

Package ‘momentuHMM’

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Type Package

Title Maximum Likelihood Analysis of Animal Movement Behavior Using Multivariate Hidden Markov Models

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Description Extended tools for analyzing telemetry data using generalized hidden Markov models. Features of momentuHMM (pronounced ``momentum'') include data pre-processing and visualization, fitting HMMs to location and auxiliary biotelemetry or environmental data, biased and correlated random walk movement models, hierarchical HMMs, multiple imputation for incorporating location measurement error and missing data, user-specified design matrices and constraints for covariate modelling of parameters, random effects, decoding of the state process, visualization of fitted models, model checking and selection, and simulation. See McClintock and Michelot (2018) <[doi:10.1111/2041-210X.12995](https://doi.org/10.1111/2041-210X.12995)>.

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<i>AIC.momentuHMM</i>	<i>AIC</i>
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Description

Akaike information criterion of momentuHMM model(s).

Usage

```
## S3 method for class 'momentuHMM'
AIC(object, ..., k = 2, n = NULL)
```

Arguments

object	A momentuHMM object.
...	Optional additional momentuHMM objects, to compare AICs of the different models. These can be passed as a list using the !!! operator (see rlang and example in AICweights).
k	Penalty per parameter. Default: 2 ; for classical AIC.
n	Optional sample size. If specified, the small sample correction AIC is used (i.e., $AICc = AIC + kp(p+1)/(n-p-1)$ where p is the number of parameters).

Value

The AIC of the model(s) provided. If several models are provided, the AICs are output in ascending order.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m
AIC(m)

## Not run:
# HMM specifications
nbStates <- 2
```

```

stepDist <- "gamma"
angleDist <- "vm"
mu0 <- c(20,70)
sigma0 <- c(10,30)
kappa0 <- c(1,1)
stepPar0 <- c(mu0,sigma0)
anglePar0 <- c(-pi/2,pi/2,kappa0)
formula <- ~cov1+cov2

# example$m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
mod1 <- fitHMM(example$m$data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
               Par0=list(step=stepPar0,angle=anglePar0),
               formula=~1,estAngleMean=list(angle=TRUE))

Par0 <- getPar0(mod1,formula=formula)
mod2 <- fitHMM(example$m$data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
               Par0=Par0$Par,beta0=Par0$beta,
               formula=formula,estAngleMean=list(angle=TRUE))

AIC(mod1,mod2)

Par0nA <- getPar0(mod1,estAngleMean=list(angle=FALSE))
mod3 <- fitHMM(example$m$data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
               Par0=Par0nA$Par,beta0=Par0nA$beta,
               formula=~1)

AIC(mod1,mod2,mod3)

# add'1 models provided as a list using the !!! operator
AIC(mod1, !!!list(mod2,mod3))

## End(Not run)

```

AICweights*Calculate Akaike information criterion model weights***Description**

Calculate Akaike information criterion model weights

Usage

```
AICweights(..., k = 2, n = NULL)
```

Arguments

...

[momentuHMM](#), [HMMfits](#), or [miHMM](#) objects, to compare AIC weights of the different models. The first object must be a [momentuHMM](#), [HMMfits](#), or [miHMM](#) object, but additional model objects can be passed as a list using the !!! operator (see [rlang](#)).

- k Penalty per parameter. Default: 2 ; for classical AIC.
- n Optional sample size. If specified, the small sample correction AIC is used (i.e., $AICc = AIC + kp(p+1)/(n-p-1)$ where p is the number of parameters).

Details

- Model objects must all be either of class `momentuHMM` or multiple imputation model objects (of class `HMMfits` and/or `miHMM`).
- AIC is only valid for comparing models fitted to the same data. The data for each model fit must therefore be identical. For multiple imputation model objects, respective model fits must have identical data.

Value

The AIC weights of the models. If multiple imputation objects are provided, then the mean model weights (and standard deviations) are provided.

Examples

```
## Not run:
# HMM specifications
nbStates <- 2
stepDist <- "gamma"
angleDist <- "vm"
mu0 <- c(20,70)
sigma0 <- c(10,30)
kappa0 <- c(1,1)
stepPar0 <- c(mu0,sigma0)
anglePar0 <- c(-pi/2,pi/2,kappa0)
formula <- ~cov1+cov2

# example$m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
mod1 <- fitHMM(example$m$data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
               Par0=list(step=stepPar0,angle=anglePar0),
               formula=~1,estAngleMean=list(angle=TRUE))

Par0 <- getPar0(mod1,formula=formula)
mod2 <- fitHMM(example$m$data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
               Par0=Par0$Par,beta0=Par0$beta,
               formula=formula,estAngleMean=list(angle=TRUE))

AICweights(mod1,mod2)

Par0nA <- getPar0(mod1,estAngleMean=list(angle=FALSE))
mod3 <- fitHMM(example$m$data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
               Par0=Par0nA$Par,beta0=Par0nA$beta,
               formula=~1)

AICweights(mod1,mod2,mod3)
```

```
# add'l models provided as a list using the !!! operator  
AICweights(mod1, !!!list(mod2,mod3))  
  
## End(Not run)
```

allProbs

Matrix of all probabilities

Description

Used in functions [viterbi](#), [logAlpha](#), [logBeta](#).

Usage

```
allProbs(m)
```

Arguments

m Object [momentuHMM](#) or [miSum](#).

Value

Matrix of all probabilities.

Examples

```
## Not run:  
P <- momentuHMM:::allProbs(m=example$m)  
  
## End(Not run)
```

checkPar0

Check parameter length and order for a [fitHMM](#) (or [MIfitHMM](#)) model

Description

Prints parameters with labels based on DM, formula, and/or formulaDelta. See [fitHMM](#) for further argument details.

Usage

```

checkPar0(data, ...)

## Default S3 method:
checkPar0(
  data,
  nbStates,
  dist,
  Par0 = NULL,
  beta0 = NULL,
  delta0 = NULL,
  estAngleMean = NULL,
  circularAngleMean = NULL,
  formula = ~1,
  formulaDelta = NULL,
  stationary = FALSE,
  mixtures = 1,
  formulaPi = NULL,
  DM = NULL,
  userBounds = NULL,
  workBounds = NULL,
  betaCons = NULL,
  betaRef = NULL,
  deltaCons = NULL,
  stateNames = NULL,
  fixPar = NULL,
  prior = NULL,
  ...
)

## S3 method for class 'hierarchical'
checkPar0(
  data,
  hierStates,
  hierDist,
  Par0 = NULL,
  hierBeta = NULL,
  hierDelta = NULL,
  estAngleMean = NULL,
  circularAngleMean = NULL,
  hierFormula = NULL,
  hierFormulaDelta = NULL,
  mixtures = 1,
  formulaPi = NULL,
  DM = NULL,
  userBounds = NULL,
  workBounds = NULL,
  betaCons = NULL,

```

```

    deltaCons = NULL,
    fixPar = NULL,
    prior = NULL,
    ...
)

```

Arguments

data	<code>momentuHMMDData</code> object, <code>momentuHierHMMDData</code> object, or a data frame containing the data stream and covariate values
...	further arguments passed to or from other methods
nbStates	Number of states of the HMM.
dist	A named list indicating the probability distributions of the data streams.
Par0	Optional named list containing vectors of state-dependent probability distribution parameters for each data stream specified in <code>dist</code> . If <code>Par0</code> is not provided, then ordered parameter indices are returned.
beta0	Optional matrix of regression coefficients for the transition probabilities. If <code>beta0</code> is not provided, then ordered parameter indices are returned.
delta0	Optional values or regression coefficients for the initial distribution of the HMM. If <code>delta0</code> is not provided, then ordered parameter indices are returned.
estAngleMean	An optional named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpcalch').
circularAngleMean	An optional named list indicating whether to use circular-linear or circular-circular regression on the mean of circular distributions ('vm' and 'wrpcalch') for turning angles.
formula	Regression formula for the transition probability covariates.
formulaDelta	Regression formula for the initial distribution.
stationary	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. Default: FALSE.
mixtures	Number of mixtures for the state transition probabilities.
formulaPi	Regression formula for the mixture distribution probabilities. Note that only the covariate values from the first row for each individual ID in <code>data</code> are used (i.e. time-varying covariates cannot be used for the mixture probabilities).
DM	An optional named list indicating the design matrices to be used for the probability distribution parameters of each data stream.
userBounds	An optional named list of 2-column matrices specifying bounds on the natural (i.e. real) scale of the probability distribution parameters for each data stream.
workBounds	An optional named list of 2-column matrices specifying bounds on the working scale of the probability distribution, transition probability, and initial distribution parameters.
betaCons	Matrix of the same dimension as <code>beta0</code> composed of integers identifying any equality constraints among the t.p.m. parameters.

<code>betaRef</code>	Numeric vector of length <code>nbStates</code> indicating the reference elements for the t.p.m. multinomial logit link.
<code>deltaCons</code>	Matrix of the same dimension as <code>delta0</code> composed of integers identifying any equality constraints among the initial distribution working scale parameters. Ignored unless a formula is provided in <code>formulaDelta</code> .
<code>stateNames</code>	Optional character vector of length <code>nbStates</code> indicating state names.
<code>fixPar</code>	An optional list of vectors indicating parameters which are assumed known prior to fitting the model.
<code>prior</code>	A function that returns the log-density of the working scale parameter prior distribution(s).
<code>hierStates</code>	A hierarchical model structure <code>Node</code> for the states ('state'). See fitHMM .
<code>hierDist</code>	A hierarchical data structure <code>Node</code> for the data streams ('dist'). See fitHMM .
<code>hierBeta</code>	A hierarchical data structure <code>Node</code> for the initial matrix of regression coefficients for the transition probabilities at each level of the hierarchy ('beta'). See fitHMM .
<code>hierDelta</code>	A hierarchical data structure <code>Node</code> for the initial values for the initial distribution at each level of the hierarchy ('delta'). See fitHMM .
<code>hierFormula</code>	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy ('formula'). See fitHMM .
<code>hierFormulaDelta</code>	A hierarchical formula structure for the initial distribution covariates for each level of the hierarchy ('formulaDelta'). See fitHMM . Default: NULL (no covariate effects and <code>fixPar\$delta</code> is specified on the working scale).

See Also

[fitHMM](#), [MIfitHMM](#)

Examples

```
m <- example$m
checkPar0(data=m$data, nbStates=2, dist=m$conditions$dist,
          estAngleMean = m$conditions$estAngleMean,
          formula = m$conditions$formula)

par <- getPar(m)
checkPar0(data=m$data, nbStates=2, dist=m$conditions$dist,
          estAngleMean = m$conditions$estAngleMean,
          formula = m$conditions$formula,
          Par0=par$Par, beta0=par$beta, delta0=par$delta)

dummyDat <- data.frame(step=0,angle=0,cov1=0,cov2=0)
checkPar0(data=dummyDat, nbStates=2, dist=m$conditions$dist,
          estAngleMean = m$conditions$estAngleMean,
          formula = m$conditions$formula)

## Not run:
simDat <- simData(nbStates=2, dist=m$conditions$dist, Par = par$Par,
                  spatialCovs = list(forest=forest),
```

```

        centers = matrix(0,1,2),
        nbCovs = 2)
checkPar0(data = simDat, nbStates=2, dist=m$conditions$dist,
          formula = ~forest,
          DM = list(step=list(mean=~cov1, sd=~cov2),
                     angle=list(mean=~center1.angle,concentration=~1)),
          estAngleMean=list(angle=TRUE),
          circularAngleMean=list(angle=TRUE))

par <- list(step=rnorm(8),angle=rnorm(4))
beta0 <- matrix(rnorm(4),2,2)
delta0 <- c(0.5,0.5)
checkPar0(data = simDat, nbStates=2, dist=m$conditions$dist,
          Par0 = par, beta0 = beta0, delta0 = delta0,
          formula = ~forest,
          DM = list(step=list(mean=~cov1, sd=~cov2),
                     angle=list(mean=~center1.angle,concentration=~1)),
          estAngleMean=list(angle=TRUE),
          circularAngleMean=list(angle=TRUE))

## End(Not run)

```

CIbeta*Confidence intervals for working (i.e., beta) parameters***Description**

Computes the standard errors and confidence intervals on the beta (i.e., working) scale of the data stream probability distribution parameters, as well as for the transition probabilities regression parameters. Working scale depends on the real (i.e., natural) scale of the parameters. For non-circular distributions or for circular distributions with `estAngleMean=FALSE`:

Usage

```
CIbeta(m, alpha = 0.95)
```

Arguments

<code>m</code>	A <code>momentuHMM</code> object
<code>alpha</code>	Significance level of the confidence intervals. Default: 0.95 (i.e. 95% CIs).

Details

- 1) if both lower and upper bounds are finite then logit is the working scale; 2) if lower bound is finite and upper bound is infinite then log is the working scale.

For circular distributions with `estAngleMean=TRUE` and no constraints imposed by a design matrix (DM) or bounds (userBounds), then the working parameters are complex functions of both the angle mean and concentrations/sd natural parameters (in this case, it's probably best just to focus on the

real parameter estimates!). However, if constraints are imposed by DM or userBounds on circular distribution parameters with `estAngleMean=TRUE` and `circularAngleMean=FALSE`:

- 1) if the natural bounds are $(-\pi, \pi]$ then tangent is the working scale, otherwise if both lower and upper bounds are finite then logit is the working scale; 2) if lower bound is finite and upper bound is infinite then log is the working scale.

When circular-circular regression is specified using `circularAngleMean`, the working scale for the mean turning angle is not as easily interpretable, but the link function is $\text{atan2}(\sin(X)*B, 1+\cos(X)*B)$, where X are the angle covariates and B the angle coefficients. Under this formulation, the reference turning angle is 0 (i.e., movement in the same direction as the previous time step). In other words, the mean turning angle is zero when the coefficient(s) B=0.

Value

A list of the following objects:

...	List(s) of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the working parameters of the data streams
beta	List of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the working parameters of the transition probabilities

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

CIBeta(m)
```

circAngles

Convert standard direction angles (in radians relative to the x-axis) to turning angle covariates suitable for circular-circular regression on the angle mean

Description

This function can be used to convert angular covariates (e.g., ocean currents, wind direction) measured in radians relative to the x-axis to turning angle covariates suitable for circular-circular regression in `fitHMM` or `MIfitHMM`.

Usage

```
circAngles(refAngle, data, coordNames = c("x", "y"))
```

Arguments

refAngle	Numeric vector of standard direction angles (in radians) relative to the x-axis, where 0 = east, pi/2 = north, pi = west, -pi/2 = south
data	data frame containing fields for the x- and y-coordinates (identified by coordNames) and 'ID' (if more than one individual)
coordNames	Names of the columns of coordinates in data. Default: c("x", "y").

Value

A vector of turning angles between the movement direction at time step t-1 and refAngle at time t

Examples

```
# extract data from momentuHMM example
data<-example$m$data

# generate fake angle covariates
u <- rnorm(nrow(data)) # horizontal component
v <- rnorm(nrow(data)) # vertical component
refAngle <- atan2(v,u)

# add turning angle covariate to data
data$cov3 <- circAngles(refAngle=refAngle,data=data)
```

Description

Computes the standard errors and confidence intervals on the real (i.e., natural) scale of the data stream probability distribution parameters, as well as for the transition probabilities parameters. If covariates are included in the probability distributions or TPM formula, the mean values of non-factor covariates are used for calculating the natural parameters. For any covariate(s) of class 'factor', then the value(s) from the first observation in the data are used.

Usage

```
CIreal(m, alpha = 0.95, covs = NULL, parms = NULL)

## Default S3 method:
CIreal(m, alpha = 0.95, covs = NULL, parms = NULL)

## S3 method for class 'hierarchical'
CIreal(m, alpha = 0.95, covs = NULL, parms = NULL)
```

Arguments

<code>m</code>	A <code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miHMM</code> , or <code>miSum</code> object
<code>alpha</code>	Significance level of the confidence intervals. Default: 0.95 (i.e. 95% CIs).
<code>covs</code>	Data frame consisting of a single row indicating the covariate values to be used in the calculations. By default, no covariates are specified.
<code>parms</code>	Optional character vector indicating which groups of real parameters to calculate confidence intervals for (e.g., 'step', 'angle', 'gamma', 'delta', etc.). Default: <code>NULL</code> , in which case confidence intervals are calculated for all groups of parameters in the model.

Details

For any covariates that are not specified using `covs`, the means of the covariate(s) are used (unless the covariate is a factor, in which case the first factor in the data is used).

Value

A list of the following objects:

<code>...</code>	List(s) of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the natural parameters of the data streams
<code>gamma</code>	List of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the transition probabilities
<code>delta</code>	List of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the initial state probabilities
<code>hierGamma</code>	A hierarchical data structure <code>Node</code> including a list of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the transition probabilities for each level of the hierarchy (only applies if <code>m</code> is a hierarchical model object)
<code>hierDelta</code>	A hierarchical data structure <code>Node</code> including a list of estimates ('est'), standard errors ('se'), and confidence intervals ('lower', 'upper') for the initial state probabilities for each level of the hierarchy (only applies if <code>m</code> is a hierarchical model object)

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

ci1<-CIreal(m)

# specify 'covs'
ci2<-CIreal(m,covs=data.frame(cov1=mean(m$data$cov1),cov2=mean(m$data$cov2)))

all.equal(ci1,ci2)
```

crawlMerge *Merge crwData or crwHierData object with additional data streams and/or covariates*

Description

This function can be used to merge `crwData` or `crwHierData` objects (as returned by `crawlWrap`) with additional data streams and/or covariates that are unrelated to location.

Usage

```
crawlMerge(crwData, data, Time.name)
```

Arguments

<code>crwData</code>	A <code>crwData</code> or <code>crwHierData</code> object
<code>data</code>	A data frame containing required columns <code>ID</code> , <code>Time.name</code> , and, if <code>crwData</code> is hierarchical, <code>level</code> , plus any additional data streams and/or covariates to merge with <code>crwData</code> .
<code>Time.name</code>	Character string indicating name of the time column to be used for merging

Details

Specifically, the function merges the `crwData$crwPredict` data frame with data based on the ID, Time.name, and, if `crwData` is hierarchical, level columns. Thus data must contain ID, Time.name, and, if `crwData` is hierarchical, level columns.

Only rows of data with ID, Time.name, and, if crwData is hierarchical, level values that exactly match crwData\$crwPredict are merged. Typically, the Time.name column in data should match predicted times of locations in crwData\$crwPredict (i.e. those corresponding to crwData\$crwPredict\$locType=="p")

Value

A `crwData` object

Examples

```
# create data frame with fake data stream
data <- data.frame(ID=rep(factor(c(1,2)),times=c(753,652)),
                     time=c(1:753,1:652),
                     fake=rpois(753+652,5))

# merge fake data stream with crwOut
crwOut <- crawlMerge(crwOut,data,"time")

## End(Not run)
```

crawlWrap*Fit and predict tracks for using crawl***Description**

Wrapper function for fitting `crawl::crwMLE` models and predicting locations with `crawl::crwPredict` for multiple individuals.

Usage

```
crawlWrap(
  obsData,
  timeStep = 1,
  ncores = 1,
  retryFits = 0,
  retrySD = 1,
  retryParallel = FALSE,
  mov.model = ~1,
  err.model = NULL,
  activity = NULL,
  drift = NULL,
  coord = c("x", "y"),
  proj = NULL,
  Time.name = "time",
  time.scale = "hours",
  theta,
  fixPar,
  method = "L-BFGS-B",
  control = NULL,
  constr = NULL,
  prior = NULL,
  need.hess = TRUE,
  initialSANN = list(maxit = 200),
  attempts = 1,
  predTime = NULL,
  fillCols = FALSE,
```

```

coordLevel = NULL,
...
)

```

Arguments

<code>obsData</code>	data.frame object containing fields for animal ID ('ID'), time of observation (identified by <code>Time.name</code> , must be numeric or POSIXct), and observed locations (x- and y- coordinates identified by <code>coord</code>), such as that returned by <code>simData</code> when temporally-irregular observed locations or measurement error are included. Alternatively, a <code>SpatialPointsDataFrame</code> or sf object will also be accepted, in which case the coord values will be taken from the spatial data set and ignored in the arguments. Note that <code>crwMLE</code> requires that longitude/latitude coordinates be projected to UTM (i.e., easting/northing). For further details see <code>crwMLE</code> .
<code>timeStep</code>	Length of the time step at which to predict regular locations from the fitted model. Unless <code>predTime</code> is specified, the sequence of times is <code>seq(a_i, b_i, timeStep)</code> where <code>a_i</code> and <code>b_i</code> are the times of the first and last observations for individual <code>i</code> . <code>timeStep</code> can be numeric (regardless of whether <code>obsData[[Time.name]]</code> is numeric or POSIXct) or a character string (if <code>obsData[[Time.name]]</code> is of class POSIXct) containing one of "sec", "min", "hour", "day", "DSTday", "week", "month", "quarter" or "year". This can optionally be preceded by a positive integer and a space, or followed by "s" (e.g., "2 hours"; see <code>seq.POSIXt</code>). <code>timeStep</code> is not used for individuals for which <code>predTime</code> is specified.
<code>ncores</code>	Number of cores to use for parallel processing. Default: 1 (no parallel processing).
<code>retryFits</code>	Number of times to attempt to achieve convergence and valid (i.e., not NaN) variance estimates after the initial model fit.
<code>retrySD</code>	An optional list of scalars or vectors for each individual indicating the standard deviation to use for normal perturbations of theta when <code>retryFits>0</code> (or <code>attempts>1</code>). Instead of a list object, <code>retrySD</code> can also be a scalar or a vector, in which case the same values are used for each each individual. If a scalar is provided, then the same value is used for each parameter. If a vector is provided, it must be of length <code>length(theta)</code> for the corresponding individual(s). Default: 1, i.e., a standard deviation of 1 is used for all parameters of all individuals. Ignored unless <code>retryFits>0</code> (or <code>attempts>1</code>).
<code>retryParallel</code>	Logical indicating whether or not to perform <code>retryFits</code> attempts for each individual in parallel. Default: FALSE. Ignored unless <code>retryFits>0</code> and <code>ncores>1</code> . Note that when attempts are done in parallel (i.e. <code>retryParallel=TRUE</code>), the current value for the log-likelihood of each individual and warnings about convergence are not printed to the console.
<code>mov.model</code>	List of mov.model objects (see <code>crwMLE</code>) containing an element for each individual. If only one movement model is provided, then the same movement model is used for each individual.
<code>err.model</code>	List of err.model objects (see <code>crwMLE</code>) containing an element for each individual. If only one error model is provided, then the same error model is used for

	each individual (in which case the names of the <code>err.model</code> components corresponding to easting/longitudinal and northing/latitudinal location error must match <code>coord</code>).
<code>activity</code>	List of activity objects (see crwMLE) containing an element for each individual. If only one activity covariate is provided, then the same activity covariate is used for each individual.
<code>drift</code>	List of drift objects (see crwMLE) containing an element for each individual. If only one drift component is provided, then the same drift component is used for each individual.
<code>coord</code>	A 2-vector of character values giving the names of the "x" and "y" coordinates in <code>data</code> . See crwMLE .
<code>proj</code>	A list of valid epsg integer codes or proj4string for <code>obsData</code> that does not inherit either 'sf' or 'sp'. A valid 'crs' list is also accepted. Otherwise, ignored. If only one <code>proj</code> is provided, then the same projection is used for each individual.
<code>Time.name</code>	Character indicating name of the location time column. See crwMLE .
<code>time.scale</code>	character. Scale for conversion of POSIX time to numeric for modeling. Defaults to "hours".
<code>theta</code>	List of theta objects (see crwMLE) containing an element for each individual. If only one theta is provided, then the same starting values are used for each individual. If theta is not specified, then crwMLE default values are used (i.e. each parameter is started at zero).
<code>fixPar</code>	List of fixPar objects (see crwMLE) containing an element for each individual. If only one <code>fixPar</code> is provided, then the same parameters are held fixed to the given value for each individual. If <code>fixPar</code> is not specified, then no parameters are fixed.
<code>method</code>	Optimization method that is passed to optim .
<code>control</code>	Control list which is passed to optim .
<code>constr</code>	List of constr objects (see crwMLE) containing an element for each individual. If only one <code>constr</code> is provided, then the same box constraints for the parameters are used for each individual.
<code>prior</code>	List of prior objects (see crwMLE) containing an element for each individual. If only one <code>prior</code> is provided, then the same prior is used for each individual.
<code>need.hess</code>	A logical value which decides whether or not to evaluate the Hessian for parameter standard errors
<code>initialSANN</code>	Control list for optim when simulated annealing is used for obtaining start values. See details
<code>attempts</code>	The number of times likelihood optimization will be attempted in cases where the fit does not converge or is otherwise non-valid. Note this is not the same as <code>retryFits</code> because <code>attempts</code> only applies when the current fit clearly does not appear to have converged; <code>retryFits</code> will proceed with additional model fitting attempts regardless of the model output.
<code>predTime</code>	List of predTime objects (see crwPredict) containing an element for each individual. <code>predTime</code> can be specified as an alternative to the automatic sequences generated according to <code>timeStep</code> . If only one <code>predTime</code> object is provided, then the same prediction times are used for each individual.

fillCols	Logical indicating whether or not to use the crawl:: fillCols function for filling in missing values in obsData for which there is a single unique value. Default: FALSE. If the output from crawlWrap is intended for analyses using fitHMM or MifitHMM , setting fillCols=TRUE should typically be avoided.
coordLevel	Character string indicating the level of the hierarchy for the location data. Ignored unless obsData includes a 'level' field.
...	Additional arguments that are ignored.

Details

- Consult [crwMLE](#) and [crwPredict](#) for further details about model fitting and prediction.
- Note that the names of the list elements corresponding to each individual in mov.model, err.model, activity, drift, theta, fixPar, constr, prior, and predTime must match the individual IDs in obsData. If only one element is provided for any of these arguments, then the same element will be applied to all individuals.

Value

A [crwData](#) or [crwHierData](#) object, i.e. a list of:

crwFits	A list of crwFit objects returned by crawl::crwMLE. See crwMLE
crwPredict	A crwPredict data frame with obsData merged with the predicted locations. See crwPredict .

The [crwData](#) object is used in [MifitHMM](#) analyses that account for temporal irregularity or location measurement error.

See Also

[MifitHMM](#), [simData](#)

Examples

