <- ifelse(ebmt4$match=="gender mismatch", 1, 0)
x2 <- ifelse(ebmt4$proph=="yes", 1, 0)
x3 <- ifelse(ebmt4$year=="1990-1994", 1, 0)
x4 <- ifelse(ebmt4$year=="1995-1998", 1, 0)
x5 <- ifelse(ebmt4$agec1=="20-40", 1, 0)
x6 <- ifelse(ebmt4$agec1==">40", 1, 0)
dt <- data.frame(ebmt4[,1:11],x1,x2,x3,x4,x5,x6)
col_names <- c("x1", "x2", "x3", "x4", "x5", "x6")
tmat <- transMat(x = list(c(2,3,5,6),c(4,5,6),c(4,5,6),c(5,6),c(),c()),
                 names = c("Tx", "Rec", "AE", "Rec+AE", "Rel", "Death"))
dat <- msprep(data = dt, trans = tmat, time = c(NA, "rec", "ae",
        "recae","rel", "srv"),status = c(NA, "rec.s", "ae.s",
        "recae.s","rel.s", "srv.s"), keep = col_names)
# select the first 450 lines for a sake of illustration
dat <- dat[1:450,]
```

```
out <- cv.l1mstateR(dat, nlambda = 100, nfolds = 10)

### Compute the cumulative hazard and the transition probabilities
## patient A: include the highest number of patients (287)
ptsA <- dat[which(dat$x1==0 & dat$x2==0 & dat$x3==1 & dat$x4==0 & dat$x5==1 & dat$x6==0),]
## observed transitions (ground truth)
# predicted time = 0
events(ptsA)
# use models to predict the transition probabilities
# L1MSTATE
ptA <- ptsA[which(ptsA$id == unique(ptsA$id)[1]),]
ptA <- ptA[which(ptA$trans %in% c(1:4)),c(4,9:14)]

cumhazA <- cumhaz.l1mstate(object = out, longdt = dat, newdata = ptA, cv.method = "pcv1")
proba_0 <- probs.l1mstate(cumhazA, longdt = longdt, tmat, predt = 0, direction = "forward")
```

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