```
## Not run:
# extract simulated obsData from example data
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x = ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.cor)

# Fit crwMLE models to obsData and predict locations
# at default intervals for both individuals
crwOut1 <- crawlWrap(obsData=obsData,
                      theta=c(4,0),fixPar=c(1,1,NA,NA),
                      err.model=err.model,attempts=100)

# Fit the same crwMLE models and predict locations
# at same intervals but specify for each individual using lists
crwOut2 <- crawlWrap(obsData=obsData,
                      theta=list(c(4,0),c(4,0)), fixPar=list(c(1,1,NA,NA),c(1,1,NA,NA)),
```

```

err.model=list(err.model,err.model),
predTime=list('1'=seq(1,633),'2'=seq(1,686)))

## End(Not run)

```

crwData*Constructor of crwData objects***Description**

Constructor of crwData objects

Usage

```
crwData(m)
```

Arguments

m A list of attributes of crawl output: `crwFits` (a list of `crwFit` objects) and `crwPredict` (a `crwPredict` object)

Value

An object `crwData`.

See Also

[crawlWrap](#), [MifitHMM](#)

crwHierData*Constructor of crwHierData objects***Description**

Constructor of `crwHierData` objects

Usage

```
crwHierData(m)
```

Arguments

m A list of attributes of crawl output: `crwFits` (a list of `crwFit` objects) and `crwPredict` (a `crwPredict` object)

Value

An object `crwHierData`.

See Also

[crawlWrap](#), [MIfitHMM](#)

`crwHierSim`

Constructor of crwHierSim objects

Description

Constructor of `crwHierSim` objects

Usage

`crwHierSim(m)`

Arguments

`m`

A list of attributes required for multiple imputation data generated from a `crwHierData` object using `MIfitHMM`: `miData` (a list of `momentuHMMData` objects), and `crwSimulator` (a list of `crwSimulator` objects).

`crwHierSim` objects are returned by `MIfitHMM` when argument `miData` is a `crwHierData` object and argument `fit=FALSE`.

Value

An object `crwHierSim`.

`crwSim`

Constructor of crwSim objects

Description

Constructor of `crwSim` objects

Usage

`crwSim(m)`

Arguments**m**

A list of attributes required for multiple imputation data generated from a `crwData` object using `MIfitHMM`: `miData` (a list of `momentuHMMData` objects), and `crwSimulator` (a list of `crwSimulator` objects).
`crwSim` objects are returned by `MIfitHMM` when argument `miData` is a `crwData` object and argument `fit=FALSE`.

Value

An object `crwSim`.

dbern_rcpp*Bernoulli density function***Description**

Probability density function of the Bernoulli distribution (written in C++)

Usage

```
dbern_rcpp(x, prob, foo)
```

Arguments

`x` Vector of quantiles

`prob` success probability

`foo` Unused (for compatibility with template)

Value

Vector of densities

dbeta_rcpp*Probability density function of the beta distribution (written in C++)***Description**

Probability density function of the beta distribution (written in C++)

Usage

```
dbeta_rcpp(x, shape1, shape2)
```

Arguments

x	Vector of quantiles
shape1	Shape1
shape2	Shape2

Value

Vector of densities

dcat_rcpp *Categorical density function*

Description

Probability density function of the categorical distribution (written in C++)

Usage

```
dcat_rcpp(x, prob, foo)
```

Arguments

x	Vector of quantiles
prob	success probability
foo	Unused (for compatibility with template)

Value

Vector of densities

dexp_rcpp *Exponential density function*

Description

Probability density function of the exponential distribution (written in C++)

Usage

```
dexp_rcpp(x, rate, foo)
```

Arguments

<code>x</code>	Vector of quantiles
<code>rate</code>	Rate
<code>foo</code>	Unused (for compatibility with template)

Value

Vector of densities

<code>dgamma_rcpp</code>	<i>Gamma density function</i>
--------------------------	-------------------------------

Description

Probability density function of the gamma distribution (written in C++)

Usage

```
dgamma_rcpp(x, mu, sigma)
```

Arguments

<code>x</code>	Vector of quantiles
<code>mu</code>	Mean
<code>sigma</code>	Standard deviation

Value

Vector of densities

<code>distAngle</code>	<i>Calculate distance between points y and z and turning angle between points x, y, and z</i>
------------------------	---

Description

Calculate distance between points y and z and turning angle between points x, y, and z

Usage

```
distAngle(x, y, z, type = "UTM", angleCov = TRUE)
```

Arguments

x	location 1
y	location 2
z	location 3
type	'UTM' if easting/northing provided (the default), 'LL' if longitude/latitude
angleCov	logical indicating to not return NA when x=y or y=z. Default: TRUE (i.e. NA is not returned if x=y or y=z).

Details

Used in [prepData](#) and [simData](#) to get distance and turning angle covariates between locations (x1,x2), (y1,y2) and activity center (z1,z2).

If type='LL' then distance is calculated as great circle distance using [spDistsN1](#), and turning angle is calculated based on initial bearings using [bearing](#).

Value

2-vector with first element the distance between y and z and second element the turning angle between (x,y) and (y,z).

dlnorm_rcpp

Log-normal density function

Description

Probability density function of the log-normal distribution (written in C++)

Usage

```
dlnorm_rcpp(x, meanlog, sdlog)
```

Arguments

x	Vector of quantiles
meanlog	Mean of the distribution on the log-scale
sdlog	Standard deviation of the distribution on the log-scale

Value

Vector of densities

dlogis_rcpp *logistic density function*

Description

Probability density function of the logistic distribution (written in C++)

Usage

```
dlogis_rcpp(x, location, scale)
```

Arguments

x	Vector of quantiles
location	mean of the distribution
scale	Dispersion parameter

Value

Vector of densities

dmvnorm_rcpp *C++ implementation of multivariate Normal probability density function for multiple inputs*

Description

C++ implementation of multivariate Normal probability density function for multiple inputs

Usage

```
dmvnorm_rcpp(x, mean, varcovM)
```

Arguments

x	data matrix of dimension p x n, p being the dimension of the data and n the number of data points.
mean	mean vectors matrix of dimension p x n
varcovM	list of length n of variance-covariance matrices, each of dimensions p x p.

Value

matrix of densities of dimension K x n.

dnbinom_rcpp	<i>negative binomial density function</i>
--------------	---

Description

Probability density function of the negative binomial distribution (written in C++)

Usage

```
dnbinom_rcpp(x, mu, size)
```

Arguments

- | | |
|------|--------------------------|
| x | Vector of quantiles |
| mu | Mean of the distribution |
| size | Dispersion parameter |

Value

Vector of densities

dnorm_rcpp	<i>Normal density function</i>
------------	--------------------------------

Description

Probability density function of the normal distribution (written in C++)

Usage

```
dnorm_rcpp(x, mean, sd)
```

Arguments

- | | |
|------|--|
| x | Vector of quantiles |
| mean | Mean of the distribution |
| sd | Standard deviation of the distribution |

Value

Vector of densities

<code>dpois_rcpp</code>	<i>Poisson density function</i>
-------------------------	---------------------------------

Description

Probability density function of the Poisson distribution (written in C++)

Usage

```
dpois_rcpp(x, rate, foo)
```

Arguments

<code>x</code>	Vector of quantiles
<code>rate</code>	Rate
<code>foo</code>	Unused (for compatibility with template)

Value

Vector of densities

<code>dt_rcpp</code>	<i>student t density function</i>
----------------------	-----------------------------------

Description

Probability density function of non-central student t (written in C++)

Usage

```
dt_rcpp(x, df, ncp)
```

Arguments

<code>x</code>	Vector of quantiles
<code>df</code>	degrees of freedom
<code>ncp</code>	non-centrality parameter

Value

Vector of densities

dvm_rcpp	<i>Von Mises density function</i>
----------	-----------------------------------

Description

Probability density function of the Von Mises distribution, defined as a function of the modified Bessel function of order 0 (written in C++)

Usage

```
dvm_rcpp(x, mu, kappa)
```

Arguments

x	Vector of quantiles
mu	Mean
kappa	Concentration

Value

Vector of densities

dweibull_rcpp	<i>Weibull density function</i>
---------------	---------------------------------

Description

Probability density function of the Weibull distribution (written in C++)

Usage

```
dweibull_rcpp(x, shape, scale)
```

Arguments

x	Vector of quantiles
shape	Shape
scale	Scale

Value

Vector of densities

`dwrpcauchy_rcpp` *Wrapped Cauchy density function*

Description

Probability density function of the wrapped Cauchy distribution (written in C++)

Usage

```
dwrpcauchy_rcpp(x, mu, rho)
```

Arguments

<code>x</code>	Vector of quantiles
<code>mu</code>	Mean
<code>rho</code>	Concentration

Value

Vector of densities

`exampleData` *Example dataset*

Description

These data are used in the examples and tests of functions to keep them as short as possible.

Usage

```
example
miExample
forest
```

Details

`example` is a list of the following objects for demonstrating [fitHMM](#):

- `m` A [momentuHMM](#) object
- `simPar` The parameters used to simulate data
- `par0` The initial parameters in the optimization to fit `m`

`miExample` is a list of the following objects for demonstrating [crawlWrap](#), [MIfitHMM](#), and [MIpool](#):

- `obsData` Simulated observation data with measurement error and temporal irregularity (generated by `simData`)
- `bPar` initial parameter estimates for `MIfitHMM` examples

`forest` is a simulated spatial covariate `raster` object of the `RasterLayer` class

`expandPar`

Expand vector of free working parameters to vector of all working parameters including any fixed parameters (used in fitHMM.R and nLogLike.R)

Description

Expand vector of free working parameters to vector of all working parameters including any fixed parameters (used in `fitHMM.R` and `nLogLike.R`)

Usage

```
expandPar(
  optPar,
  optInd,
  fixPar,
  wparIndex,
  betaCons,
  deltaCons,
  nbStates,
  nbCovsDelta,
  stationary,
  nbCovs,
  nbRecovs = 0,
  mixtures = 1,
  nbCovsPi = 0
)
```

Arguments

<code>optPar</code>	vector of free working parameters
<code>optInd</code>	indices of constrained parameters
<code>fixPar</code>	Vector of working parameters which are assumed known prior to fitting the model (NA indicates parameters is to be estimated)
<code>wparIndex</code>	Vector of indices for the elements of <code>fixPar</code> that are not NA
<code>betaCons</code>	Matrix of the same dimension as <code>beta0</code> composed of integers identifying any equality constraints among the t.p.m. parameters.
<code>deltaCons</code>	Matrix of the same dimension as <code>delta0</code> composed of integers identifying any equality constraints among the initial distribution working scale parameters.

<code>nbStates</code>	Number of states of the HMM
<code>nbCovsDelta</code>	Number of initial distribution covariates
<code>stationary</code>	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. Default: FALSE.
<code>nbCovs</code>	Number of t.p.m. covariates
<code>nbRecovs</code>	Number of recharge covariates
<code>mixtures</code>	Number of mixtures for the state transition probabilities
<code>nbCovsPi</code>	Number of mixture probability covariates

Value

A vector of all working parameters including any fixed parameters

Examples

```
## Not run:
nbStates <- 2
stepDist <- "gamma" # step distribution
angleDist <- "vm" # turning angle distribution

# extract data from momentuHMM example
data <- example$m$data

#### 1. fit the model to the simulated data
# define initial values for the parameters
mu0 <- c(20,70)
sigma0 <- c(10,30)
kappa0 <- c(1,1)
stepPar <- c(mu0,sigma0) # no zero-inflation, so no zero-mass included
anglePar <- kappa0 # not estimating angle mean, so not included
formula <- ~cov1+cos(cov2)

# constrain cov1 effect to state 1 -> 2 and cov2 effect to state 2 -> 1
fixPar <- list(beta=c(NA,NA,0,NA,0,NA))

m <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
             Par0=list(step=stepPar,angle=anglePar),formula=formula,fixPar=fixPar)

# convert free parameter vector (m$mod$wpar) to full set of working parameters (m$mod$estimate)
est <- momentuHMM:::expandPar(m$mod$wpar,m$conditions$optInd,unlist(m$conditions$fixPar),
                               m$conditions$wparIndex,m$conditions$betaCons,m$conditions$deltaCons,
                               nbStates,
                               ncol(m$covsDelta)-1,m$conditions$stationary,nrow(m$mle$beta)-1)

all(est==m$mod$estimate)

## End(Not run)
```

`fitHMM`*Fit a multivariate HMM to the data*

Description

Fit a (multivariate) hidden Markov model to the data provided, using numerical optimization of the log-likelihood function.

Usage

```
fitHMM(data, ...)

## S3 method for class 'momentuHMMData'
fitHMM(
  data,
  nbStates,
  dist,
  Par0,
  beta0 = NULL,
  delta0 = NULL,
  estAngleMean = NULL,
  circularAngleMean = NULL,
  formula = ~1,
  formulaDelta = NULL,
  stationary = FALSE,
  mixtures = 1,
  formulaPi = NULL,
  nlmPar = list(),
  fit = TRUE,
  DM = NULL,
  userBounds = NULL,
  workBounds = NULL,
  betaCons = NULL,
  betaRef = NULL,
  deltaCons = NULL,
  mvnCoords = NULL,
  stateNames = NULL,
  knownStates = NULL,
  fixPar = NULL,
  retryFits = 0,
  retrySD = NULL,
  optMethod = "nlm",
  control = list(),
  prior = NULL,
  modelName = NULL,
  ...
)
```

```
## S3 method for class 'momentuHierHMMData'
fitHMM(
  data,
  hierStates,
  hierDist,
  Par0,
  hierBeta = NULL,
  hierDelta = NULL,
  estAngleMean = NULL,
  circularAngleMean = NULL,
  hierFormula = NULL,
  hierFormulaDelta = NULL,
  mixtures = 1,
  formulaPi = NULL,
  nlmPar = list(),
  fit = TRUE,
  DM = NULL,
  userBounds = NULL,
  workBounds = NULL,
  betaCons = NULL,
  deltaCons = NULL,
  mvnCoords = NULL,
  knownStates = NULL,
  fixPar = NULL,
  retryFits = 0,
  retrySD = NULL,
  optMethod = "nlm",
  control = list(),
  prior = NULL,
  modelName = NULL,
  ...
)
```

Arguments

<code>data</code>	A <code>momentuHMMData</code> (as returned by <code>prepData</code> or <code>simData</code>) or a <code>momentuHierHMMData</code> (as returned by <code>prepData</code> or <code>simHierData</code>) object.
<code>...</code>	further arguments passed to or from other methods
<code>nbStates</code>	Number of states of the HMM.
<code>dist</code>	A named list indicating the probability distributions of the data streams. Currently supported distributions are 'bern', 'beta', 'cat', 'exp', 'gamma', 'Inorm', 'logis', 'negbinom', 'norm', 'mvnrm2' (bivariate normal distribution), 'mvnrm3' (trivariate normal distribution), 'pois', 'rw_norm' (normal random walk), 'rw_mvnorm2' (bivariate normal random walk), 'rw_mvnorm3' (trivariate normal random walk), 'vm', 'vmConsensus', 'weibull', and 'wrcapuchy'. For example, <code>dist=list(step='gamma', angle='vm', dives='pois')</code> indicates 3 data streams ('step', 'angle', and 'dives') and their respective probability distributions ('gamma', 'vm', and 'pois'). The

names of the data streams (e.g., 'step', 'angle', 'dives') must match component names in `data`.

<code>Par0</code>	A named list containing vectors of initial state-dependent probability distribution parameters for each data stream specified in <code>dist</code> . The parameters should be in the order expected by the pdfs of <code>dist</code> , and any zero-mass and/or one-mass parameters should be the last (if both are present, then zero-mass parameters must precede one-mass parameters). Note that zero-mass parameters are mandatory if there are zeros in data streams with a 'gamma', 'weibull', 'exp', 'lnorm', or 'beta' distribution, and one-mass parameters are mandatory if there are ones in data streams with a 'beta' distribution. For example, for a 2-state model using the Von Mises (<code>vm</code>) distribution for a data stream named 'angle' and the zero-inflated gamma distribution for a data stream named 'step', the vector of initial parameters would be something like: <code>Par0=list(step=c(mean_1,mean_2,sd_1,sd_2,zeromass_1,zero mass_2),angle=c(mean_1,mean_2,concentration_1,concentration_2))</code> . If <code>DM</code> is not specified for a given data stream, then <code>Par0</code> is on the natural (i.e., real) scale of the parameters. However, if <code>DM</code> is specified for a given data stream, then <code>Par0</code> must be on the working (i.e., beta) scale of the parameters, and the length of <code>Par0</code> must match the number of columns in the design matrix. See details below.
<code>beta0</code>	Initial matrix of regression coefficients for the transition probabilities (more information in 'Details'). Default: <code>NULL</code> . If not specified, <code>beta0</code> is initialized such that the diagonal elements of the transition probability matrix are dominant.
<code>delta0</code>	Initial value for the initial distribution of the HMM. Default: <code>rep(1/nbStates,nbStates)</code> . If <code>formulaDelta</code> includes a formula, then <code>delta0</code> must be specified as a <code>k x (nbStates-1)</code> matrix, where <code>k</code> is the number of covariates and the columns correspond to states <code>2:nbStates</code> . See details below.
<code>estAngleMean</code>	An optional named list indicating whether or not to estimate the angle mean for data streams with angular distributions (' <code>vm</code> ' and ' <code>wrpcauchy</code> '). For example, <code>estAngleMean=list(angle=TRUE)</code> indicates the angle mean is to be estimated for 'angle'. Default is <code>NULL</code> , which assumes any angle means are fixed to zero and are not to be estimated. Any <code>estAngleMean</code> elements corresponding to data streams that do not have angular distributions are ignored. <code>estAngleMean</code> is also ignored for any ' <code>vmConsensus</code> ' data streams (because the angle mean must be estimated in consensus models).
<code>circularAngleMean</code>	An optional named list indicating whether to use circular-linear (FALSE) or circular-circular (TRUE) regression on the mean of circular distributions (' <code>vm</code> ' and ' <code>wrpcauchy</code> ') for turning angles. For example, <code>circularAngleMean=list(angle=TRUE)</code> indicates the angle mean is to be estimated for 'angle' using circular-circular regression. Whenever circular-circular regression is used for an angular data stream, a corresponding design matrix (<code>DM</code>) must be specified for the data stream, and the previous movement direction (i.e., a turning angle of zero) is automatically used as the reference angle (i.e., the intercept). Any circular-circular regression covariates in <code>data</code> should therefore be relative to the previous direction of movement (instead of standard directions relative to the x-axis; see <code>prepData</code> and <code>circAngles</code>). See Duchesne et al. (2015) for specifics on the circular-circular regression model using previous movement direction as the

reference angle. Default is NULL, which assumes circular-linear regression is used for any angular distributions for which the mean angle is to be estimated. `circularAngleMean` elements corresponding to angular data streams are ignored unless the corresponding element of `estAngleMean` is TRUE. Any `circularAngleMean` elements corresponding to data streams that do not have angular distributions are ignored. `circularAngleMean` is also ignored for any 'vmConsensus' data streams (because the consensus model is a circular-circular regression model). Alternatively, `circularAngleMean` can be specified as a numeric scalar, where the value specifies the coefficient for the reference angle (i.e., directional persistence) term in the circular-circular regression model. For example, setting `circularAngleMean` to 0 specifies a circular-circular regression model with no directional persistence term (thus specifying a biased random walk instead of a biased correlated random walk). Setting `circularAngleMean` to 1 is equivalent to setting it to TRUE, i.e., a circular-circular regression model with a coefficient of 1 for the directional persistence reference angle.

<code>formula</code>	Regression formula for the transition probability covariates. Default: ~1 (no covariate effect). In addition to allowing standard functions in R formulas (e.g., <code>cos(cov)</code> , <code>cov1*cov2</code> , <code>I(cov^2)</code>), special functions include <code>cosinor(cov, period)</code> for modeling cyclical patterns, spline functions (<code>bs</code> , <code>ns</code> , <code>bSpline</code> , <code>cSpline</code> , <code>iSpline</code> , and <code>mSpline</code>), and state- or parameter-specific formulas (see details). Any formula terms that are not state- or parameter-specific are included on all of the transition probabilities.
<code>formulaDelta</code>	Regression formula for the initial distribution. Default for <code>fitHMM.momentuHMMDATA</code> : NULL (no covariate effects; both <code>delta0</code> and <code>fixPar\$delta</code> are specified on the real scale). Default for <code>fitHMM.momentuHierHMMDATA</code> : ~1 (both <code>delta0</code> and <code>fixPar\$delta</code> are specified on the working scale). Standard functions in R formulas are allowed (e.g., <code>cos(cov)</code> , <code>cov1*cov2</code> , <code>I(cov^2)</code>). When any formula is provided, then both <code>delta0</code> and <code>fixPar\$delta</code> are specified on the working scale.
<code>stationary</code>	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. Default: FALSE.
<code>mixtures</code>	Number of mixtures for the state transition probabilities (i.e. discrete random effects *sensu* DeRuiter et al. 2017). Default: <code>mixtures=1</code> .
<code>formulaPi</code>	Regression formula for the mixture distribution probabilities. Default: NULL (no covariate effects; both <code>beta0\$pi</code> and <code>fixPar\$pi</code> are specified on the real scale). Standard functions in R formulas are allowed (e.g., <code>cos(cov)</code> , <code>cov1*cov2</code> , <code>I(cov^2)</code>). When any formula is provided, then both <code>beta0\$pi</code> and <code>fixPar\$pi</code> are specified on the working scale. Note that only the covariate values from the first row for each individual ID in <code>data</code> are used (i.e. time-varying covariates cannot be used for the mixture probabilities).
<code>nlmPar</code>	List of parameters to pass to the optimization function <code>nlm</code> (which should be either <code>print.level</code> , <code>gradtol</code> , <code>stepmax</code> , <code>steptol</code> , <code>iterlim</code> , or <code>hessian</code> – see <code>nlm</code> 's documentation for more detail). For <code>print.level</code> , the default value of 0 means that no printing occurs, a value of 1 means that the first and last iterations of the optimization are detailed, and a value of 2 means that each iteration of the optimization is detailed. Ignored unless <code>optMethod="nlm"</code> .

	equality constraints among the initial distribution working scale parameters. Ignored unless a formula is provided in <code>formulaDelta</code> . See details.
<code>mvnCoords</code>	Character string indicating the name of location data that are to be modeled using a multivariate normal distribution. For example, if <code>mu="mvnorm2"</code> was included in <code>dist</code> and <code>(mu.x, mu.y)</code> are location data, then <code>mvnCoords="mu"</code> needs to be specified in order for these data to be properly treated as locations in functions such as <code>plot.momentuHMM</code> , <code>plot.miSum</code> , <code>plot.miHMM</code> , <code>plotSpatialCov</code> , and <code>MIpool</code> .
<code>stateNames</code>	Optional character vector of length <code>nbStates</code> indicating state names.
<code>knownStates</code>	Vector of values of the state process which are known prior to fitting the model (if any). Default: <code>NULL</code> (states are not known). This should be a vector with length the number of rows of <code>'data'</code> ; each element should either be an integer (the value of the known states) or <code>NA</code> if the state is not known.
<code>fixPar</code>	An optional list of vectors indicating parameters which are assumed known prior to fitting the model. Default: <code>NULL</code> (no parameters are fixed). For data streams, each element of <code>fixPar</code> should be a vector of the same name and length as the corresponding element of <code>Par0</code> . For transition probability parameters, the corresponding element of <code>fixPar</code> must be named “beta” and have the same dimensions as <code>beta0</code> . For initial distribution parameters, the corresponding element of <code>fixPar</code> must be named “delta” and have the same dimensions as <code>delta0</code> . Each parameter should either be numeric (the fixed value of the parameter) or <code>NA</code> if the parameter is to be estimated. Corresponding <code>fixPar</code> parameters must be on the same scale as <code>Par0</code> (e.g. if <code>DM</code> is specified for a given data stream, any fixed parameters for this data stream must be on the working scale), <code>beta0</code> , and <code>delta0</code> .
<code>retryFits</code>	Non-negative integer indicating the number of times to attempt to iteratively fit the model using random perturbations of the current parameter estimates as the initial values for likelihood optimization. $\text{Normal}(0, \text{retrySD}^2)$ perturbations are used on the working scale parameters. Default: 0. When <code>retryFits>0</code> , the model with the largest log likelihood value is returned. Ignored if <code>fit=FALSE</code> .
<code>retrySD</code>	An optional list of scalars or vectors indicating the standard deviation to use for normal perturbations of each working scale parameter when <code>retryFits>0</code> . For data streams, each element of <code>retrySD</code> should be a vector of the same name and length as the corresponding element of <code>Par0</code> (if a scalar is provided, then this value will be used for all working parameters of the data stream). For transition probability parameters, the corresponding element of <code>retrySD</code> must be named “beta” and have the same dimensions as <code>beta0</code> . For initial distribution parameters, the corresponding element of <code>retrySD</code> must be named “delta” and have the same dimensions as <code>delta0</code> (if <code>delta0</code> is on the working scale) or be of length <code>nbStates-1</code> (if <code>delta0</code> is on the natural scale). Alternatively <code>retrySD</code> can be a scalar, in which case this value is used for all parameters. Default: <code>NULL</code> (in which case <code>retrySD=1</code> for data stream parameters and <code>retrySD=10</code> for initial distribution and state transition probabilities). Ignored unless <code>retryFits>0</code> .
<code>optMethod</code>	The optimization method to be used. Can be “nlm” (the default; see <code>nls</code>), “Nelder-Mead” (see <code>optim</code>), or “SANN” (see <code>optim</code>).
<code>control</code>	A list of control parameters to be passed to <code>optim</code> (ignored unless <code>optMethod="Nelder-Mead"</code> or <code>optMethod="SANN"</code>).

<code>prior</code>	A function that returns the log-density of the working scale parameter prior distribution(s). See 'Details'.
<code>modelName</code>	An optional character string providing a name for the fitted model. If provided, <code>modelName</code> will be returned in <code>print.momentuHMM</code> , <code>AIC.momentuHMM</code> , <code>AICweights</code> , and other functions.
<code>hierStates</code>	A hierarchical model structure <code>Node</code> for the states ('state'). See details.
<code>hierDist</code>	A hierarchical data structure <code>Node</code> for the data streams ('dist'). See details.
<code>hierBeta</code>	A hierarchical data structure <code>Node</code> for the matrix of initial values for the regression coefficients of the transition probabilities at each level of the hierarchy ('beta'). See details.
<code>hierDelta</code>	A hierarchical data structure <code>Node</code> for the matrix of initial values for the regression coefficients of the initial distribution at each level of the hierarchy ('delta'). See details.
<code>hierFormula</code>	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy ('formula'). Default: NULL (only hierarchical-level effects, with no covariate effects). Any formula terms that are not state- or parameter-specific are included on all of the transition probabilities within a given level of the hierarchy. See details.
<code>hierFormulaDelta</code>	A hierarchical formula structure for the initial distribution covariates for each level of the hierarchy ('formulaDelta'). Default: NULL (no covariate effects and <code>fixPar\$delta</code> is specified on the working scale).

Details

- By default the matrix `beta0` of regression coefficients for the transition probabilities has one row for the intercept, plus one row for each covariate, and one column for each non-diagonal element of the transition probability matrix. For example, in a 3-state HMM with 2 formula covariates, the matrix `beta` has three rows (intercept + two covariates) and six columns (six non-diagonal elements in the 3x3 transition probability matrix - filled in row-wise). In a covariate-free model (default), `beta0` has one row, for the intercept. While the diagonal elements are by default the reference elements, other elements can serve as the reference using the `betaRef` argument. For example, in a 3-state model, setting `betaRef=c(3,2,3)` changes the reference elements to state transition 1 -> 3 for state 1 (instead of 1 -> 1), state transition 2 -> 2 for state 2 (same as default), and state transition 3 -> 3 for state 3 (same as default).
- When covariates are not included in `formulaDelta` (i.e. `formulaDelta=NULL`), then `delta0` (and `fixPar$delta`) are specified as a vector of length `nbStates` that sums to 1. When any formula is specified for `formulaDelta` (e.g. `formulaDelta=~1`, `formulaDelta=~cov1`), then `delta0` (and `fixPar$delta`) must be specified as a `k x (nbStates-1)` matrix of working parameters, where `k` is the number of regression coefficients and the columns correspond to states 2:`nbStates`. For example, in a 3-state HMM with `formulaDelta=~cov1+cov2`, the matrix `delta0` has three rows (intercept + two covariates) and 2 columns (corresponding to states 2 and 3). The initial distribution working parameters are transformed to the real scale as `exp(covsDelta*Delta)/rowSums(exp(covsDelta*Delta))`, where `covsDelta` is the `N x k` design matrix, `Delta=cbind(rep(0,k),delta0)` is a `k x nbStates` matrix of working parameters, and `N=length(unique(data$ID))`.

- The choice of initial parameters (particularly `Par0` and `beta0`) is crucial to fit a model. The algorithm might not find the global optimum of the likelihood function if the initial parameters are poorly chosen.
- If `DM` is specified for a particular data stream, then the initial values are specified on the working (i.e., beta) scale of the parameters. The working scale of each parameter is determined by the link function used. If a parameter `P` is bound by (0,Inf) then the working scale is the `log(P)` scale. If the parameter bounds are (-pi,pi) then the working scale is `tan(P/2)` unless circular-circular regression is used. Otherwise if the parameter bounds are finite then `logit(P)` is the working scale. However, when both zero- and one-inflation are included, then a multinomial logit link is used because the sum of the zeromass and onemass probability parameters cannot exceed 1. The function `getParDM` is intended to help with obtaining initial values on the working scale when specifying a design matrix and other parameter constraints (see example below). When circular-circular regression is specified using `circularAngleMean`, the working scale for the mean turning angle is not as easily interpretable, but the link function is `atan2(sin(X)*B,1+cos(X)*B)`, where `X` are the angle covariates and `B` the angle coefficients (see Duchesne et al. 2015). Under this formulation, the reference turning angle is 0 (i.e., movement in the same direction as the previous time step). In other words, the mean turning angle is zero when the coefficient(s) `B=0`.
- Circular-circular regression in `momentuHMM` is designed for turning angles (not bearings) as computed by `simData` and `prepData`. Any circular-circular regression angle covariates for time step `t` should therefore be relative to the previous direction of movement for time step `t-1`. In other words, circular-circular regression covariates for time step `t` should be the turning angle between the direction of movement for time step `t-1` and the standard direction of the covariate relative to the x-axis for time step `t`. If provided standard directions in radians relative to the x-axis (where 0 = east, $\pi/2$ = north, π = west, and $-\pi/2$ = south), `circAngles` or `prepData` can perform this calculation for you.

When the circular-circular regression model is used, the special function `angleFormula(cov, strength, by)` can be used in `DM` for the mean of angular distributions (i.e. '`vm`', '`vmConsensus`', and '`wr-pcauchy`'), where `cov` is an angle covariate (e.g. wind direction), `strength` is an optional positive real covariate (e.g. wind speed), and `by` is an optional factor variable for individual- or group-level effects (e.g. ID, sex). The `strength` argument allows angle covariates to be weighted based on their relative strength or importance at time step `t` as in Rivest et al. (2016). In this case, the link function for the mean angle is `atan2((Z * sin(X)) %*% B,1+(Z * cos(X)) %*% B)`, where `X` are the angle covariates, `Z` the strength covariates, and `B` the angle coefficients (see Rivest et al. 2016).

- State-specific formulas can be specified in `DM` using special formula functions. These special functions can take the names `paste0("state",1:nbStates)` (where the integer indicates the state-specific formula). For example, `DM=list(step=list(mean=~cov1+state1(cov2),sd=~cov2+state2(cov1)))` includes `cov1` on the mean parameter for all states, `cov2` on the mean parameter for state 1, `cov2` on the `sd` parameter for all states, and `cov1` on the `sd` parameter for state 2.
- State- and parameter-specific formulas can be specified for transition probabilities in `formula` using special formula functions. These special functions can take the names `paste0("state",1:nbStates)` (where the integer indicates the current state from which transitions occur), `paste0("toState",1:nbStates)` (where the integer indicates the state to which transitions occur), or `paste0("betaCol",nbStates*(nbStates-1))` (where the integer indicates the column of the beta matrix). For example with `nbStates=3`, `formula=~cov1+betaCol1(cov2)+state3(cov3)+toState1(cov4)` includes `cov1` on all transition probability parameters, `cov2` on the beta column corresponding to the transition from

state 1->2, cov3 on transition probabilities from state 3 (i.e., beta columns corresponding to state transitions 3->1 and 3->2), and cov4 on transition probabilities to state 1 (i.e., beta columns corresponding to state transitions 2->1 and 3->1).

- `betaCons` can be used to impose equality constraints among the t.p.m. parameters. It must be a matrix of the same dimension as `beta0` and be composed of integers, where each beta parameter is sequentially indexed in a column-wise fashion (see [checkPar0](#)). Parameter indices in `betaCons` must therefore be integers between 1 and `nbStates*(nbStates-1)`.

Use of `betaCons` is perhaps best demonstrated by example. If no constraints are imposed (the default), then `betaCons=matrix(1:length(beta0), nrow(beta0), ncol(beta0))` such that each beta parameter is (column-wise) sequentially identified by a unique integer. Suppose we wish to fit a model with `nbStates=3` states and a covariate ('cov1') on the t.p.m. With no constraints on the t.p.m., we would have `betaCons=matrix(1:(2*(nbStates*(nbStates-1))), nrow=2, ncol=nbStates-2, "1 -> 3", "2 -> 1", "2 -> 3", "3 -> 1", "3 -> 2"))`. If we then wanted to constrain the t.p.m. such that the covariate effect is identical for transitions from state 1 to states 2 and 3 (and vice versa), we have `betaCons=matrix(c(1,2,3,2,5,6,7,8,9,6,11,12), nrow=2, ncol=nbStates*(nbStates-2), "1 -> 3", "2 -> 1", "2 -> 3", "3 -> 1", "3 -> 2"))`; this results in 10 estimated beta parameters (instead of 12), the "cov1" effects indexed by a "2" ("1 -> 2" and "1 -> 3") constrained to be equal, and the "cov1" effects indexed by a "6" ("2 -> 1" and "3 -> 1") constrained to be equal.

Now suppose we instead wish to constrain these sets of state transition probabilities to be equal, i.e., $\text{Pr}(1 \rightarrow 2) = \text{Pr}(1 \rightarrow 3)$ and $\text{Pr}(2 \rightarrow 1) = \text{Pr}(3 \rightarrow 1)$; then we have `betaCons=matrix(c(1,2,1,2,5,6,7,8,5,6 -> 2, "1 -> 3", "2 -> 1", "2 -> 3", "3 -> 1", "3 -> 2"))`

- Cyclical relationships (e.g., hourly, monthly) may be modeled in DM or formula using the `cosinor(x, period)` special formula function for covariate `x` and sine curve period of time length `period`. For example, if the data are hourly, a 24-hour cycle can be modeled using `~cosinor(cov1, 24)`, where the covariate `cov1` is a repeating sequential series of integers indicating the hour of day ($0, 1, \dots, 23, 0, 1, \dots, 23, 0, 1, \dots$) (note that `fitHMM` will not do this for you, the appropriate covariate must be included in `data`; see example below). The `cosinor(x, period)` function converts `x` to 2 covariates `cosinorCos(x)=cos(2*pi*x/period)` and `cosinorSin(x)=sin(2*pi*x/period)` for inclusion in the model (i.e., 2 additional parameters per state). The amplitude of the sine wave is thus `sqrt(B_cos^2 + B_sin^2)`, where `B_cos` and `B_sin` are the working parameters corresponding to `cosinorCos(x)` and `cosinorSin(x)`, respectively (e.g., see Cornelissen 2014).
- Similar to that used in [crawlWrap](#), the `prior` argument is a user-specified function that returns the log-density of the working scale parameter prior distribution(s). In addition to including prior information about parameters, one area where priors can be particularly useful is for handling numerical issues that can arise when parameters are near a boundary. When parameters are near boundaries, they can wander into the "nether regions" of the parameter space during optimization. For example, setting `prior=function(par) {sum(dnorm(par, 0, sd, log=TRUE))}` with a reasonably large `sd` (e.g. 100 or 1000) can help prevent working parameters from straying too far along the real line. Here `par` is the vector of working scale parameters (as returned by `fitHMM`, e.g., see `examplemodestimate`) in the following order: data stream working parameters (in order `names(dist)`), beta working parameters, and delta working parameters. Instead of specifying the same prior on all parameters, different priors could be specified on different parameters (and not all parameters must have user-specified priors). For example, `prior=function(par){dnorm(par[3], 0, 100, log=TRUE)}` would only specify a prior for the third working parameter. Note that the `prior` function must return a scalar on the log

scale. See 'harbourSealExample.R' in the "vignettes" source directory for an example using the `prior` argument.

- `fitHMM.momentuHierHMMDATA` is very similar to `fitHMM.momentuHMMDATA` except that instead of simply specifying the number of states (`nbStates`), distributions (`dist`), and a single t.p.m. formula (`formula`), the `hierStates` argument specifies the hierarchical nature of the states, the `hierDist` argument specifies the hierarchical nature of the data streams, and the `hierFormula` argument specifies a t.p.m. formula for each level of the hierarchy. All are specified as `Node` objects from the `data.tree` package.

Value

A `momentuHMM` or `momentuHierHMM` object, i.e. a list of:

<code>mle</code>	A named list of the maximum likelihood estimates of the parameters of the model (if the numerical algorithm has indeed identified the global maximum of the likelihood function). Elements are included for the parameters of each data stream, as well as <code>beta</code> (transition probabilities regression coefficients - more information in 'Details'), <code>gamma</code> (transition probabilities on real scale, based on mean covariate values if <code>formula</code> includes covariates), and <code>delta</code> (initial distribution).
<code>CIreal</code>	Standard errors and 95% confidence intervals on the real (i.e., natural) scale of parameters
<code>CIbeta</code>	Standard errors and 95% confidence intervals on the beta (i.e., working) scale of parameters
<code>data</code>	The <code>momentuHMMDATA</code> or <code>momentuHierHMMDATA</code> object
<code>mod</code>	List object returned by the numerical optimizer <code>nlm</code> or <code>optim</code> . Items in <code>mod</code> include the best set of free working parameters found (<code>wpar</code>), the best full set of working parameters including any fixed parameters (<code>estimate</code>), the value of the likelihood at <code>estimate</code> (<code>minimum</code>), the estimated variance-covariance matrix at <code>estimate</code> (<code>Sigma</code>), and the elapsed time in seconds for the optimization (<code>elapsedTime</code>).
<code>conditions</code>	Conditions used to fit the model, e.g., bounds (parameter bounds), distributions, <code>zeroInflation</code> , <code>estAngleMean</code> , <code>stationary</code> , <code>formula</code> , <code>DM</code> , <code>fullDM</code> (full design matrix), etc.
<code>rawCovs</code>	Raw covariate values for transition probabilities, as found in the data (if any). Used in <code>plot.momentuHMM</code> .
<code>stateNames</code>	The names of the states.
<code>knownStates</code>	Vector of values of the state process which are known.
<code>covsDelta</code>	Design matrix for initial distribution.

References

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See Also

[getParDM](#), [prepData](#), [simData](#)
[simHierData](#)

Examples

```
nbStates <- 2
stepDist <- "gamma" # step distribution
angleDist <- "vm" # turning angle distribution

# extract data from momentuHMM example
data <- example$m$data

#### 1. fit the model to the simulated data
# define initial values for the parameters
mu0 <- c(20,70)
sigma0 <- c(10,30)
kappa0 <- c(1,1)
stepPar <- c(mu0,sigma0) # no zero-inflation, so no zero-mass included
anglePar <- kappa0 # not estimating angle mean, so not included
formula <- ~cov1+cos(cov2)
```

```

m <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
             Par0=list(step=stepPar,angle=anglePar),formula=formula)

print(m)

## Not run:
### 2. fit the exact same model to the simulated data using DM formulas
# Get initial values for the parameters on working scale
Par0 <- getParDM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                  Par=list(step=stepPar,angle=anglePar),
                  DM=list(step=list(mean=~1,sd=~1),angle=list(concentration=~1)))

mDMf <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                 Par0=Par0,formula=formula,
                 DM=list(step=list(mean=~1,sd=~1),angle=list(concentration=~1)))

print(mDMf)

### 3. fit the exact same model to the simulated data using DM matrices
# define DM
DMm <- list(step=diag(4),angle=diag(2))

# user-specified dimnames not required but are recommended
dimnames(DMm$step) <- list(c("mean_1","mean_2","sd_1","sd_2"),
                           c("mean_1:(Intercept)","mean_2:(Intercept)",
                             "sd_1:(Intercept)","sd_2:(Intercept)"))
dimnames(DMm$angle) <- list(c("concentration_1","concentration_2"),
                           c("concentration_1:(Intercept)","concentration_2:(Intercept)"))

mDMm <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                 Par0=Par0,formula=formula,
                 DM=DMm)

print(mDMm)

### 4. fit step mean parameter covariate model to the simulated data using DM
stepDMf <- list(mean=~cov1,sd=~1)
Par0 <- getParDM(data,nbStates,list(step=stepDist,angle=angleDist),
                  Par=list(step=stepPar,angle=anglePar),
                  DM=list(step=stepDMf,angle=DMm$angle))
mDMfcov <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                     Par0=Par0,
                     formula=formula,
                     DM=list(step=stepDMf,angle=DMm$angle))

print(mDMfcov)

### 5. fit the exact same step mean parameter covariate model using DM matrix
stepDMm <- matrix(c(1,0,0,0,"cov1",0,0,0,0,1,0,0,0,"cov1",0,0,
                     0,0,1,0,0,0,0,1),4,6,dimnames=list(c("mean_1","mean_2","sd_1","sd_2"),
                     c("mean_1:(Intercept)","mean_1:cov1","mean_2:(Intercept)","mean_2:cov1",
                       "sd_1:(Intercept)","sd_2:(Intercept)")))
Par0 <- getParDM(data,nbStates,list(step=stepDist,angle=angleDist),

```

```

    Par=list(step=stepPar,angle=anglePar),
    DM=list(step=stepDMm,angle=DMm$angle))
mDMmcov <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                     Par0=Par0,
                     formula=formula,
                     DM=list(step=stepDMm,angle=DMm$angle))

print(mDMmcov)

### 6. fit circular-circular angle mean covariate model to the simulated data using DM

# Generate fake circular covariate using circAngles
data$cov3 <- circAngles(refAngle=2*atan(rnorm(nrow(data))),data)

# Fit circular-circular regression model for angle mean
# Note no intercepts are estimated for angle means because these are by default
# the previous movement direction (i.e., a turning angle of zero)
mDMcircf <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                     Par0=list(step=stepPar,angle=c(0,0,Par0$angle)),
                     formula=formula,
                     estAngleMean=list(angle=TRUE),
                     circularAngleMean=list(angle=TRUE),
                     DM=list(angle=list(mean=~cov3,concentration=~1)))

print(mDMcircf)

### 7. fit the exact same circular-circular angle mean model using DM matrices

# Note no intercept terms are included in DM for angle means because the intercept is
# by default the previous movement direction (i.e., a turning angle of zero)
mDMcircm <- fitHMM(data=data,nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                     Par0=list(step=stepPar,angle=c(0,0,Par0$angle)),
                     formula=formula,
                     estAngleMean=list(angle=TRUE),
                     circularAngleMean=list(angle=TRUE),
                     DM=list(angle=matrix(c("cov3",0,0,0,0,"cov3",0,0,0,0,1,0,0,0,0,1),4,4)))

print(mDMcircm)

### 8. Cosinor and state-dependent formulas
nbStates<-2
dist<-list(step="gamma")
Par<-list(step=c(100,1000,50,100))

# include 24-hour cycle on all transition probabilities
# include 12-hour cycle on transitions from state 2
formula=~cosinor(hour24,24)+state2(cosinor(hour12,12))

# specify appropriate covariates
covs<-data.frame(hour24=0:23,hour12=0:11)

beta<-matrix(c(-1.5,1,1,NA,NA,-1.5,-1,-1,1,1),5,2)
# row names for beta not required but can be helpful

```

```

rownames(beta)<-c("(Intercept)",
                     "cosinorCos(hour24, 24)",
                     "cosinorSin(hour24, 24)",
                     "cosinorCos(hour12, 12)",
                     "cosinorSin(hour12, 12)")
data.cos<-simData(nbStates=nbStates,dist=dist,Par=Par,
                  beta=beta,formula=formula,covs=covs)

m.cosinor<-fitHMM(data.cos,nbStates=nbStates,dist=dist,Par0=Par,formula=formula)
m.cosinor

#### 9. Piecewise constant B-spline on step length mean and angle concentration
nObs <- 1000 # length of simulated track
cov <- data.frame(time=1:nObs) # time covariate for splines
dist <- list(step="gamma",angle="vm")
stepDM <- list(mean=~splines2::bspline(time,df=2,degree=0),sd=~1)
angleDM <- list(mean=~1,concentration=~splines2::bspline(time,df=2,degree=0))
DM <- list(step=stepDM,angle=angleDM)
Par <- list(step=c(log(1000),1,-1,log(100)),angle=c(0,log(10),2,-5))

data.spline<-simData(obsPerAnimal=nObs,nbStates=1,dist=dist,Par=Par,DM=DM,covs=cov)

Par0 <- list(step=Par$step,angle=Par$angle[-1])
m.spline<-fitHMM(data.spline,nbStates=1,dist=dist,Par0=Par0,
                    DM=list(step=stepDM,
                            angle=angleDM["concentration"]))

#### 10. Initial state (delta) based on covariate
nObs <- 100
dist <- list(step="gamma",angle="vm")
Par <- list(step=c(100,1000,50,100),angle=c(0,0,0.01,0.75))

# create sex covariate
cov <- data.frame(sex=factor(rep(c("F","M"),each=nObs))) # sex covariate
formulaDelta <- ~ sex + 0

# Female begins in state 1, male begins in state 2
delta <- matrix(c(-100,100),2,1,dimnames=list(c("sexF","sexM"),"state 2"))

data.delta<-simData(nbAnimals=2,obsPerAnimal=nObs,nbStates=2,dist=dist,Par=Par,
                     delta=delta,formulaDelta=formulaDelta,covs=cov)

Par0 <- list(step=Par$step, angle=Par$angle[3:4])
m.delta <- fitHMM(data.delta, nbStates=2, dist=dist, Par0 = Par0,
                     formulaDelta=formulaDelta)

#### 11. Two mixtures based on covariate
nObs <- 100
nbAnimals <- 20
dist <- list(step="gamma",angle="vm")
Par <- list(step=c(100,1000,50,100),angle=c(0,0,0.1,2))

# create sex covariate

```

```

cov <- data.frame(sex=factor(rep(c("F","M"),each=nObs*nbAnimals/2)))
formulaPi <- ~ sex + 0

# Females more likely in mixture 1, males more likely in mixture 2
beta <- list(beta=matrix(c(-1.5,-0.5,-1.5,-3),2,2),
              pi=matrix(c(-2,2),2,1,dimnames=list(c("sexF","sexM"),"mix2")))

data.mix<-simData(nbAnimals=nbAnimals,obsPerAnimal=nObs,nbStates=2,dist=dist,Par=Par,
                  beta=beta,formulaPi=formulaPi,mixtures=2,covs=cov)

Par0 <- list(step=Par$step, angle=Par$angle[3:4])
m.mix <- fitHMM(data.mix, nbStates=2, dist=dist, Par0 = Par0,
                  beta0=beta, formulaPi=formulaPi,mixtures=2)

## End(Not run)

```

formatHierHMM

*Convert hierarchical HMM structure to a conventional HMM***Description**

Convert hierarchical HMM structure to a conventional HMM

Usage

```

formatHierHMM(
  data,
  hierStates,
  hierDist,
  hierBeta = NULL,
  hierDelta = NULL,
  hierFormula = NULL,
  hierFormulaDelta = NULL,
  mixtures = 1,
  workBounds = NULL,
  betaCons = NULL,
  deltaCons = NULL,
  fixPar = NULL,
  checkData = TRUE
)

```

Arguments

- | | |
|-------------------|---|
| data | <code>momentuHierHMMData</code> object or a data frame containing the data streams and covariates. |
| hierStates | A hierarchical data structure <code>Node</code> for the states ('state'). See fitHMM . |
| hierDist | A hierarchical data structure <code>Node</code> for the data streams ('dist'). See fitHMM . |

<code>hierBeta</code>	A hierarchical data structure <code>Node</code> for the matrix of initial values for the regression coefficients of the transition probabilities at each level of the hierarchy ('beta'). See fitHMM .
<code>hierDelta</code>	A hierarchical data structure <code>Node</code> for the matrix of initial values for the regression coefficients of the initial distribution at each level of the hierarchy ('delta'). See fitHMM .
<code>hierFormula</code>	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy ('formula'). See fitHMM . Default: NULL (only hierarchical-level effects, with no covariate effects).
<code>hierFormulaDelta</code>	A hierarchical formula structure for the initial distribution covariates for each level of the hierarchy ('formulaDelta'). See fitHMM . Default: NULL (no covariate effects and <code>fixPar\$delta</code> is specified on the working scale).
<code>mixtures</code>	Number of mixtures for the state transition probabilities (i.e. discrete random effects *sensu* DeRuiter et al. 2017). See fitHMM . Default: <code>mixtures=1</code> .
<code>workBounds</code>	A list with elements named 'beta' and/or 'delta', where each element is a hierarchical data structure <code>Node</code> indicating t.p.m. and initial distribution working parameter bounds ('workBounds') for parameters in <code>hierBeta</code> and <code>hierDelta</code> , respectively.
<code>betaCons</code>	A hierarchical data structure <code>Node</code> indicating t.p.m. constraints ('betaCons') among parameters in <code>hierBeta</code> at each level of the hierarchy.
<code>deltaCons</code>	A hierarchical data structure <code>Node</code> indicating initial distribution constraints ('deltaCons') among parameters in <code>hierDelta</code> at each level of the hierarchy.
<code>fixPar</code>	A list with elements named 'beta' and/or 'delta', where each element is a hierarchical data structure <code>Node</code> indicating t.p.m. and initial distribution parameters in <code>hierBeta</code> and <code>hierDelta</code> , respectively, which are assumed known.
<code>checkData</code>	logical indicating whether or not to check the suitability of data for the specified hierarchy. Ignored unless data is provided. Default: TRUE.

Value

A list of arguments needed for specifying a hierarchical HMM as a conventional HMM in [fitHMM](#) or [MIfitHMM](#), including:

<code>nbStates</code>	See fitHMM .
<code>dist</code>	See fitHMM .
<code>formula</code>	See fitHMM .
<code>formulaDelta</code>	See fitHMM .
<code>beta0</code>	See fitHMM .
<code>delta0</code>	See fitHMM .
<code>betaRef</code>	See fitHMM .
<code>betaCons</code>	See fitHMM .
<code>deltaCons</code>	See fitHMM .
<code>fixPar</code>	See fitHMM .
<code>workBounds</code>	See fitHMM .
<code>stateNames</code>	See fitHMM .

getCovNames	<i>Get names of any covariates used in probability distribution parameters</i>
-------------	--

Description

Get names of any covariates used in probability distribution parameters

Usage

```
getCovNames(m, p, distname)
```

Arguments

m	<code>momentuHMM</code> object
p	list returned by <code>parDef</code>
distname	Name of the data stream

Value

A list of:

DMterms	Names of all covariates included in the design matrix for the data stream
DMpartems	A list of the names of all covariates for each of the probability distribution parameters

getDM_rcpp	<i>Get design matrix</i>
------------	--------------------------

Description

Loop for creating full design matrix (X) from pseudo-design matrix (DM). Written in C++. Used in `getDM`.

Usage

```
getDM_rcpp(X, covs, DM, nr, nc, cov, nbObs)
```

Arguments

X	full design matrix
covs	matrix of covariates
DM	pseudo design matrix
nr	number of rows in design matrix
nc	number of column in design matrix
cov	covariate names
nbObs	number of observations

Value

full design matrix (X)

`getPar`

Get starting values from momentuHMM, miHMM, or miSum object returned by fitHMM, MI-fitHMM, or MIpool

Description

Get starting values from momentuHMM, miHMM, or miSum object returned by fitHMM, MI-fitHMM, or MIpool

Usage

`getPar(m)`

Arguments

`m` A `momentuHMM`, `miHMM`, or `miSum` object.

Value

A list of parameter values (Par, beta, delta) that can be used as starting values in `fitHMM` or `MIfitHMM`

See Also

[getPar0](#), [getParDM](#)

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m
Par <- getPar(m)
```

getPar0	<i>Get starting values for new model from existing momentuHMM or momentuHierHMM model fit</i>
---------	---

Description

For nested models, this function will extract starting parameter values (i.e., `Par0` in `fitHMM` or `MifitHMM`) from an existing `momentuHMM` or `momentuHierHMM` model fit based on the provided arguments for the new model. Any parameters that are not in common between `model` and the new model (as specified by the arguments) are set to `0`. This function is intended to help users incrementally build and fit more complicated models from simpler nested models (and vice versa).

Usage

```
getPar0(model, ...)

## Default S3 method:
getPar0(
  model,
  nbStates = length(model$stateNames),
  estAngleMean = model$conditions$estAngleMean,
  circularAngleMean = model$conditions$circularAngleMean,
  formula = model$conditions$formula,
  formulaDelta = model$conditions$formulaDelta,
  stationary = model$conditions$stationary,
  mixtures = model$conditions$mixtures,
  formulaPi = model$conditions$formulaPi,
  DM = model$conditions$DM,
  betaRef = model$conditions$betaRef,
  stateNames = model$stateNames,
  ...
)

## S3 method for class 'hierarchical'
getPar0(
  model,
  hierStates = model$conditions$hierStates,
  estAngleMean = model$conditions$estAngleMean,
  circularAngleMean = model$conditions$circularAngleMean,
  hierFormula = model$conditions$hierFormula,
  hierFormulaDelta = model$conditions$hierFormulaDelta,
  mixtures = model$conditions$mixtures,
  formulaPi = model$conditions$formulaPi,
  DM = model$conditions$DM,
  ...
)
```

Arguments

<code>model</code>	A <code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miHMM</code> , or <code>miSum</code> object (as returned by <code>fitHMM</code> , <code>MIfitHMM</code> , or <code>MIpool</code>)
<code>...</code>	further arguments passed to or from other methods
<code>nbStates</code>	Number of states in the new model. Default: <code>nbStates=length(model\$stateNames)</code>
<code>estAngleMean</code>	Named list indicating whether or not the angle mean for data streams with angular distributions ('vm' and 'wrpcauchy') are to be estimated in the new model. Default: <code>estAngleMean=model\$conditions\$estAngleMean</code>
<code>circularAngleMean</code>	Named list indicating whether circular-linear or circular-circular regression on the mean of circular distributions ('vm' and 'wrpcauchy') for turning angles are to be used in the new model. See <code>fitHMM</code> . Default: <code>circularAngleMean=model\$conditions\$circularAngleMean</code>
<code>formula</code>	Regression formula for the transition probability covariates of the new model (see <code>fitHMM</code>). Default: <code>formula=model\$conditions\$formula</code> .
<code>formulaDelta</code>	Regression formula for the initial distribution covariates of the new model (see <code>fitHMM</code>). Default: <code>formulaDelta=model\$conditions\$formulaDelta</code> .
<code>stationary</code>	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. Default: FALSE.
<code>mixtures</code>	Number of mixtures for the state transition probabilities (see <code>fitHMM</code>). Default: <code>formula=model\$conditions\$mixtures</code> .
<code>formulaPi</code>	Regression formula for the mixture distribution probabilities (see <code>fitHMM</code>). Default: <code>formula=model\$conditions\$formulaPi</code> .
<code>DM</code>	Named list indicating the design matrices to be used for the probability distribution parameters of each data stream in the new model (see <code>fitHMM</code>). Only parameters with design matrix column names that match those in <code>model\$conditions\$fullDM</code> are extracted, so care must be taken in naming columns if any elements of <code>DM</code> are specified as matrices instead of formulas. Default: <code>DM=model\$conditions\$DM</code> .
<code>betaRef</code>	Numeric vector of length <code>nbStates</code> indicating the reference elements for the t.p.m. multinomial logit link. Default: <code>formula=model\$conditions\$betaRef</code> .
<code>stateNames</code>	Character vector of length <code>nbStates</code> indicating the names and order of the states in the new model. Default: <code>stateNames=model\$stateNames[1:nbStates]</code> .
<code>hierStates</code>	A hierarchical model structure <code>Node</code> for the states (see <code>fitHMM</code>). Default: <code>hierStates=model\$conditions\$hierStates</code> .
<code>hierFormula</code>	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy (see <code>fitHMM</code>). Default: <code>hierFormula=model\$conditions\$hierFormula</code> .
<code>hierFormulaDelta</code>	A hierarchical formula structure for the initial distribution covariates for each level of the hierarchy (' <code>formulaDelta</code> '). Default: NULL (no covariate effects and <code>fixPar\$delta</code> is specified on the working scale).

Details

All other `fitHMM` (or `MIfitHMM`) model specifications (e.g., `dist`, `hierDist`, `userBounds`, `workBounds`, etc.) and data are assumed to be the same for `model` and the new model (as specified by `nbStates`,

`hierStates, estAngleMean, circularAngleMean, formula, hierFormula, formulaDelta, hierFormulaDelta, DM, etc.).`

Note that for hierarchical models, `getPar0` will return hierarchical data.tree objects (`hierBeta` and `hierDelta`) with the default values for `fixPar`, `betaCons`, and `deltaCons`; if hierarchical t.p.m. or initial distribution parameters are subject to constraints, then these must be set manually on the list object returned by `getPar0`.

Value

A named list containing starting values suitable for `Par0` and `beta0` arguments in [fitHMM](#) or [MIfitHMM](#):

<code>Par</code>	A list of vectors of state-dependent probability distribution parameters for each data stream specified in <code>model\$conditions\$dist</code>
<code>beta</code>	Matrix of regression coefficients for the transition probabilities
<code>delta</code>	Initial distribution of the HMM. Only returned if <code>stateNames</code> has the same membership as the state names for <code>model</code>

See Also

[getPar](#), [getParDM](#), [fitHMM](#), [MIfitHMM](#)

Examples

```
# model is a momentuHMM object, automatically loaded with the package
model <- example$m
data <- model$data
dist <- model$conditions$dist
nbStates <- length(model$stateNames)
estAngleMean <- model$conditions$estAngleMean

newformula <- ~cov1+cov2
Par0 <- getPar0(model, formula=newformula)

## Not run:
newModel <- fitHMM(model$data, dist=dist, nbStates=nbStates,
                     Par0=Par0$Par, beta0=Par0$beta,
                     formula=newformula,
                     estAngleMean=estAngleMean)

## End(Not run)

newDM1 <- list(step=list(mean=~cov1, sd=~cov1))
Par0 <- getPar0(model, DM=newDM1)

## Not run:
newModel1 <- fitHMM(model$data, dist=dist, nbStates=nbStates,
                     Par0=Par0$Par, beta0=Par0$beta,
                     formula=model$conditions$formula,
                     estAngleMean=estAngleMean,
                     DM=newDM1)
```

```

## End(Not run)

# same model but specify DM for step using matrices
newDM2 <- list(step=matrix(c(1,0,0,0,
                           "cov1",0,0,0,
                           0,1,0,0,
                           0,"cov1",0,0,
                           0,0,1,0,
                           0,0,"cov1",0,
                           0,0,0,1,
                           0,0,0,"cov1"),nrow=nbStates*2))

# to be extracted, new design matrix column names must match
# column names of model$conditions$fullDM
colnames(newDM2$step)<-paste0(rep(c("mean_","sd_"),each=2*nbStates),
                               rep(1:nbStates,each=2),
                               rep(c(":Intercept"),":cov1"),2*nbStates))
Par0 <- getPar0(model,DM=newDM2)

## Not run:
newModel2 <- fitHMM(model$data,dist=dist,nbStates=nbStates,
                     Par0=Par0$Par,beta0=Par0$beta,
                     formula=model$conditions$formula,
                     estAngleMean=estAngleMean,
                     DM=newDM2)

## End(Not run)

```

getParDM

Get starting values on working scale based on design matrix and other parameter constraints

Description

Convert starting values on the natural scale of data stream probability distributions to a feasible set of working scale parameters based on a design matrix and other parameter constraints.

Usage

```

getParDM(data, ...)

## Default S3 method:
getParDM(
  data = data.frame(),
  nbStates,
  dist,
  Par,
  zeroInflation = NULL,

```

```

oneInflation = NULL,
estAngleMean = NULL,
circularAngleMean = NULL,
DM = NULL,
userBounds = NULL,
workBounds = NULL,
...
)

## S3 method for class 'hierarchical'
getParDM(
  data = data.frame(),
  hierStates,
  hierDist,
  Par,
  zeroInflation = NULL,
  oneInflation = NULL,
  estAngleMean = NULL,
  circularAngleMean = NULL,
  DM = NULL,
  userBounds = NULL,
  workBounds = NULL,
  ...
)

```

Arguments

<code>data</code>	Optional <code>momentuHMMData</code> object, <code>momentuHierHMMData</code> object, or a data frame containing the covariate values. <code>data</code> must be specified if covariates are included in <code>DM</code> . If a data frame is provided, then either <code>nbStates</code> and <code>dist</code> must be specified (for a regular HMM) or <code>hierStates</code> and <code>hierDist</code> must be specified (for a hierarchical HMM).
<code>...</code>	further arguments passed to or from other methods
<code>nbStates</code>	Number of states of the HMM.
<code>dist</code>	A named list indicating the probability distributions of the data streams. Currently supported distributions are ' <code>bern</code> ', ' <code>beta</code> ', ' <code>exp</code> ', ' <code>gamma</code> ', ' <code>Inorm</code> ', ' <code>norm</code> ', ' <code>mvnrm2</code> ' (bivariate normal distribution), ' <code>mvnrm3</code> ' (trivariate normal distribution), ' <code>pois</code> ', ' <code>rw_norm</code> ' (normal random walk), ' <code>rw_mvnorm2</code> ' (bivariate normal random walk), ' <code>rw_mvnorm3</code> ' (trivariate normal random walk), ' <code>vm</code> ', ' <code>vm-Consensus</code> ', ' <code>weibull</code> ', and ' <code>wrpcauchy</code> '. For example, <code>dist=list(step='gamma', angle='vm', dives='pois')</code> indicates 3 data streams (' <code>step</code> ', ' <code>angle</code> ', and ' <code>dives</code> ') and their respective probability distributions (' <code>gamma</code> ', ' <code>vm</code> ', and ' <code>pois</code> ').
<code>Par</code>	A named list containing vectors of state-dependent probability distribution parameters for each data stream specified in <code>dist</code> . The parameters should be on the natural scale, in the order expected by the pdfs of <code>dist</code> , and any zero-mass parameters should be the last.

<code>zeroInflation</code>	A named list of logicals indicating whether the probability distributions of the data streams should be zero-inflated. If <code>zeroInflation</code> is TRUE for a given data stream, then values for the zero-mass parameters should be included in the corresponding element of <code>Par</code> . Ignored if data is a <code>momentuHMMDData</code> or <code>momentuHierHMMDData</code> object.
<code>oneInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams are one-inflated. If <code>oneInflation</code> is TRUE for a given data stream, then values for the one-mass parameters should be included in the corresponding element of <code>Par</code> . Ignored if data is a <code>momentuHMMDData</code> or <code>momentuHierHMMDData</code> object.
<code>estAngleMean</code>	An optional named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpcauchy'). Any <code>estAngleMean</code> elements corresponding to data streams that do not have angular distributions are ignored.
<code>circularAngleMean</code>	An optional named list indicating whether to use circular-linear or circular-circular regression on the mean of circular distributions ('vm' and 'wrpcauchy') for turning angles. See <code>fitHMM</code> . <code>circularAngleMean</code> elements corresponding to angular data streams are ignored unless the corresponding element of <code>estAngleMean</code> is TRUE. Any <code>circularAngleMean</code> elements corresponding to data streams that do not have angular distributions are ignored.
<code>DM</code>	A named list indicating the design matrices to be used for the probability distribution parameters of each data stream. Each element of <code>DM</code> can either be a named list of linear regression formulas or a matrix. For example, for a 2-state model using the gamma distribution for a data stream named 'step', <code>DM=list(step=list(mean=~cov1, sd=~1))</code> specifies the mean parameters as a function of the covariate 'cov1' for each state. This model could equivalently be specified as a 4x6 matrix using character strings for the covariate: <code>DM=list(step=matrix(c(1,0,0,0, 'cov1', 0,0,0,0,1,0,0,0, 'cov1', 0,0,0,0,1,0,0,0,0,1),4,6))</code> where the 4 rows correspond to the state-dependent paramaters (mean_1,mean_2,sd_1,sd_2) and the 6 columns correspond to the regression coefficients.
<code>userBounds</code>	An optional named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream. For example, for a 2-state model using the wrapped Cauchy ('wrpcauchy') distribution for a data stream named 'angle' with <code>estAngleMean\$angle=TRUE</code>), <code>userBounds=list(angle=matrix(c(-pi,-pi,-1,-1,pi,pi,1,1),4,2))</code> specifies (-1,1) bounds for the concentration parameters instead of the default [0,1) bounds.
<code>workBounds</code>	An optional named list of 2-column matrices specifying bounds on the working scale of the probability distribution, transition probability, and initial distribution parameters. For each matrix, the first column pertains to the lower bound and the second column the upper bound. For data streams, each element of <code>workBounds</code> should be a k x 2 matrix with the same name of the corresponding element of <code>Par0</code> , where k is the number of parameters. For transition probability parameters, the corresponding element of <code>workBounds</code> must be a k x 2 matrix named "beta", where k=length(beta0). For initial distribution parameters, the

corresponding element of workBounds must be a k x 2 matrix named “delta”, where k=length(delta0).

- | | |
|------------|---|
| hierStates | A hierarchical model structure Node for the states. See fitHMM . |
| hierDist | A hierarchical data structure Node for the data streams. See fitHMM . |

Details

If design matrix includes non-factor covariates, then natural scale parameters are assumed to correspond to the mean value(s) for the covariate(s) (if nrow(data)>1) and getParDM simply returns one possible solution to the system of linear equations defined by Par, DM, and any other constraints using singular value decomposition. This can be helpful for exploring relationships between the natural and working scale parameters when covariates are included, but getParDM will not necessarily return “good” starting values (i.e., Par0) for [fitHMM](#) or [MIfitHMM](#).

Value

A list of parameter values that can be used as starting values (Par0) in [fitHMM](#) or [MIfitHMM](#)

See Also

[getPar](#), [getPar0](#), [fitHMM](#), [MIfitHMM](#)

Examples

```
# data is a momentuHMMData object, automatically loaded with the package
data <- example$m$data
stepDist <- "gamma"
angleDist <- "vm"
nbStates <- 2
stepPar0 <- c(15,50,10,20) # natural scale mean_1, mean_2, sd_1, sd_2
anglePar0 <- c(0.7,1.5) # natural scale conentration_1, concentration_2

# get working parameters for 'DM' that constrains step length mean_1 < mean_2
stepDM <- matrix(c(1,1,0,0,0,1,0,0,0,1,0,0,0,0,1),4,4,
                  dimnames=list(NULL,c("mean:(Intercept)","mean_2",
                                      "sd_1:(Intercept)","sd_2:(Intercept)")))
stepworkBounds <- matrix(c(-Inf,Inf),4,2,byrow=TRUE,
                           dimnames=list(colnames(stepDM),c("lower","upper")))
stepworkBounds["mean_2","lower"] <- 0 #coefficient for 'mean_2' constrained to be positive
wPar0 <- getParDM(nbStates=2,dist=list(step=stepDist),
                  Par=list(step=stepPar0),
                  DM=list(step=stepDM),workBounds=list(step=stepworkBounds))

## Not run:
# Fit HMM using wPar0 as initial values for the step data stream
mPar <- fitHMM(data,nbStates=2,dist=list(step=stepDist,angle=angleDist),
                Par0=list(step=wPar0$step,angle=anglePar0),
                DM=list(step=stepDM),workBounds=list(step=stepworkBounds))

## End(Not run)
```

```

# get working parameters for 'DM' using 'cov1' and 'cov2' covariates
stepDM2 <- list(mean=~cov1, sd=~cov2)
wPar20 <- getParDM(data, nbStates=2, dist=list(step=stepDist),
                     Par=list(step=stepPar0),
                     DM=list(step=stepDM2))

## Not run:
# Fit HMM using wPar20 as initial values for the step data stream
mPar2 <- fitHMM(data, nbStates=2, dist=list(step=stepDist, angle=angleDist),
                  Par0=list(step=wPar20$step, angle=anglePar0),
                  DM=list(step=stepDM2))

## End(Not run)

```

getTrProbs

*Transition probability matrix***Description**

Computation of the transition probability matrix for each time step as a function of the covariates and the regression parameters.

Usage

```

getTrProbs(data, ...)

## Default S3 method:
getTrProbs(
  data,
  nbStates,
  beta,
  workBounds = NULL,
  formula = ~1,
  mixtures = 1,
  betaRef = NULL,
  stateNames = NULL,
  getCI = FALSE,
  covIndex = NULL,
  alpha = 0.95,
  ...
)

## S3 method for class 'hierarchical'
getTrProbs(
  data,
  hierStates,
  hierBeta,

```

```

workBounds = NULL,
hierFormula = NULL,
mixtures = 1,
hierDist,
getCI = FALSE,
covIndex = NULL,
alpha = 0.95,
...
)

```

Arguments

data	<code>momentuHMM</code> object, <code>momentuHierHMM</code> object, <code>miSum</code> object, <code>miHMM</code> object, <code>momentuHMMData</code> object, <code>momentuHierHMMData</code> object, or a data frame containing the covariate values.
	If a data frame is provided, then either <code>nbStates</code> must be specified (for a regular HMM) or <code>hierStates</code> and <code>hierDist</code> must be specified (for a hierarchical HMM).
...	further arguments passed to or from other methods; most are ignored if <code>data</code> is a <code>momentuHMM</code> or <code>momentuHierHMM</code> object
nbStates	Number of states. Ignored unless <code>data</code> is a data frame.
beta	Matrix of regression coefficients for the transition probabilities
workBounds	An optional named list of 2-column matrices specifying bounds on the working scale of the transition probability parameters ('beta' and, for recharge models, 'g0' and 'theta'). <code>workBounds\$beta</code> must be a $k \times 2$ matrix, where $k=\text{length}(\text{beta})$. The first column pertains to the lower bound and the second column the upper bound. Ignored unless <code>data</code> is a data frame.
formula	Regression formula for the transition probability covariates. Ignored unless <code>data</code> is a data frame.
mixtures	Number of mixtures for the state transition probabilities. Ignored unless <code>data</code> is a data frame.
betaRef	Indices of reference elements for t.p.m. multinomial logit link. Ignored unless <code>data</code> is a data frame.
stateNames	Optional character vector of length <code>nbStates</code> indicating state names. Ignored unless <code>data</code> is a data frame.
getCI	Logical indicating whether to calculate standard errors and logit-transformed confidence intervals based on fitted <code>momentuHMM</code> or <code>momentuHierHMM</code> object. Default: FALSE.
covIndex	Integer vector indicating specific rows of the data to be used in the calculations. This can be useful for reducing unnecessarily long computation times (particularly when <code>getCI=TRUE</code>), e.g., when <code>formula</code> includes factor covariates (such as ID) but no temporal covariates. Ignored if <code>data</code> is not a <code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miSum</code> , or <code>miHMM</code> object.
alpha	Significance level of the confidence intervals (if <code>getCI=TRUE</code>). Default: 0.95 (i.e. 95% CIs).

<code>hierStates</code>	A hierarchical model structure Node for the states ('state'). See fitHMM .
<code>hierBeta</code>	A hierarchical data structure Node for the matrix of regression coefficients for the transition probabilities at each level of the hierarchy, including initial values ('beta'), parameter equality constraints ('betaCons'), fixed parameters ('fixPar'), and working scale bounds ('workBounds'). See details.
<code>hierFormula</code>	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy ('formula'). See fitHMM .
<code>hierDist</code>	A hierarchical data structure Node for the data streams ('dist'). See fitHMM .

Value

If `mixtures=1`, an array of dimension `nbStates x nbStates x nrow(data)` containing the t.p.m for each observation in `data`. If `mixtures>1`, a list of length `mixtures`, where each element is an array of dimension `nbStates x nbStates x nrow(data)` containing the t.p.m for each observation in `data`.

If `getCI=TRUE` then a list of arrays is returned (with elements `est`, `se`, `lower`, and `upper`).

If a hierarchical HMM structure is provided, then a hierarchical data structure containing the state transition probabilities for each time step at each level of the hierarchy ('gamma') is returned.

Examples

```
m <- example$m
trProbs <- getTrProbs(m)

# equivalent
trProbs <- getTrProbs(m$data, nbStates=2, beta=m$mle$beta, formula=m$conditions$formula)

## Not run:
# calculate SEs and 95% CIs
trProbsSE <- getTrProbs(m, getCI=TRUE)

# plot estimates and CIs for each state transition
par(mfrow=c(2,2))
for(i in 1:2){
  for(j in 1:2){
    plot(trProbsSE$est[i,j,], type="l",
         ylim=c(0,1), ylab=paste(i,"->",j))
    arrows(1:dim(trProbsSE$est)[3],
           trProbsSE$lower[i,j,],
           1:dim(trProbsSE$est)[3],
           trProbsSE$upper[i,j,],
           length=0.025, angle=90, code=3, col=gray(.5), lwd=1.3)
  }
}

# limit calculations to first 10 observations
trProbsSE_10 <- getTrProbs(m, getCI=TRUE, covIndex=1:10)

## End(Not run)
```

HMMfits*Constructor of HMMfits objects*

Description

Constructor of HMMfits objects

Usage

```
HMMfits(m)
```

Arguments

m A list of [momentuHMM](#) objects.

HMMfits objects are returned by [MIfitHMM](#) when arguments `fit=TRUE` and `poolEstimates=FALSE`.

Value

An object HMMfits.

is.crwData*Is crwData*

Description

Check that an object is of class [crwData](#). Used in [MIfitHMM](#).

Usage

```
is.crwData(x)
```

Arguments

x An R object

Value

TRUE if x is of class [crwData](#), FALSE otherwise.

is.crwHierData *Is crwHierData*

Description

Check that an object is of class [crwHierData](#). Used in [MIfitHMM](#).

Usage

```
is.crwHierData(x)
```

Arguments

x An R object

Value

TRUE if x is of class [crwHierData](#), FALSE otherwise.

is.crwHierSim *Is crwHierSim*

Description

Check that an object is of class [crwHierSim](#).

Usage

```
is.crwHierSim(x)
```

Arguments

x An R object

Value

TRUE if x is of class [crwHierSim](#), FALSE otherwise.

`is.crwSim`*Is crwSim*

Description

Check that an object is of class `crwSim`.

Usage

```
is.crwSim(x)
```

Arguments

x An R object

Value

TRUE if x is of class `crwSim`, FALSE otherwise.

`is.HMMfits`*Is HMMfits*

Description

Check that an object is of class `HMMfits`.

Usage

```
is.HMMfits(x)
```

Arguments

x An R object

Value

TRUE if x is of class `HMMfits`, FALSE otherwise.

`is.miHMM`*Is miHMM*

Description

Check that an object is of class `miHMM`.

Usage

```
is.miHMM(x)
```

Arguments

x An R object

Value

TRUE if x is of class `miHMM`, FALSE otherwise.

`is.miSum`*Is miSum*

Description

Check that an object is of class `miSum`.

Usage

```
is.miSum(x)
```

Arguments

x An R object

Value

TRUE if x is of class `miSum`, FALSE otherwise.

is.momentuHierHMM *Is momentuHierHMM*

Description

Check that an object is of class [momentuHierHMM](#). Used in [CIreal](#), [CIBeta](#), [plotPR](#), [plotStates](#), [pseudoRes](#), [stateProbs](#), and [viterbi](#).

Usage

```
is.momentuHierHMM(x)
```

Arguments

x An R object

Value

TRUE if x is of class [momentuHierHMM](#), FALSE otherwise.

is.momentuHierHMMDATA *Is momentuHierHMMDATA*

Description

Check that an object is of class [momentuHierHMMDATA](#). Used in [fitHMM](#).

Usage

```
is.momentuHierHMMDATA(x)
```

Arguments

x An R object

Value

TRUE if x is of class [momentuHierHMMDATA](#), FALSE otherwise.

is.momentuHMM*Is momentuHMM*

Description

Check that an object is of class `momentuHMM`. Used in `CIreal`, `CIbeta`, `plotPR`, `plotStates`, `pseudoRes`, `stateProbs`, and `viterbi`.

Usage

```
is.momentuHMM(x)
```

Arguments

x An R object

Value

TRUE if x is of class `momentuHMM`, FALSE otherwise.

is.momentuHMMData*Is momentuHMMData*

Description

Check that an object is of class `momentuHMMData`. Used in `fitHMM`.

Usage

```
is.momentuHMMData(x)
```

Arguments

x An R object

Value

TRUE if x is of class `momentuHMMData`, FALSE otherwise.

logAlpha	<i>Forward log-probabilities</i>
----------	----------------------------------

Description

Used in `stateProbs` and `pseudoRes`.

Usage

```
logAlpha(m)
```

Arguments

`m` A `momentuHMM`, `miHMM`, or `miSum` object.

Value

A list of length `model$conditions$mixtures` where each element is a matrix of forward log-probabilities for each mixture.

Examples

```
## Not run:  
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package  
m <- example$m  
  
la <- momentuHMM:::logAlpha(m)  
  
## End(Not run)
```

logBeta	<i>Backward log-probabilities</i>
---------	-----------------------------------

Description

Used in `stateProbs`.

Usage

```
logBeta(m)
```

Arguments

`m` A `momentuHMM`, `miHMM`, or `miSum` object.

Value

A list of length `model$conditions$mixtures` where each element is a matrix of backward log-probabilities for each mixture.

Examples

```
## Not run:
## m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

lb <- momentuHMM:::logBeta(m)

## End(Not run)
```

MIfitHMM

*Fit HMMs to multiple imputation data***Description**

Fit a (multivariate) hidden Markov model to multiple imputation data. Multiple imputation is a method for accommodating missing data, temporal-irregularity, or location measurement error in hidden Markov models, where pooled parameter estimates reflect uncertainty attributable to observation error.

Usage

```
MIfitHMM(miData, ...)

## Default S3 method:
MIfitHMM(
  miData,
  nSims,
  ncores = 1,
  poolEstimates = TRUE,
  alpha = 0.95,
  na.rm = FALSE,
  nbStates,
  dist,
  Par0,
  beta0 = NULL,
  delta0 = NULL,
  estAngleMean = NULL,
  circularAngleMean = NULL,
  formula = ~1,
  formulaDelta = NULL,
  stationary = FALSE,
  mixtures = 1,
```

```
formulaPi = NULL,
nlmPar = NULL,
fit = TRUE,
useInitial = FALSE,
DM = NULL,
userBounds = NULL,
workBounds = NULL,
betaCons = NULL,
betaRef = NULL,
deltaCons = NULL,
mvnCoords = NULL,
stateNames = NULL,
knownStates = NULL,
fixPar = NULL,
retryFits = 0,
retrySD = NULL,
optMethod = "nlm",
control = list(),
prior = NULL,
modelName = NULL,
covNames = NULL,
spatialCovs = NULL,
centers = NULL,
centroids = NULL,
angleCovs = NULL,
altCoordNames = NULL,
method = "IS",
parIS = 1000,
dfSim = Inf,
grid.eps = 1,
crit = 2.5,
scaleSim = 1,
quad.ask = FALSE,
force.quad = TRUE,
fullPost = TRUE,
dfPostIS = Inf,
scalePostIS = 1,
thetaSamp = NULL,
...
)

## S3 method for class 'hierarchical'
MIfitHMM(
  miData,
  nSims,
  ncores = 1,
  poolEstimates = TRUE,
  alpha = 0.95,
```

```
na.rm = FALSE,  
hierStates,  
hierDist,  
Par0,  
hierBeta = NULL,  
hierDelta = NULL,  
estAngleMean = NULL,  
circularAngleMean = NULL,  
hierFormula = NULL,  
hierFormulaDelta = NULL,  
mixtures = 1,  
formulaPi = NULL,  
nlmPar = NULL,  
fit = TRUE,  
useInitial = FALSE,  
DM = NULL,  
userBounds = NULL,  
workBounds = NULL,  
betaCons = NULL,  
deltaCons = NULL,  
mvnCoords = NULL,  
knownStates = NULL,  
fixPar = NULL,  
retryFits = 0,  
retrySD = NULL,  
optMethod = "nlm",  
control = list(),  
prior = NULL,  
modelName = NULL,  
covNames = NULL,  
spatialCovs = NULL,  
centers = NULL,  
centroids = NULL,  
angleCovs = NULL,  
altCoordNames = NULL,  
method = "IS",  
parIS = 1000,  
dfSim = Inf,  
grid.eps = 1,  
crit = 2.5,  
scaleSim = 1,  
quad.ask = FALSE,  
force.quad = TRUE,  
fullPost = TRUE,  
dfPostIS = Inf,  
scalePostIS = 1,  
thetaSamp = NULL,  
...
```

)

Arguments

miData	A <code>crwData</code> object, a <code>crwHierData</code> object, a <code>crwSim</code> object, a <code>crwHierSim</code> object, a list of <code>momentuHMMDa</code> objects, or a list of <code>momentuHierHMMDa</code> objects.
...	further arguments passed to or from other methods
nSims	Number of imputations in which to fit the HMM using <code>fitHMM</code> . If <code>miData</code> is a list of <code>momentuHMMDa</code> objects, <code>nSims</code> cannot exceed the length of <code>miData</code> .
ncores	Number of cores to use for parallel processing. Default: 1 (no parallel processing).
poolEstimates	Logical indicating whether or not to calculate pooled parameter estimates across the <code>nSims</code> imputations using <code>MIpool</code> . Default: TRUE.
alpha	Significance level for calculating confidence intervals of pooled estimates when <code>poolEstimates</code> =TRUE (see <code>MIpool</code>). Default: 0.95.
na.rm	Logical indicating whether or not to exclude model fits with NA parameter estimates or standard errors from pooling when <code>poolEstimates</code> =TRUE (see <code>MIpool</code>). Default: FALSE.
nbStates	Number of states of the HMM. See <code>fitHMM</code> .
dist	A named list indicating the probability distributions of the data streams. See <code>fitHMM</code> .
Par0	A named list containing vectors of initial state-dependent probability distribution parameters for each data stream specified in <code>dist</code> . See <code>fitHMM</code> . <code>Par0</code> may also be a list of length <code>nSims</code> , where each element is a named list containing vectors of initial state-dependent probability distribution parameters for each imputation. Note that if <code>useInitial</code> =TRUE then <code>Par0</code> is ignored after the first imputation.
beta0	Initial matrix of regression coefficients for the transition probabilities. See <code>fitHMM</code> . <code>beta0</code> may also be a list of length <code>nSims</code> , where each element is an initial matrix of regression coefficients for the transition probabilities for each imputation.
delta0	Initial values for the initial distribution of the HMM. See <code>fitHMM</code> . <code>delta0</code> may also be a list of length <code>nSims</code> , where each element is the initial values for the initial distribution of the HMM for each imputation.
estAngleMean	An optional named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpcalch'). See <code>fitHMM</code> .
circularAngleMean	An optional named list indicating whether to use circular-linear or circular-circular regression on the mean of circular distributions ('vm' and 'wrpcalch') for turning angles. See <code>fitHMM</code> .
formula	Regression formula for the transition probability covariates. See <code>fitHMM</code> .
formulaDelta	Regression formula for the initial distribution. See <code>fitHMM</code> .
stationary	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. See <code>fitHMM</code> .

mixtures	Number of mixtures for the state transition probabilities (i.e. discrete random effects <i>sensu</i> DeRuiter et al. 2017). Default: <code>mixtures=1</code> .
formulaPi	Regression formula for the mixture distribution probabilities. See fitHMM .
nlmPar	List of parameters to pass to the optimization function <code>nlm</code> (which should be either <code>print.level</code> , <code>gradtol</code> , <code>stepmax</code> , <code>steptol</code> , <code>iterlim</code> , or <code>hessian</code> – see <code>nlm</code> 's documentation for more detail). For <code>print.level</code> , the default value of 0 means that no printing occurs, a value of 1 means that the first and last iterations of the optimization are detailed, and a value of 2 means that each iteration of the optimization is detailed. Ignored unless <code>optMethod="nlm"</code> .
fit	TRUE if the HMM should be fitted to the data, FALSE otherwise. See fitHMM . If <code>fit=FALSE</code> and <code>miData</code> is a <code>crwData</code> object, then <code>MIfitHMM</code> returns a list containing a <code>momentuHMMData</code> object (if <code>nSims=1</code>) or, if <code>nSims>1</code> , a <code>crwSim</code> object.
useInitial	Logical indicating whether or not to use parameter estimates for the first model fit as initial values for all subsequent model fits. If <code>ncores>1</code> then the first model is fit on a single core and then used as the initial values for all subsequent model fits on each core (in this case, the progress of the initial model fit can be followed using the <code>print.level</code> option in <code>nlmPar</code>). Default: FALSE. Ignored if <code>nSims<2</code> .
DM	An optional named list indicating the design matrices to be used for the probability distribution parameters of each data stream. See fitHMM .
userBounds	An optional named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream. See fitHMM .
workBounds	An optional named list of 2-column matrices specifying bounds on the working scale of the probability distribution, transition probability, and initial distribution parameters. See fitHMM .
betaCons	Matrix of the same dimension as <code>beta0</code> composed of integers identifying any equality constraints among the t.p.m. parameters. See fitHMM .
betaRef	Numeric vector of length <code>nbStates</code> indicating the reference elements for the t.p.m. multinomial logit link. See fitHMM .
deltaCons	Matrix of the same dimension as <code>delta0</code> composed of integers identifying any equality constraints among the initial distribution working scale parameters. Ignored unless a formula is provided in <code>formulaDelta</code> . See fitHMM .
mvnCoords	Character string indicating the name of location data that are to be modeled using a multivariate normal distribution. For example, if <code>mu="mvnorm2"</code> was included in <code>dist</code> and (<code>mu.x</code> , <code>mu.y</code>) are location data, then <code>mvnCoords="mu"</code> needs to be specified in order for these data to be properly treated as locations in functions such as <code>plot.momentuHMM</code> , <code>plot.miSum</code> , <code>plot.miHMM</code> , <code>plotSpatialCov</code> , and <code>MIpool</code> .
stateNames	Optional character vector of length <code>nbStates</code> indicating state names.
knownStates	Vector of values of the state process which are known prior to fitting the model (if any). See fitHMM . If <code>miData</code> is a list of <code>momentuHMMData</code> objects, then <code>knownStates</code> can alternatively be a list of vectors containing the known values for the state process for each element of <code>miData</code> .

<code>fixPar</code>	An optional list of vectors indicating parameters which are assumed known prior to fitting the model. See fitHMM .
<code>retryFits</code>	Non-negative integer indicating the number of times to attempt to iteratively fit the model using random perturbations of the current parameter estimates as the initial values for likelihood optimization. See fitHMM .
<code>retrySD</code>	An optional list of scalars or vectors indicating the standard deviation to use for normal perturbations of each working scale parameter when <code>retryFits > 0</code> . See fitHMM .
<code>optMethod</code>	The optimization method to be used. Can be “nlm” (the default; see nlm), “Nelder-Mead” (see optim), or “SANN” (see optim).
<code>control</code>	A list of control parameters to be passed to optim (ignored unless <code>optMethod = "Nelder-Mead"</code> or <code>optMethod = "SANN"</code>).
<code>prior</code>	A function that returns the log-density of the working scale parameter prior distribution(s). See fitHMM .
<code>modelName</code>	An optional character string providing a name for the fitted model. If provided, <code>modelName</code> will be returned in print.momentuHMM , AIC.momentuHMM , AICweights , and other functions.
<code>covNames</code>	Names of any covariates in <code>miData\$crwPredict</code> (if <code>miData</code> is a crwData object; otherwise <code>covNames</code> is ignored). See prepData . If <code>miData</code> is a crwData object, any covariate in <code>miData\$crwPredict</code> that is used in <code>formula</code> , <code>formulaDelta</code> , <code>formulaPi</code> , or <code>DM</code> must be included in <code>covNames</code> .
<code>spatialCovs</code>	List of raster layer(s) for any spatial covariates. See prepData .
<code>centers</code>	2-column matrix providing the x-coordinates (column 1) and y-coordinates (column 2) for any activity centers (e.g., potential centers of attraction or repulsion) from which distance and angle covariates will be calculated based on realizations of the position process. See prepData . Ignored unless <code>miData</code> is a crwData object.
<code>centroids</code>	List where each element is a data frame containing the x-coordinates ('x'), y-coordinates ('y'), and times (with user-specified name, e.g., 'time') for centroids (i.e., dynamic activity centers where the coordinates can change over time) from which distance and angle covariates will be calculated based on the location data. See prepData . Ignored unless <code>miData</code> is a crwData object.
<code>angleCovs</code>	Character vector indicating the names of any circular-circular regression angular covariates in <code>miData\$crwPredict</code> that need conversion from standard direction (in radians relative to the x-axis) to turning angle (relative to previous movement direction) See prepData . Ignored unless <code>miData</code> is a crwData or crwHierData object.
<code>altCoordNames</code>	Character string indicating an alternative name for the returned location data. See prepData . Ignored unless <code>miData</code> is a crwData or crwHierData object.
<code>method</code>	Method for obtaining weights for movement parameter samples. See crwSimulator . Ignored unless <code>miData</code> is a crwData object.
<code>parIS</code>	Size of the parameter importance sample. See crwSimulator . Ignored unless <code>miData</code> is a crwData object.
<code>dfSim</code>	Degrees of freedom for the t approximation to the parameter posterior. See 'df' argument in crwSimulator . Ignored unless <code>miData</code> is a crwData object.

grid.eps	Grid size for method="quadrature". See crwSimulator . Ignored unless miData is a crwData object.
crit	Criterion for deciding "significance" of quadrature points (difference in log-likelihood). See crwSimulator . Ignored unless miData is a crwData object.
scaleSim	Scale multiplier for the covariance matrix of the t approximation. See 'scale' argument in crwSimulator . Ignored unless miData is a crwData object.
quad.ask	Logical, for method='quadrature'. Whether or not the sampler should ask if quadrature sampling should take place. It is used to stop the sampling if the number of likelihood evaluations would be extreme. Default: FALSE. Ignored if ncores>1.
force.quad	A logical indicating whether or not to force the execution of the quadrature method for large parameter vectors. See crwSimulator . Default: TRUE. Ignored unless miData is a crwData object and method='`quadrature`'.
fullPost	Logical indicating whether to draw parameter values as well to simulate full posterior. See crwPostIS . Ignored unless miData is a crwData object.
dfPostIS	Degrees of freedom for multivariate t distribution approximation to parameter posterior. See 'df' argument in crwPostIS . Ignored unless miData is a crwData object.
scalePostIS	Extra scaling factor for t distribution approximation. See 'scale' argument in crwPostIS . Ignored unless miData is a crwData object.
thetaSamp	If multiple parameter samples are available in crwSimulator objects, setting thetaSamp=n will use the nth sample. Defaults to the last. See crwSimulator and crwPostIS . Ignored unless miData is a crwData object.
hierStates	A hierarchical model structure Node for the states. See fitHMM .
hierDist	A hierarchical data structure Node for the data streams. See fitHMM .
hierBeta	A hierarchical data structure Node for the matrix of initial values for the regression coefficients of the transition probabilities at each level of the hierarchy ('beta'). See fitHMM .
hierDelta	A hierarchical data structure Node for the matrix of initial values for the regression coefficients of the initial distribution at each level of the hierarchy ('delta'). See fitHMM .
hierFormula	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy. See fitHMM .
hierFormulaDelta	A hierarchical formula structure for the initial distribution covariates for each level of the hierarchy ('formulaDelta'). Default: NULL (no covariate effects and fixPar\$delta is specified on the working scale). See fitHMM .

Details

miData can either be a [crwData](#) or [crwHierData](#) object (as returned by [crawlWrap](#)), a [crwSim](#) or [crwHierSim](#) object (as returned by MIfitHMM when fit=FALSE), or a list of [momentuHMMData](#) or [momentuHierHMMData](#) objects (e.g., each element of the list as returned by [prepData](#)).

If miData is a [crwData](#) (or [crwHierData](#)) object, MIfitHMM uses a combination of [crwSimulator](#), [crwPostIS](#), [prepData](#), and [fitHMM](#) to draw nSims realizations of the position process and fit the

specified HMM to each imputation of the data. The vast majority of MIfitHMM arguments are identical to the corresponding arguments from these functions.

If miData is a `crwData` or `crwHierData` object, nSims determines both the number of realizations of the position process to draw (using `crwSimulator` and `crwPostIS`) as well as the number of HMM fits.

If miData is a `crwSim` (or `crwHierSim`) object or a list of `momentuHMMData` (or `momentuHierHMMData`) object(s), the specified HMM will simply be fitted to each of the `momentuHMMData` (or `momentuHierHMMData`) objects and all arguments related to `crwSimulator`, `crwPostIS`, or `prepData` are ignored.

Value

If nSims>1, poolEstimates=TRUE, and fit=TRUE, a `miHMM` object, i.e., a list consisting of:

`miSum` `miSum` object returned by `MIpool`.

`HMMfits` List of length nSims comprised of `momentuHMM` objects.

If poolEstimates=FALSE and fit=TRUE, a list of length nSims consisting of `momentuHMM` objects is returned.

However, if fit=FALSE and miData is a `crwData` object, then MIfitHMM returns a `crwSim` object, i.e., a list containing miData (a list of `momentuHMMData` objects) and crwSimulator (a list of `crwSimulator` objects),and most other arguments related to `fitHMM` are ignored.

References

Hooten M.B., Johnson D.S., McClintock B.T., Morales J.M. 2017. Animal Movement: Statistical Models for Telemetry Data. CRC Press, Boca Raton.

McClintock B.T. 2017. Incorporating telemetry error into hidden Markov movement models using multiple imputation. Journal of Agricultural, Biological, and Environmental Statistics.

See Also

`crawlWrap`, `crwPostIS`, `crwSimulator`, `fitHMM`, `getParDM`, `MIpool`, `prepData`

Examples

```
# Don't run because it takes too long on a single core
## Not run:
# extract simulated obsData from example data
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x= ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)

# create crwData object by fitting crwMLE models to obsData and predict locations
# at default intervals for both individuals
crwOut <- crawlWrap(obsData=obsData,
                      theta=c(4,0),fixPar=c(1,1,NA,NA),
                      err.model=err.model)
```

```

# HMM specifications
nbStates <- 2
stepDist <- "gamma"
angleDist <- "vm"
mu0 <- c(20,70)
sigma0 <- c(10,30)
kappa0 <- c(1,1)
stepPar0 <- c(mu0,sigma0)
anglePar0 <- c(-pi/2,pi/2,kappa0)
formula <- ~cov1+cos(cov2)
nbCovs <- 2
beta0 <- matrix(c(rep(-1.5,nbStates*(nbStates-1)),rep(0,nbStates*(nbStates-1)*nbCovs)),
nrow=nbCovs+1,byrow=TRUE)

# first fit HMM to best predicted position process
bestData<-prepData(crwOut,covNames=c("cov1","cov2"))
bestFit<-fitHMM(bestData,
                  nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                  Par0=list(step=stepPar0,angle=anglePar0),beta0=beta0,
                  formula=formula,estAngleMean=list(angle=TRUE))

print(bestFit)

# extract estimates from 'bestFit'
bPar0 <- getPar(bestFit)

# Fit nSims=5 imputations of the position process
miFits<-MIfitHMM(miData=crwOut,nSims=5,
                    nbStates=nbStates,dist=list(step=stepDist,angle=angleDist),
                    Par0=bPar0$Par,beta0=bPar0$beta,delta0=bPar0$delta,
                    formula=formula,estAngleMean=list(angle=TRUE),
                    covNames=c("cov1","cov2"))

# print pooled estimates
print(miFits)

## End(Not run)

```

miHMM*Constructor of miHMM objects*

Description

Constructor of miHMM objects

Usage

```
miHMM(m)
```

Arguments

- m** A list with attributes `miSum` (a `miSum` object) and `HMMfits` (a list of `momentuHMM` objects).
`miHMM` objects are returned by `MIfitHMM` when arguments `fit=TRUE`, `nSims>1`, and `poolEstimates=TRUE`.

Value

An object `miHMM`.

MIpool	<i>Calculate pooled parameter estimates and states across multiple imputations</i>
--------	--

Description

Calculate pooled parameter estimates and states across multiple imputations

Usage

```
MIpool(im, alpha = 0.95, ncores = 1, covs = NULL, na.rm = FALSE)
```

Arguments

- im** List comprised of `momentuHMM` or `momentuHierHMM` objects
- alpha** Significance level for calculating confidence intervals of pooled estimates (including location error ellipses). Default: 0.95.
- ncores** Number of cores to use for parallel processing. Default: 1 (no parallel processing).
- covs** Data frame consisting of a single row indicating the covariate values to be used in the calculation of pooled natural parameters. For any covariates that are not specified using `covs`, the means of the covariate(s) across the imputations are used (unless the covariate is a factor, in which case the first factor in the data is used). By default, no covariates are specified.
- na.rm** Logical indicating whether or not to exclude model fits with NA parameter estimates or standard errors from pooling. Default: FALSE.

Details

Pooled estimates, standard errors, and confidence intervals are calculated using standard multiple imputation formulas. Working scale parameters are pooled using `MIcombine` and t-distributed confidence intervals. Natural scale parameters and normally-distributed confidence intervals are calculated by transforming the pooled working scale parameters and, if applicable, are based on covariate means across all imputations (and/or values specified in `covs`).

The calculation of pooled error ellipses uses `dataEllipse` from the `car` package. The suggested package `car` is not automatically imported by `momentuHMM` and must be installed in order to calculate error ellipses. A warning will be triggered if the `car` package is required but not installed.

Note that pooled estimates for `timeInStates` and `stateProbs` do not include within-model uncertainty and are based entirely on across-model variability.

Value

A `miSum` object, i.e., a list comprised of model and pooled parameter summaries, including data (averaged across imputations), conditions, Par, and MIcombine (as returned by `MIcombine` for working parameters).

`miSum$Par` is a list comprised of:

<code>beta</code>	Pooled estimates for the working parameters
<code>real</code>	Estimates for the natural parameters based on pooled working parameters and covariate means (or <code>covs</code>) across imputations (if applicable)
<code>timeInStates</code>	The proportion of time steps assigned to each state
<code>states</code>	The most frequent state assignment for each time step based on the <code>viterbi</code> algorithm for each model fit
<code>stateProbs</code>	Pooled state probability estimates for each time step
<code>mixtureProbs</code>	Pooled mixture probabilities for each individual (only applies if <code>mixtures>1</code>)
<code>hierStateProbs</code>	Pooled state probability estimates for each time step at each level of the hierarchy (only applies if <code>im</code> is comprised of <code>momentuHierHMM</code> objects)

Examples

```
## Not run:
# Extract data and crawl inputs from miExample
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x= ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)

# Fit crawl to obsData
crwOut <- crawlWrap(obsData,theta=c(4,0),fixPar=c(1,1,NA,NA),
                      err.model=err.model)

# Fit four imputations
bPar <- miExample$bPar
HMMfits <- MIfitHMM(crwOut,nSims=4,poolEstimates=FALSE,
                      nbStates=2,dist=list(step="gamma",angle="vm"),
                      Par0=bPar$Par,beta0=bPar$beta,
                      formula=~cov1+cos(cov2),
                      estAngleMean=list(angle=TRUE),
                      covNames=c("cov1","cov2"))

# Pool estimates
miSum <- MIpool(HMMfits)
```

```
print(miSum)
## End(Not run)
```

miSum *Constructor of miSum objects*

Description

Constructor of **miSum** objects

Usage

```
miSum(m)
```

Arguments

- m** A list of attributes required for multiple imputation summaries: **data** (averaged across imputations), **Par** (the pooled estimates of the parameters of the model), **conditions** (conditions used to fit the model), and **MIcombine** (as returned by **MIcombine** for the working parameters).

Value

An object **miSum**.

mixtureProbs *Mixture probabilities*

Description

For a fitted model, this function computes the probability of each individual being in a particular mixture

Usage

```
mixtureProbs(m, getCI = FALSE, alpha = 0.95)
```

Arguments

- m** **momentuHMM** or **momentuHierHMM** object
- getCI** Logical indicating whether to calculate standard errors and logit-transformed confidence intervals for fitted **momentuHMM** or **momentuHierHMM** object. Default: FALSE.
- alpha** Significance level of the confidence intervals (if **getCI=TRUE**). Default: 0.95 (i.e. 95% CIs).

Details

When `getCI=TRUE`, it can take a while for large data sets and/or a large number of mixtures because the model likelihood for each individual must be repeatedly evaluated in order to numerically approximate the SEs.

Value

The matrix of individual mixture probabilities, with element [i,j] the probability of individual i being in mixture j

References

Maruotti, A., and T. Ryden. 2009. A semiparametric approach to hidden Markov models under longitudinal observations. *Statistics and Computing* 19: 381-393.

Examples

```
## Not run:
nObs <- 100
nbAnimals <- 20
dist <- list(step="gamma",angle="vm")
Par <- list(step=c(100,1000,50,100),angle=c(0,0,0.1,2))

# create sex covariate
cov <- data.frame(sex=factor(rep(c("F","M"),each=nObs*nbAnimals/2)))
formulaPi <- ~ sex + 0

# Females more likely in mixture 1, males more likely in mixture 2
beta <- list(beta=matrix(c(-1.5,-0.5,-1.5,-3),2,2),
             pi=matrix(c(-2,2),2,1,dimnames=list(c("sexF","sexM"),"mix2")))

data.mix<-simData(nbAnimals=nbAnimals,obsPerAnimal=nObs,nbStates=2,dist=dist,Par=Par,
                  beta=beta,formulaPi=formulaPi,mixtures=2,covs=cov)

Par0 <- list(step=Par$step, angle=Par$angle[3:4])
m.mix <- fitHMM(data.mix, nbStates=2, dist=dist, Par0 = Par0,
                  beta0=beta,formulaPi=formulaPi,mixtures=2)

mixProbs <- mixtureProbs(m.mix, getCI=TRUE)

## End(Not run)
```

Description

Constructor of `momentuHierHMM` objects

Usage

```
momentuHierHMM(m)
```

Arguments

- m** A list of attributes of the fitted model: `mle` (the maximum likelihood estimates of the parameters of the model), `data` (the `fitHMM` data), `mod` (the object returned by the `fitHMM` numerical optimizer `nlm` or `optim`), `conditions` (conditions used to fit the model: `hierStates`, `hierDist`, `zeroInflation`, `estAngleMean`, `circularAngleMean` `stationary`, `formula`, `userBounds`, `bounds`, `workBounds`, `DM`, etc.), `stateNames`, and `rawCovs` (optional – only if there are transition probability matrix covariates in the data).

Value

An object `momentuHierHMM`.

`momentuHierHMMData`

Constructor of momentuHierHMMData objects

Description

Constructor of `momentuHierHMMData` objects

Usage

```
momentuHierHMMData(data)
```

Arguments

- data** A dataframe containing: `ID` (the ID(s) of the observed animal(s)), `level` (the level of the hierarchy for each observation), and the data streams such as `step` (the step lengths, if any), `angle` (the turning angles, if any), `x` (either easting or longitude, if any), `y` (either norting or latitude, if any), and covariates (if any).

Value

An object `momentuHierHMMData`.

momentuHMM*Constructor of momentuHMM objects***Description**

Constructor of `momentuHMM` objects

Usage

```
momentuHMM(m)
```

Arguments

- | | |
|----------------|---|
| <code>m</code> | A list of attributes of the fitted model: <code>mle</code> (the maximum likelihood estimates of the parameters of the model), <code>data</code> (the <code>fitHMM</code> data), <code>mod</code> (the object returned by the <code>fitHMM</code> numerical optimizer <code>nlm</code> or <code>optim</code>), <code>conditions</code> (conditions used to fit the model: <code>dist</code> , <code>zeroInflation</code> , <code>estAngleMean</code> , <code>circularAngleMean</code> , <code>stationary</code> , <code>formula</code> , <code>userBounds</code> , <code>bounds</code> , <code>workBounds</code> , <code>DM</code> , etc.), <code>stateNames</code> , and <code>rawCovs</code> (optional – only if there are transition probability matrix covariates in the data). |
|----------------|---|

Value

An object `momentuHMM`.

momentuHMMData*Constructor of momentuHMMData objects***Description**

Constructor of `momentuHMMData` objects

Usage

```
momentuHMMData(data)
```

Arguments

- | | |
|-------------------|---|
| <code>data</code> | A dataframe containing: <code>ID</code> (the ID(s) of the observed animal(s)) and the data streams such as <code>step</code> (the step lengths, if any), <code>angle</code> (the turning angles, if any), <code>x</code> (either easting or longitude, if any), <code>y</code> (either norting or latitude, if any), and covariates (if any). |
|-------------------|---|

Value

An object `momentuHMMData`.

n2w*Scaling function: natural to working parameters.*

Description

Scales each data stream probability distribution parameter from its natural interval to the set of real numbers, to allow for unconstrained optimization. Used during the optimization of the log-likelihood. Parameters of any data streams for which a design matrix is specified are ignored.

Usage

```
n2w(par, bounds, beta, delta = NULL, nbStates, estAngleMean, DM, Bndind, dist)
```

Arguments

par	Named list of vectors containing the initial parameter values for each data stream.
bounds	Named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream.
beta	List of regression coefficients for the transition probabilities.
delta	Initial distribution. Default: NULL ; if the initial distribution is not estimated.
nbStates	The number of states of the HMM.
estAngleMean	Named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpcapuchy').
DM	An optional named list indicating the design matrices to be used for the probability distribution parameters of each data stream. Each element of DM can either be a named list of linear regression formulas or a matrix.
Bndind	Named list indicating whether DM is NULL with default parameter bounds for each data stream.
dist	A named list indicating the probability distributions of the data streams.

Value

A vector of unconstrained parameters.

Examples

```
## Not run:
m<-example$m
nbStates <- 2
nbCovs <- 2
parSize <- list(step=2,angle=2)
par <- list(step=c(t(m$mle$step)),angle=c(t(m$mle$angle)))
bounds <- m$conditions$bounds
beta <- matrix(rnorm(6),ncol=2,nrow=3)
delta <- c(0.6,0.4)
```

```

#working parameters
wpar <- momentuHMM::n2w(par,bounds,list(beta=beta),log(delta[-1]/delta[1]),nbStates,
m$conditions$estAngleMean,NULL,m$conditions$Bndind,
m$conditions$dist)

#natural parameter
p <-  momentuHMM:::w2n(wpar,bounds,parSize,nbStates,nbCovs,m$conditions$estAngleMean,
m$conditions$circularAngleMean,lapply(m$conditions$dist,function(x) x=="vmConsensus"),
m$conditions$stationary,m$conditions$fullDM,
m$conditions$DMind,1,m$conditions$dist,m$conditions$Bndind,
matrix(1,nrow=length(unique(m$data$ID)),ncol=1),covsDelta=m$covsDelta,
workBounds=m$conditions$workBounds)

## End(Not run)

```

nLogLike

*Negative log-likelihood function***Description**

Negative log-likelihood function

Usage

```

nLogLike(
  optPar,
  nbStates,
  formula,
  bounds,
  parSize,
  data,
  dist,
  covs,
  estAngleMean,
  circularAngleMean,
  consensus,
  zeroInflation,
  oneInflation,
  stationary = FALSE,
  fullDM,
  DMind,
  Bndind,
  knownStates,
  fixPar,
  wparIndex,
  nc,
  meanind,

```

```

covsDelta,
workBounds,
prior = NULL,
betaCons = NULL,
betaRef,
deltaCons = NULL,
optInd = NULL,
recovs = NULL,
g0covs = NULL,
mixtures = 1,
covsPi,
recharge = NULL,
aInd
)

```

Arguments

<code>optPar</code>	Vector of working parameters.
<code>nbStates</code>	Number of states of the HMM.
<code>formula</code>	Regression formula for the transition probability covariates.
<code>bounds</code>	Named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream.
<code>parSize</code>	Named list indicating the number of natural parameters of the data stream probability distributions
<code>data</code>	An object <code>momentuHMMData</code> .
<code>dist</code>	Named list indicating the probability distributions of the data streams.
<code>covs</code>	data frame containing the beta model covariates (if any)
<code>estAngleMean</code>	Named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpc Cauchy').
<code>circularAngleMean</code>	Named list indicating whether to use circular-linear or circular-circular regression on the mean of circular distributions ('vm' and 'wrpc Cauchy') for turning angles. See fitHMM .
<code>consensus</code>	Named list indicating whether to use the circular-circular regression consensus model
<code>zeroInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams are zero-inflated.
<code>oneInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams are one-inflated.
<code>stationary</code>	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. Default: FALSE.
<code>fullDM</code>	Named list containing the full (i.e. not shorthand) design matrix for each data stream.

DMind	Named list indicating whether fullDM includes individual- and/or temporal-covariates for each data stream specifies (-1,1) bounds for the concentration parameters instead of the default [0,1) bounds.
Bndind	Named list indicating whether DM is NULL with default parameter bounds for each data stream.
knownStates	Vector of values of the state process which are known prior to fitting the model (if any).
fixPar	Vector of working parameters which are assumed known prior to fitting the model (NA indicates parameters is to be estimated).
wparIndex	Vector of indices for the elements of fixPar that are not NA.
nc	indicator for zeros in fullDM
meanind	index for circular-circular regression mean angles with at least one non-zero entry in fullDM
covsDelta	data frame containing the delta model covariates (if any)
workBounds	named list of 2-column matrices specifying bounds on the working scale of the probability distribution, transition probability, and initial distribution parameters
prior	A function that returns the log-density of the working scale parameter prior distribution(s)
betaCons	Matrix of the same dimension as beta0 composed of integers identifying any equality constraints among the t.p.m. parameters.
betaRef	Indices of reference elements for t.p.m. multinomial logit link.
deltaCons	Matrix of the same dimension as delta0 composed of integers identifying any equality constraints among the initial distribution working scale parameters.
optInd	indices of constrained parameters
recovs	data frame containing the recharge model theta covariates (if any)
g0covs	data frame containing the recharge model g0 covariates (if any)
mixtures	Number of mixtures for the state transition probabilities
covsPi	data frame containing the pi model covariates
recharge	recharge model specification (only used for hierarchical models)
aInd	vector of indices of first observation for each animal

Value

The negative log-likelihood of the parameters given the data.

Examples

```
## Not run:
# data is a momentuHMMData object (as returned by prepData), automatically loaded with the package
data <- example$m$data
m<-example$m
Par <- getPar(m)
nbStates <- length(m$stateNames)
```

```

inputs <- momentuHMM:::checkInputs(nbStates,m$conditions$dist,Par$Par,m$conditions$estAngleMean,
                                  m$conditions$circularAngleMean,m$conditions$zeroInflation,m$conditions$oneInflation,
                                  m$conditions$DM,m$conditions$userBounds,
                                  m$stateNames)

wpar <- momentuHMM:::n2w(Par$Par,m$conditions$bounds,list(beta=Par$beta),
                           log(Par$delta[-1]/Par$delta[1]),nbStates,m$conditions$estAngleMean,
                           m$conditions$DM,m$conditions$Bndind,
                           m$conditions$dist)

l <- momentuHMM:::nLogLike(wpar,nbStates,m$conditions$formula,m$conditions$bounds,
                             inputs$p$parSize,data,inputs$dist,model.matrix(m$conditions$formula,data),
                             m$conditions$estAngleMean,m$conditions$circularAngleMean,inputs$consensus,
                             m$conditions$zeroInflation,m$conditions$oneInflation,m$conditions$stationary,
                             m$conditions$fullDM,m$conditions$DMind,
                             m$conditions$Bndind,m$knownStates,unlist(m$conditions$fixPar),
                             m$conditions$wparIndex,covsDelta=m$covsDelta,workBounds=m$conditions$workBounds,
                             betaRef=m$conditions$betaRef,covsPi=m$covsPi)

## End(Not run)

```

nLogLike_rcpp*Negative log-likelihood***Description**

Computation of the negative log-likelihood (forward algorithm - written in C++)

Usage

```

nLogLike_rcpp(
  nbStates,
  covs,
  data,
  dataNames,
  dist,
  Par,
  aInd,
  zeroInflation,
  oneInflation,
  stationary,
  knownStates,
  betaRef,
  mixtures
)

```

Arguments

<code>nbStates</code>	Number of states,
<code>covs</code>	Covariates,
<code>data</code>	A <code>momentuHMMData</code> object of the observations,
<code>dataNames</code>	Character vector containing the names of the data streams,
<code>dist</code>	Named list indicating the probability distributions of the data streams.
<code>Par</code>	Named list containing the state-dependent parameters of the data streams, matrix of regression coefficients for the transition probabilities ('beta'), and initial distribution ('delta').
<code>aInd</code>	Vector of indices of the rows at which the data switches to another animal
<code>zeroInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams are zero-inflated.
<code>oneInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams are one-inflated.
<code>stationary</code>	<code>false</code> if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If true, the initial distribution is considered equal to the stationary distribution. Default: <code>false</code> .
<code>knownStates</code>	Vector of values of the state process which are known prior to fitting the model (if any). Default: <code>NULL</code> (states are not known). This should be a vector with length the number of rows of 'data'; each element should either be an integer (the value of the known states) or <code>NA</code> if the state is not known.
<code>betaRef</code>	Indices of reference elements for t.p.m. multinomial logit link.
<code>mixtures</code>	Number of mixtures for the state transition probabilities

Value

Negative log-likelihood

<code>parDef</code>	<i>Parameters definition</i>
---------------------	------------------------------

Description

Parameters definition

Usage

```
parDef(
  dist,
  nbStates,
  estAngleMean,
  zeroInflation,
  oneInflation,
  DM,
  userBounds = NULL
)
```

Arguments

<code>dist</code>	Named list indicating the probability distributions of the data streams.
<code>nbStates</code>	Number of states of the HMM.
<code>estAngleMean</code>	Named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpcauchy').
<code>zeroInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams should be zero-inflated.
<code>oneInflation</code>	Named list of logicals indicating whether the probability distributions of the data streams are one-inflated.
<code>DM</code>	An optional named list indicating the design matrices to be used for the probability distribution parameters of each data stream. Each element of <code>DM</code> can either be a named list of linear regression formulas or a matrix.
<code>userBounds</code>	An optional named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream. For example, for a 2-state model using the wrapped Cauchy ('wrpcauchy') distribution for a data stream named 'angle' with <code>estAngleMean\$angle=TRUE</code>), <code>userBounds=list(angle=matrix(c(-pi,-pi,-1,-1,pi,pi,1,1),4,2))</code> specifies (-1,1) bounds for the concentration parameters instead of the default [0,1) bounds.

Value

A list of:

<code>parSize</code>	Named list indicating the number of natural parameters of the data stream probability distributions.
<code>bounds</code>	Named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream.
<code>parNames</code>	Names of parameters of the probability distribution for each data stream.
<code>Bndind</code>	Named list indicating whether <code>DM</code> is NULL with default parameter bounds for each data stream.

Examples

```
## Not run:
pD<-momentuHMM:::parDef(list(step="gamma",angle="wrpcauchy"),
  nbStates=2,list(step=FALSE,angle=FALSE),list(step=FALSE,angle=FALSE),
  list(step=FALSE,angle=FALSE),NULL,NULL)

## End(Not run)
```

plot.crwData*Plot crwData***Description**

Plot observed locations, error ellipses (if applicable), predicted locations, and prediction intervals from [crwData](#) or [crwHierData](#) object.

Usage

```
## S3 method for class 'crwData'
plot(
  x,
  animals = NULL,
  compact = FALSE,
  ask = TRUE,
  plotEllipse = TRUE,
  crawlPlot = FALSE,
  ...
)
```

Arguments

<code>x</code>	An object crwData or crwHierData (as returned by crawlWrap).
<code>animals</code>	Vector of indices or IDs of animals for which information will be plotted. Default: <code>NULL</code> ; all animals are plotted.
<code>compact</code>	<code>TRUE</code> for a compact plot (all individuals at once), <code>FALSE</code> otherwise (default – one individual at a time). Ignored unless crwPredictPlot=FALSE .
<code>ask</code>	If <code>TRUE</code> , the execution pauses between each plot.
<code>plotEllipse</code>	If <code>TRUE</code> (the default) then error ellipses are plotted (if applicable). Ignored unless crwPredictPlot=FALSE .
<code>crawlPlot</code>	Logical indicating whether or not to create individual plots using crwPredictPlot . See crwPredictPlot for details.
<code>...</code>	Further arguments for passing to crwPredictPlot

Details

In order for error ellipses to be plotted, the names for the semi-major axis, semi-minor axis, and orientation in `x$crwPredict` must respectively be `error_semimajor_axis`, `error_seminor_axis`, and `error_ellipse_orientation`.

If the [crwData](#) (or [crwHierData](#)) object was created using data generated by [simData](#) (or [simHierData](#)) or [simObsData](#), then the true locations (`mx`,`my`) are also plotted.

See Also

[crwPredictPlot](#)

Examples

```
## Not run:
# extract simulated obsData from example data
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x= ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)

# create crwData object
crwOut <- crawlWrap(obsData=obsData,
                      theta=c(4,0),fixPar=c(1,1,NA,NA),
                      err.model=err.model)

plot(crwOut,compact=TRUE,ask=FALSE,plotEllipse=FALSE)

## End(Not run)
```

plot.miHMM

Plot miHMM

Description

For multiple imputation analyses, plot the pooled data stream densities over histograms of the data, probability distribution parameters and transition probabilities as functions of the covariates, and maps of the animals' tracks colored by the decoded states.

Usage

```
## S3 method for class 'miHMM'
plot(
  x,
  animals = NULL,
  cova = NULL,
  ask = TRUE,
  breaks = "Sturges",
  hist.ylim = NULL,
  sepAnimals = FALSE,
  sepStates = FALSE,
  col = NULL,
  cumul = TRUE,
  plotTracks = TRUE,
  plotCI = FALSE,
  alpha = 0.95,
  plotStationary = FALSE,
  plotEllipse = TRUE,
  ...
)
```

Arguments

<code>x</code>	Object <code>miHMM</code> (as returned by MIfithMM)
<code>animals</code>	Vector of indices or IDs of animals for which information will be plotted. Default: <code>NULL</code> ; all animals are plotted.
<code>covs</code>	Data frame consisting of a single row indicating the covariate values to be used in plots. If none are specified, the means of any covariates appearing in the model are used (unless covariate is a factor, in which case the first factor appearing in the data is used).
<code>ask</code>	If <code>TRUE</code> , the execution pauses between each plot.
<code>breaks</code>	Histogram parameter. See <code>hist</code> documentation.
<code>hist.ylim</code>	Parameter <code>ylim</code> for the step length histograms. See <code>hist</code> documentation. Default: <code>NULL</code> ; the function sets default values.
<code>sepAnimals</code>	If <code>TRUE</code> , the data is split by individuals in the histograms. Default: <code>FALSE</code> .
<code>sepStates</code>	If <code>TRUE</code> , the data is split by states in the histograms. Default: <code>FALSE</code> .
<code>col</code>	Vector or colors for the states (one color per state).
<code>cumul</code>	If <code>TRUE</code> , the sum of weighted densities is plotted (default).
<code>plotTracks</code>	If <code>TRUE</code> , the Viterbi-decoded tracks are plotted (default).
<code>plotCI</code>	Logical indicating whether to include confidence intervals in natural parameter plots (default: <code>FALSE</code>)
<code>alpha</code>	Significance level of the confidence intervals (if <code>plotCI=TRUE</code>). Default: 0.95 (i.e. 95% CIs).
<code>plotStationary</code>	Logical indicating whether to plot the stationary state probabilities as a function of any covariates (default: <code>FALSE</code>)
<code>plotEllipse</code>	Logical indicating whether to plot error ellipses around imputed location means. Default: <code>TRUE</code> .
<code>...</code>	Additional arguments passed to <code>graphics::plot</code> and <code>graphics::hist</code> functions. These can currently include <code>asp</code> , <code>cex</code> , <code>cex.axis</code> , <code>cex.lab</code> , <code>cex.legend</code> , <code>cex.main</code> , <code>legend.pos</code> , and <code>lwd</code> . See par . <code>legend.pos</code> can be a single keyword from the list “bottomright”, “bottom”, “bottomleft”, “left”, “topleft”, “top”, “topright”, “right”, and “center”. Note that <code>asp</code> and <code>cex</code> only apply to plots of animal tracks.

Details

The state-dependent densities are weighted by the frequency of each state in the most probable state sequence (decoded with the function [viterbi](#) for each imputation). For example, if the most probable state sequence indicates that one third of observations correspond to the first state, and two thirds to the second state, the plots of the densities in the first state are weighted by a factor 1/3, and in the second state by a factor 2/3.

Examples

```

## Not run:
# Extract data from miExample
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x= ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)

# Fit crawl to obsData
crwOut <- crawlWrap(obsData,theta=c(4,0),fixPar=c(1,1,NA,NA),
                      err.model=err.model)

# Fit four imputations
bPar <- miExample$bPar
HMMfits <- MIfitHMM(crwOut,nSims=4,poolEstimates=FALSE,
                      nbStates=2,dist=list(step="gamma",angle="vm"),
                      Par0=bPar$Par,beta0=bPar$beta,
                      formula=~cov1+cos(cov2),
                      estAngleMean=list(angle=TRUE),
                      covNames=c("cov1","cov2"))

miHMM <- momentuHMM:::miHMM(list(miSum=MIpool(HMMfits),HMMfits=HMMfits))
plot(miHMM)

## End(Not run)

```

plot.miSum

Plot miSum

Description

Plot the fitted step and angle densities over histograms of the data, transition probabilities as functions of the covariates, and maps of the animals' tracks colored by the decoded states.

Usage

```

## S3 method for class 'miSum'
plot(
  x,
  animals = NULL,
  covs = NULL,
  ask = TRUE,
  breaks = "Sturges",
  hist.ylim = NULL,
  sepAnimals = FALSE,
  sepStates = FALSE,
  col = NULL,
  cumul = TRUE,

```

```

plotTracks = TRUE,
plotCI = FALSE,
alpha = 0.95,
plotStationary = FALSE,
plotEllipse = TRUE,
...
)

```

Arguments

<code>x</code>	Object <code>miSum</code> (as return by MIpool)
<code>animals</code>	Vector of indices or IDs of animals for which information will be plotted. Default: <code>NULL</code> ; all animals are plotted.
<code>covs</code>	Data frame consisting of a single row indicating the covariate values to be used in plots. If none are specified, the means of any covariates appearing in the model are used (unless covariate is a factor, in which case the first factor appearing in the data is used).
<code>ask</code>	If <code>TRUE</code> , the execution pauses between each plot.
<code>breaks</code>	Histogram parameter. See <code>hist</code> documentation.
<code>hist.ylim</code>	Parameter <code>ylim</code> for the step length histograms. See <code>hist</code> documentation. Default: <code>NULL</code> ; the function sets default values.
<code>sepAnimals</code>	If <code>TRUE</code> , the data is split by individuals in the histograms. Default: <code>FALSE</code> .
<code>sepStates</code>	If <code>TRUE</code> , the data is split by states in the histograms. Default: <code>FALSE</code> .
<code>col</code>	Vector or colors for the states (one color per state).
<code>cumul</code>	If <code>TRUE</code> , the sum of weighted densities is plotted (default).
<code>plotTracks</code>	If <code>TRUE</code> , the Viterbi-decoded tracks are plotted (default).
<code>plotCI</code>	Logical indicating whether to include confidence intervals in natural parameter plots (default: <code>FALSE</code>)
<code>alpha</code>	Significance level of the confidence intervals (if <code>plotCI=TRUE</code>). Default: 0.95 (i.e. 95% CIs).
<code>plotStationary</code>	Logical indicating whether to plot the stationary state probabilities as a function of any covariates (default: <code>FALSE</code>)
<code>plotEllipse</code>	Logical indicating whether to plot error ellipses around imputed location means. Default: <code>TRUE</code> .
<code>...</code>	Additional arguments passed to <code>graphics::plot</code> and <code>graphics::hist</code> functions. These can currently include <code>asp</code> , <code>cex</code> , <code>cex.axis</code> , <code>cex.lab</code> , <code>cex.legend</code> , <code>cex.main</code> , <code>legend.pos</code> , and <code>lwd</code> . See par . <code>legend.pos</code> can be a single keyword from the list “bottomright”, “bottom”, “bottomleft”, “left”, “topleft”, “top”, “topright”, “right”, and “center”. Note that <code>asp</code> and <code>cex</code> only apply to plots of animal tracks.

Details

The state-dependent densities are weighted by the frequency of each state in the most probable state sequence (decoded with the function `viterbi` for each imputation). For example, if the most probable state sequence indicates that one third of observations correspond to the first state, and two thirds to the second state, the plots of the densities in the first state are weighted by a factor 1/3, and in the second state by a factor 2/3.

Examples

```
## Not run:
# Extract data from miExample
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x= ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)

# Fit crawl to obsData
crwOut <- crawlWrap(obsData,theta=c(4,0),fixPar=c(1,1,NA,NA),
                      err.model=err.model)

# Fit four imputations
bPar <- miExample$bPar
HMMfits <- MIifitHMM(crwOut,nSims=4,poolEstimates=FALSE,
                      nbStates=2,dist=list(step="gamma",angle="vm"),
                      Par0=bPar$Par,beta0=bPar$beta,
                      formula=~cov1+cos(cov2),
                      estAngleMean=list(angle=TRUE),
                      covNames=c("cov1","cov2"))

# Pool estimates
miSum <- MIpool(HMMfits)
plot(miSum)

## End(Not run)
```

plot.momentuHMM

Plot momentuHMM

Description

Plot the fitted step and angle densities over histograms of the data, transition probabilities as functions of the covariates, and maps of the animals' tracks colored by the decoded states.

Usage

```
## S3 method for class 'momentuHMM'
plot(
  x,
```

```

  animals = NULL,
  covs = NULL,
  ask = TRUE,
  breaks = "Sturges",
  hist.ylim = NULL,
  sepAnimals = FALSE,
  sepStates = FALSE,
  col = NULL,
  cumul = TRUE,
  plotTracks = TRUE,
  plotCI = FALSE,
  alpha = 0.95,
  plotStationary = FALSE,
  ...
)

```

Arguments

<code>x</code>	Object <code>momentuHMM</code>
<code>animals</code>	Vector of indices or IDs of animals for which information will be plotted. Default: <code>NULL</code> ; all animals are plotted.
<code>covs</code>	Data frame consisting of a single row indicating the covariate values to be used in plots. If none are specified, the means of any covariates appearing in the model are used (unless covariate is a factor, in which case the first factor in the data is used).
<code>ask</code>	If <code>TRUE</code> , the execution pauses between each plot.
<code>breaks</code>	Histogram parameter. See <code>hist</code> documentation.
<code>hist.ylim</code>	An optional named list of vectors specifying <code>ylim=c(ymin,ymax)</code> for the data stream histograms. See <code>hist</code> documentation. Default: <code>NULL</code> ; the function sets default values for all data streams.
<code>sepAnimals</code>	If <code>TRUE</code> , the data is split by individuals in the histograms. Default: <code>FALSE</code> .
<code>sepStates</code>	If <code>TRUE</code> , the data is split by states in the histograms. Default: <code>FALSE</code> .
<code>col</code>	Vector or colors for the states (one color per state).
<code>cumul</code>	If <code>TRUE</code> , the sum of weighted densities is plotted (default).
<code>plotTracks</code>	If <code>TRUE</code> , the Viterbi-decoded tracks are plotted (default).
<code>plotCI</code>	Logical indicating whether to include confidence intervals in natural parameter plots (default: <code>FALSE</code>)
<code>alpha</code>	Significance level of the confidence intervals (if <code>plotCI=TRUE</code>). Default: 0.95 (i.e. 95% CIs).
<code>plotStationary</code>	Logical indicating whether to plot the stationary state probabilities as a function of any covariates (default: <code>FALSE</code>). Ignored unless covariate are included in formula.
<code>...</code>	Additional arguments passed to <code>graphics::plot</code> and <code>graphics::hist</code> functions. These can currently include <code>asp</code> , <code>cex</code> , <code>cex.axis</code> , <code>cex.lab</code> , <code>cex.legend</code> ,

`cex.main`, `legend.pos`, and `lwd`. See [par](#). `legend.pos` can be a single keyword from the list “bottomright”, “bottom”, “bottomleft”, “left”, “topleft”, “top”, “topright”, “right”, and “center”. Note that `asp` and `cex` only apply to plots of animal tracks.

Details

The state-dependent densities are weighted by the frequency of each state in the most probable state sequence (decoded with the function [viterbi](#)). For example, if the most probable state sequence indicates that one third of observations correspond to the first state, and two thirds to the second state, the plots of the densities in the first state are weighted by a factor 1/3, and in the second state by a factor 2/3.

Confidence intervals for natural parameters are calculated from the working parameter point and covariance estimates using finite-difference approximations of the first derivative for the transformation (see [grad](#)). For example, if `dN` is the numerical approximation of the first derivative of the transformation $N = \exp(x_1 * B_1 + x_2 * B_2)$ for covariates (x_1, x_2) and working parameters (B_1, B_2), then $\text{var}(N) = dN \%*\% \text{Sigma} \%*\% dN$, where $\text{Sigma} = \text{cov}(B_1, B_2)$, and normal confidence intervals can be constructed as $N +/- qnorm(1 - (1 - \alpha)/2) * \text{se}(N)$.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

plot(m, ask=TRUE, animals=1, breaks=20, plotCI=TRUE)
```

`plot.momentuHMMData` *Plot momentuHMMData or momentuHierHMMData*

Description

Plot `momentuHMMData` or `momentuHierHMMData`

Usage

```
## S3 method for class 'momentuHMMData'
plot(
  x,
  dataNames = c("step", "angle"),
  animals = NULL,
  compact = FALSE,
  ask = TRUE,
  breaks = "Sturges",
  ...
)
```

Arguments

x	An object <code>momentuHMMData</code> or <code>momentuHierHMMData</code>
dataNames	Names of the variables to plot. Default is <code>dataNames=c("step", "angle")</code> .
animals	Vector of indices or IDs of animals for which information will be plotted. Default: <code>NULL</code> ; all animals are plotted.
compact	TRUE for a compact plot (all individuals at once), FALSE otherwise (default – one individual at a time).
ask	If TRUE, the execution pauses between each plot.
breaks	Histogram parameter. See <code>hist</code> documentation.
...	Currently unused. For compatibility with generic method.

Examples

```
# data is a momentuHMMData object (as returned by prepData), automatically loaded with the package
data <- example$m$data

plot(data,dataNames=c("step","angle","cov1","cov2"),
      compact=TRUE, breaks=20, ask=FALSE)
```

plotPR

*Plot pseudo-residuals***Description**

Plots time series, qq-plots (against the standard normal distribution) using `qqPlot`, and sample ACF functions of the pseudo-residuals for each data stream

Usage

```
plotPR(m, lag.max = NULL, ncores = 1)
```

Arguments

m	A <code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miHMM</code> , <code>HMMfits</code> , or <code>miSum</code> object.
lag.max	maximum lag at which to calculate the acf. See <code>acf</code> .
ncores	number of cores to use for parallel processing

Details

- If some turning angles in the data are equal to pi, the corresponding pseudo-residuals will not be included. Indeed, given that the turning angles are defined on $(-\pi, \pi]$, an angle of pi results in a pseudo-residual of +Inf (check Section 6.2 of reference for more information on the computation of pseudo-residuals).

- If some data streams are zero-inflated and/or one-inflated, the corresponding pseudo-residuals are shown as segments, because pseudo-residuals for discrete data are defined as segments (see Zucchini and MacDonald, 2009, Section 6.2).
- For multiple imputation analyses, if `m` is a `miHMM` object or a list of `momentuHMM` objects, then the pseudo-residuals are individually calculated and plotted for each model fit. Note that pseudo-residuals for `miSum` objects (as returned by `MIpool`) are based on pooled parameter estimates and the means of the data values across all imputations (and therefore may not be particularly meaningful).

References

Zucchini, W. and MacDonald, I.L. 2009. Hidden Markov Models for Time Series: An Introduction Using R. Chapman & Hall (London).

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

plotPR(m)
```

`plotSat`

Plot observations on satellite image

Description

Plot tracking data on a satellite map. This function plots coordinates in longitude and latitude (not UTM), so if data coordinates are not provided in longitude and latitude, then the coordinate reference system must be provided using the `projargs` argument. This function uses the package `ggmap` to fetch a satellite image from Google. An Internet connection is required to use this function.

Usage

```
plotSat(
  data,
  zoom = NULL,
  location = NULL,
  segments = TRUE,
  compact = TRUE,
  col = NULL,
  alpha = 1,
  size = 1,
  shape = 16,
  states = NULL,
  animals = NULL,
  ask = TRUE,
```

```

    return = FALSE,
    stateNames = NULL,
    projargs = NULL
)

```

Arguments

data	Data frame or <code>momentuHMMData</code> object, with necessary fields 'x' (longitudinal direction) and 'y' (latitudinal direction). A <code>momentuHMM</code> , <code>miHMM</code> , or <code>miSum</code> object is also permitted, from which the data will be extracted. If <code>states=NULL</code> and a <code>momentuHMM</code> , <code>miHMM</code> , or <code>miSum</code> object is provided, the decoded states are automatically plotted.
zoom	The zoom level, as defined for <code>get_map</code> . Integer value between 3 (continent) and 21 (building).
location	Location of the center of the map to be plotted (this must be in the same coordinate reference system as <code>data</code>).
segments	TRUE if segments should be plotted between the observations (default), FALSE otherwise.
compact	FALSE if tracks should be plotted separately, TRUE otherwise (default).
col	Palette of colours to use for the dots and segments. If not specified, uses default palette.
alpha	Transparency argument for <code>geom_point</code> .
size	Size argument for <code>geom_point</code> .
shape	Shape argument for <code>geom_point</code> . If <code>states</code> is provided, then <code>shape</code> must either be a scalar or a vector of length <code>length(unique(states))</code> . If <code>states=NULL</code> , then <code>shape</code> must either be a scalar or a vector consisting of a value for each individual to be plotted.
states	A sequence of integers, corresponding to the decoded states for these data (such that the observations are colored by <code>states</code>).
animals	Vector of indices or IDs of animals/tracks to be plotted. Default: <code>NULL</code> ; all animals are plotted.
ask	If TRUE, the execution pauses between each plot.
return	If TRUE, the function returns a ggplot object (which can be edited and plotted manually). If FALSE, the function automatically plots the map (default).
stateNames	Optional character vector of length <code>max(states)</code> indicating state names. Ignored unless <code>states</code> is provided.
projargs	A character string of PROJ.4 projection arguments indicating the coordinate reference system for <code>data</code> and <code>location</code> coordinates (if not longitude and latitude). A <code>CRS</code> object is also permitted. If <code>projargs</code> is provided, the coordinates will be internally transformed to longitude and latitude for plotting.

Details

If the plot displays the message "Sorry, we have no imagery here", try a lower level of zoom.

References

D. Kahle and H. Wickham. `ggmap`: Spatial Visualization with `ggplot2`. *The R Journal*, 5(1), 144–161. URL: <http://journal.r-project.org/archive/2013-1/kahle-wickham.pdf>

plotSpatialCov	<i>Plot observations on raster image</i>
----------------	--

Description

Plot tracking data over a raster layer.

Usage

```
plotSpatialCov(
  data,
  spatialCov,
  segments = TRUE,
  compact = TRUE,
  col = NULL,
  alpha = 1,
  size = 1,
  shape = 16,
  states = NULL,
  animals = NULL,
  ask = TRUE,
  return = FALSE,
  stateNames = NULL
)
```

Arguments

<code>data</code>	Data frame or <code>momentuHMMData</code> object, with necessary fields 'x' (longitudinal direction) and 'y' (latitudinal direction). A <code>momentuHMM</code> , <code>miHMM</code> , or <code>miSum</code> object is also permitted, from which the data will be extracted. If <code>states=NULL</code> and a <code>momentuHMM</code> , <code>miHMM</code> , or <code>miSum</code> object is provided, the decoded states are automatically plotted.
<code>spatialCov</code>	<code>raster</code> object of the <code>RasterLayer</code> class on which to plot the location data
<code>segments</code>	TRUE if segments should be plotted between the observations (default), FALSE otherwise.
<code>compact</code>	FALSE if tracks should be plotted separately, TRUE otherwise (default).
<code>col</code>	Palette of colours to use for the dots and segments. If not specified, uses default palette.
<code>alpha</code>	Transparency argument for <code>geom_point</code> .
<code>size</code>	Size argument for <code>geom_point</code> .

<code>shape</code>	Shape argument for <code>geom_point</code> . If <code>states</code> is provided, then <code>shape</code> must either be a scalar or a vector of length <code>length(unique(states))</code> . If <code>states=NULL</code> , then <code>shape</code> must either be a scalar or a vector consisting of a value for each individual to be plotted.
<code>states</code>	A sequence of integers, corresponding to the decoded states for these data. If specified, the observations are colored by states.
<code>animals</code>	Vector of indices or IDs of animals/tracks to be plotted. Default: <code>NULL</code> ; all animals are plotted.
<code>ask</code>	If <code>TRUE</code> , the execution pauses between each plot.
<code>return</code>	If <code>TRUE</code> , the function returns a <code>ggplot</code> object (which can be edited and plotted manually). If <code>FALSE</code> , the function automatically plots the map (default).
<code>stateNames</code>	Optional character vector of length <code>max(states)</code> indicating state names. Ignored unless <code>states</code> is provided.

Examples

```
## Not run:
stepDist <- "gamma"
angleDist <- "vm"

# plot simulated data over forest raster automatically loaded with the package
spatialCov<-list(forest=forest)
data <- simData(nbAnimals=2,nbStates=2,dist=list(step=stepDist,angle=angleDist),
               Par=list(step=c(100,1000,50,100),angle=c(0,0,0.1,5)),
               beta=matrix(c(5,-10,-25,50),nrow=2,ncol=2,byrow=TRUE),
               formula=~forest,spatialCovs=spatialCov,
               obsPerAnimal=225,states=TRUE)

plotSpatialCov(data,forest,states=data$states)

## End(Not run)
```

`plotStates`

Plot states

Description

Plot the states and states probabilities.

Usage

```
plotStates(m, animals = NULL, ask = TRUE)
```

Arguments

<code>m</code>	A <code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miHMM</code> , or <code>miSum</code> object
<code>animals</code>	Vector of indices or IDs of animals for which states will be plotted.
<code>ask</code>	If <code>TRUE</code> , the execution pauses between each plot.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

# plot states for first and second animals
plotStates(m, animals=c(1,2))
```

plotStationary

Plot stationary state probabilities

Description

Plot stationary state probabilities

Usage

```
plotStationary(
  model,
  covs = NULL,
  col = NULL,
  plotCI = FALSE,
  alpha = 0.95,
  return = FALSE,
  ...
)
```

Arguments

<code>model</code>	<code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miHMM</code> , or <code>miSum</code> object
<code>covs</code>	Optional data frame consisting of a single row indicating the covariate values to be used in plots. If none are specified, the means of any covariates appearing in the model are used (unless covariate is a factor, in which case the first factor in the data is used).
<code>col</code>	Vector or colors for the states (one color per state).
<code>plotCI</code>	Logical indicating whether to include confidence intervals in plots (default: FALSE)
<code>alpha</code>	Significance level of the confidence intervals (if <code>plotCI=TRUE</code>). Default: 0.95 (i.e. 95% CIs).
<code>return</code>	Logical indicating whether to return a list containing estimates, SEs, CIs, and covariate values used to create the plots for each mixture and state. Ignored if <code>plotCI=FALSE</code> . Default: FALSE.
<code>...</code>	Additional arguments passed to <code>graphics::plot</code> . These can currently include <code>cex.axis</code> , <code>cex.lab</code> , <code>cex.main</code> , <code>legend</code> , <code>legend.pos</code> , and <code>lwd</code> . See <code>par</code> . <code>legend.pos</code> can be a single keyword from the list “bottomright”, “bottom”, “bottomleft”, “left”, “topleft”, “top”, “topright”, “right”, and “center”.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

plotStationary(m)
```

prepData

*Preprocessing of the data streams and covariates***Description**

Preprocessing of the data streams, including calculation of step length, turning angle, and covariates from location data to be suitable for analysis using [fitHMM](#).

Usage

```
prepData(data, ...)

## Default S3 method:
prepData(
  data,
  type = c("UTM", "LL"),
  coordNames = c("x", "y"),
  covNames = NULL,
  spatialCovs = NULL,
  centers = NULL,
  centroids = NULL,
  angleCovs = NULL,
  altCoordNames = NULL,
  ...
)

## S3 method for class 'hierarchical'
prepData(
  data,
  type = c("UTM", "LL"),
  coordNames = c("x", "y"),
  covNames = NULL,
  spatialCovs = NULL,
  centers = NULL,
  centroids = NULL,
  angleCovs = NULL,
  altCoordNames = NULL,
  hierLevels,
  coordLevel,
  ...
)
```

Arguments

<code>data</code>	Either a data frame of data streams or a <code>crwData</code> (or <code>crwHierData</code>) object (as returned by <code>crawlWrap</code>). If data is a data frame, it can optionally include a field ID (identifiers for the observed individuals), coordinates from which step length ('step') and turning angle ('angle') are calculated, and any covariates (with names matching <code>covNames</code> and/or <code>angleCovs</code>). If step length and turning angle are to be calculated from coordinates, the <code>coordNames</code> argument must identify the names for the x- (longitudinal) and y- (latitudinal) coordinates, and, for hierarchical data, the <code>coordLevel</code> argument must identify the level of the hierarchy at which the location data are obtained. With the exception of ID, <code>coordNames</code> , and, for hierarchical data, <code>level</code> , all variables in <code>data</code> are treated as data streams unless identified as covariates in <code>covNames</code> and/or <code>angleCovs</code> .
<code>...</code>	further arguments passed to or from other methods
<code>type</code>	'UTM' if easting/northing provided (the default), 'LL' if longitude/latitude. If <code>type='LL'</code> then step lengths are calculated in kilometers and turning angles are based on initial bearings (see <code>turnAngle</code>). Ignored if <code>data</code> is a <code>crwData</code> object.
<code>coordNames</code>	Names of the columns of coordinates in the <code>data</code> data frame. Default: <code>c("x", "y")</code> . If <code>coordNames=NULL</code> then step lengths, turning angles, and location covariates (i.e., those specified by <code>spatialCovs</code> , <code>centers</code> , and <code>angleCovs</code>) are not calculated. Ignored if <code>data</code> is a <code>crwData</code> object.
<code>covNames</code>	Character vector indicating the names of any covariates in <code>data</code> data frame. Any variables in <code>data</code> (other than ID) that are not identified in <code>covNames</code> and/or <code>angleCovs</code> are assumed to be data streams (i.e., missing values will not be accounted for).
<code>spatialCovs</code>	List of <code>raster</code> objects for spatio-temporally referenced covariates. Covariates specified by <code>spatialCovs</code> are extracted from the raster layer(s) based on the location data (and the z values for a raster <code>stack</code> or <code>brick</code>) for each time step. If an element of <code>spatialCovs</code> is a raster <code>stack</code> or <code>brick</code> , then z values must be set using <code>raster::setZ</code> and <code>data</code> must include column(s) of the corresponding z value(s) for each observation (e.g., 'time').
<code>centers</code>	2-column matrix providing the x-coordinates (column 1) and y-coordinates (column 2) for any activity centers (e.g., potential centers of attraction or repulsion) from which distance and angle covariates will be calculated based on the location data. If no row names are provided, then generic names are generated for the distance and angle covariates (e.g., 'center1.dist', 'center1.angle', 'center2.dist', 'center2.angle'); otherwise the covariate names are derived from the row names of <code>centers</code> as <code>paste0(rep(rownames(centers), each=2), c(".dist", ".angle"))</code> . As with covariates identified in <code>angleCovs</code> , note that the angle covariates for each activity center are calculated relative to the previous movement direction (instead of standard direction relative to the x-axis); this is to allow the mean turning angle to be modelled as a function of these covariates using circular-circular regression in <code>fitHMM</code> or <code>MIfitHMM</code> .
<code>centroids</code>	List where each element is a data frame containing the x-coordinates ('x'), y-coordinates ('y'), and times (with user-specified name, e.g., 'time') for centroids (i.e., dynamic activity centers where the coordinates can change over time) from which distance and angle covariates will be calculated based on the location

		data. If any centroids are specified, then data must include a column indicating the time of each observation, and this column name must match the corresponding user-specified name of the time column in <code>centroids</code> (e.g. ‘time’). Times can be numeric or POSIXt. If no list names are provided, then generic names are generated for the distance and angle covariates (e.g., ‘centroid1.dist’, ‘centroid1.angle’, ‘centroid2.dist’, ‘centroid2.angle’); otherwise the covariate names are derived from the list names of <code>centroids</code> as <code>paste0(rep(names(centroids), each=2), c(".dist", ".angle"))</code> . As with covariates identified in <code>angleCovs</code> , note that the angle covariates for each centroid are calculated relative to the previous movement direction (instead of standard direction relative to the x-axis); this is to allow the mean turning angle to be modelled as a function of these covariates using circular-circular regression in <code>fitHMM</code> or <code>MIfitHMM</code> .
angleCovs		Character vector indicating the names of any circular-circular regression angular covariates in <code>data</code> or <code>spatialCovs</code> that need conversion from standard direction (in radians relative to the x-axis) to turning angle (relative to previous movement direction) using <code>circAngles</code> .
altCoordNames		Character string indicating an alternative name for the returned location data. If provided, then <code>prepData</code> will return easting (or longitude) coordinate names as <code>paste0(altCoordNames, ".x")</code> and northing (or latitude) as <code>paste0(altCoordNames, ".y")</code> instead of <code>x</code> and <code>y</code> , respectively. This can be useful for location data that are intended to be modeled using a bivariate normal distribution (see <code>fitHMM</code>). Ignored unless <code>coordNames</code> are provided.
hierLevels		Character vector indicating the levels of the hierarchy and their order, from top (coarsest scale) to bottom (finest scale), that are included in <code>data\$level</code> . For example, for a 2-level hierarchy then <code>hierLevels=c("1", "2i", "2")</code> indicates <code>data\$level</code> for each observation can be one of three factor levels: “1” (coarse scale), “2i” (initial fine scale), and “2” (fine scale). Ignored if <code>data</code> is a <code>crwHierData</code> object.
coordLevel		Character string indicating the level of the hierarchy for the location data. If specified, then <code>data</code> must include a ‘level’ field indicating the level of the hierarchy for each observation. Ignored if <code>coordNames</code> is <code>NULL</code> or <code>data</code> is a <code>crwHierData</code> object.

Details

- If `data` is a `crwData` (or `crwHierData`) object, the `momentuHMMData` (or `momentuHierHMMData`) object created by `prepData` includes step lengths and turning angles calculated from the best predicted locations (i.e., `crwData$crwPredict$mu.x` and `crwData$crwPredict$mu.y`). Prior to using `prepData`, additional data streams or covariates unrelated to location (including z-values associated with `spatialCovs` raster stacks or bricks) can be merged with the `crwData` (or `crwHierData`) object using `crawlMerge`.
- For hierarchical data, `data` must include a ‘level’ field indicating the level of the hierarchy for each observation, and, for location data identified by `coordNames`, the `coordLevel` argument must indicate the level of the hierarchy at which the location data are obtained.

Value

An object `momentuHMMData` or `momentuHierHMMData`, i.e., a dataframe of:

ID	The ID(s) of the observed animal(s)
...	Data streams (e.g., 'step', 'angle', etc.)
x	Either easting or longitude (if coordNames is specified or data is a crwData object)
y	Either norting or latitude (if coordNames is specified or data is a crwData object)
...	Covariates (if any)

See Also

[crawlMerge](#), [crawlWrap](#), [crwData](#)
[crwHierData](#)

Examples

```

coord1 <- c(1,2,3,4,5,6,7,8,9,10)
coord2 <- c(1,1,1,2,2,2,1,1,1,2)
cov1 <- rnorm(10)

data <- data.frame(coord1=coord1,coord2=coord2,cov1=cov1)
d <- prepData(data,coordNames=c("coord1","coord2"),covNames="cov1")

# include additional data stream named 'omega'
omega <- rbeta(10,1,1)
data <- data.frame(coord1=coord1,coord2=coord2,omega=omega,cov1=cov1)
d <- prepData(data,coordNames=c("coord1","coord2"),covNames="cov1")

# include 'forest' example raster layer as covariate
data <- data.frame(coord1=coord1*1000,coord2=coord2*1000)
spatialCov <- list(forest=forest)
d <- prepData(data,coordNames=c("coord1","coord2"),spatialCovs=spatialCov)

# include 2 activity centers
data <- data.frame(coord1=coord1,coord2=coord2,cov1=cov1)
d <- prepData(data,coordNames=c("coord1","coord2"),covNames="cov1",
              centers=matrix(c(0,10,0,10),2,2,dimnames=list(c("c1","c2"),NULL)))

# include centroid
data <- data.frame(coord1=coord1,coord2=coord2,cov1=cov1,time=1:10)
d <- prepData(data,coordNames=c("coord1","coord2"),covNames="cov1",
              centroid=list(centroid=data.frame(x=coord1+rnorm(10),
                                                y=coord2+rnorm(10),
                                                time=1:10)))

# Include angle covariate that needs conversion to
# turning angle relative to previous movement direction
u <- rnorm(10) # horizontal component
v <- rnorm(10) # vertical component
cov2 <- atan2(v,u)
data <- data.frame(coord1=coord1,coord2=coord2,cov1=cov1,cov2=cov2)

```

```
d <- prepData(data,coordNames=c("coord1","coord2"),covNames="cov1",
angleCovs="cov2")
```

print.miHMM

Print miHMM

Description

Print miHMM

Usage

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

Arguments

x	A miHMM object.
...	Currently unused. For compatibility with generic method.

Examples

```
## Not run:
# Extract data from miExample
obsData <- miExample$obsData

# error ellipse model
err.model <- list(x= ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)

# Fit crawl to obsData
crwOut <- crawlWrap(obsData,theta=c(4,0),fixPar=c(1,1,NA,NA),
err.model=err.model)

# Fit four imputations
bPar <- miExample$bPar
HMMfits <- MIfitHMM(crwOut,nSims=4,poolEstimates=FALSE,
nbStates=2,dist=list(step="gamma",angle="vm"),
Par0=bPar$Par,beta0=bPar$beta,
formula=~cov1+cos(cov2),
estAngleMean=list(angle=TRUE),
covNames=c("cov1","cov2"))

miHMM <- momentuHMM:::miHMM(list(miSum=MIpool(HMMfits),HMMfits=HMMfits))
print(miHMM)

## End(Not run)
```

print.miSum	<i>Print miSum</i>
-------------	--------------------

Description

Print miSum

Usage

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

Arguments

x	A miSum object.
...	Currently unused. For compatibility with generic method.

Examples

```
## Not run:  
# Extract data from miExample  
obsData <- miExample$obsData  
  
# error ellipse model  
err.model <- list(x = ~ ln.sd.x - 1, y = ~ ln.sd.y - 1, rho = ~ error.corr)  
  
# Fit crawl to obsData  
crwOut <- crawlWrap(obsData, theta=c(4,0), fixPar=c(1,1,NA,NA),  
                      err.model=err.model)  
  
# Fit four imputations  
bPar <- miExample$bPar  
HMMfits <- MIfitHMM(crwOut, nSims=4, poolEstimates=FALSE,  
                      nbStates=2, dist=list(step="gamma", angle="vm"),  
                      Par0=bPar$Par, beta0=bPar$beta,  
                      formula=~cov1+cos(cov2),  
                      estAngleMean=list(angle=TRUE),  
                      covNames=c("cov1", "cov2"))  
  
# Pool estimates  
miSum <- MIpool(HMMfits)  
print(miSum)  
  
## End(Not run)
```

`print.momentuHMM` *Print momentuHMM*

Description

Print `momentuHMM`

Usage

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

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

Arguments

- `x` A `momentuHMM` object.
- `...` Currently unused. For compatibility with generic method.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

print(m)
```

`pseudoRes` *Pseudo-residuals*

Description

The pseudo-residuals of `momentuHMM` models, as described in Zucchini and McDonad (2009).

Usage

```
pseudoRes(m, ncores = 1)
```

Arguments

- `m` A `momentuHMM`, `miHMM`, `HMMfits`, or `miSum` object.
- `ncores` number of cores to use for parallel processing

Details

If some turning angles in the data are equal to pi, the corresponding pseudo-residuals will not be included. Indeed, given that the turning angles are defined on $(-\pi, \pi]$, an angle of π results in a pseudo-residual of $+\infty$ (check Section 6.2 of reference for more information on the computation of pseudo-residuals).

A continuity adjustment (adapted from Harte 2017) is made for discrete probability distributions. When the data are near the boundary (e.g. 0 for “pois”; 0 and 1 for “bern”), then the pseudo residuals can be a poor indicator of lack of fit.

For multiple imputation analyses, if m is a `miHMM` object or a list of `momentuHMM` objects, then the pseudo-residuals are individually calculated for each model fit. Note that pseudo-residuals for `miSum` objects (as returned by `MIpool`) are based on pooled parameter estimates and the means of the data values across all imputations (and therefore may not be particularly meaningful).

Value

If m is a `momentuHMM`, `miHMM`, or `miSum` object, a list of pseudo-residuals for each data stream (e.g., ‘stepRes’, ‘angleRes’) is returned. If m is a list of `momentuHMM` objects, then a list of length `length(m)` is returned where each element is a list of pseudo-residuals for each data stream.

References

Harte, D. 2017. HiddenMarkov: Hidden Markov Models. R package version 1.8-8.

Zucchini, W. and MacDonald, I.L. 2009. Hidden Markov Models for Time Series: An Introduction Using R. Chapman & Hall (London).

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m
res <- pseudoRes(m)
stats::qqnorm(res$stepRes)
stats::qqnorm(res$angleRes)
```

Description

Approximate individual-level random effects estimation for state transition probabilities based on Burnham & White (2002)

Usage

```
randomEffects(
  m,
  Xformula = ~1,
  alpha = 0.95,
  ncores = 1,
  nlmPar = list(),
  fit = TRUE,
  retryFits = 0,
  retrySD = NULL,
  optMethod = "nlm",
  control = list(),
  modelName = NULL,
  ...
)
```

Arguments

<code>m</code>	A momentuHMM object.
<code>Xformula</code>	Formula for the design matrix of the random effects model. The default <code>Xformula=~1</code> specifies an intercept-only model with no additional individual covariate effects.
<code>alpha</code>	Significance level of the confidence intervals. Default: 0.95 (i.e. 95% CIs).
<code>ncores</code>	number of cores to use for parallel processing
<code>nlmPar</code>	List of parameters to pass to the optimization function <code>nlm</code> . See fitHMM .
<code>fit</code>	TRUE if the HMM should be re-fitted at the shrinkage estimates, FALSE otherwise.
<code>retryFits</code>	Non-negative integer indicating the number of times to attempt to iteratively fit the model using random perturbations of the current parameter estimates as the initial values for likelihood optimization. See fitHMM .
<code>retrySD</code>	An optional list of scalars or vectors indicating the standard deviation to use for normal perturbations of each working scale parameter when <code>retryFits>0</code> . See fitHMM .
<code>optMethod</code>	The optimization method to be used. See fitHMM .
<code>control</code>	A list of control parameters to be passed to <code>optim</code> (ignored unless <code>optMethod="Nelder-Mead"</code> or <code>optMethod="SANN"</code>).
<code>modelName</code>	An optional character string providing a name for the fitted model. See fitHMM .
<code>...</code>	further arguments passed to or from other methods. Not currently used.

Value

A `randomEffects` model similar to a [momentuHMM](#) object, but including the additional random effect components:

<code>varcomp</code>	A list of length <code>nbStates*(nbStates-1)</code> with each element containing the random effect mean coefficient(s) (<code>mu</code>), random effect variance (<code>sigma</code>), and logit-scale shrinkage estimates for the state transition probability parameters (<code>ztilde</code>).
----------------------	---

traceG The trace of the projection matrix for each random effect.

References

- Burnham, K.P. and White, G.C. 2002. Evaluation of some random effects methodology applicable to bird ringing data. *Journal of Applied Statistics* 29: 245-264.
- McClintock, B.T. 2021. Worth the effort? A practical examination of random effects in hidden Markov models for animal telemetry data. *Methods in Ecology and Evolution* doi:10.1111/2041-210X.13619.

Examples

```
## Not run:
# simulated data with normal random effects
# and binary individual covariate

nbAnimals <- 5 # should be larger for random effects estimation
obsPerAnimal <- 110
indCov <- rbinom(nbAnimals,1,0.5) # individual covariate
betaCov <- c(-0.5,0.5) # covariate effects
mu <- c(-0.1,0.1) # mean for random effects
sigma <- c(0.2,0.4) # sigma for random effects
beta0 <- cbind(rnorm(nbAnimals,mu[1],sigma[1]),
               rnorm(nbAnimals,mu[2],sigma[2]))

reData <- simData(nbAnimals=nbAnimals,obsPerAnimal=obsPerAnimal,nbStates=2,
                  dist=list(step="gamma"),formula=~0+ID+indCov,
                  Par=list(step=c(1,10,1,2)),
                  beta=rbind(beta0,betaCov),
                  covs=data.frame(indCov=rep(indCov,each=obsPerAnimal)))

# fit null model
nullFit <- fitHMM(reData,nbStates=2,
                     dist=list(step="gamma"),
                     Par0=list(step=c(1,10,1,2)))

# fit covariate model
covFit <- fitHMM(reData,nbStates=2,
                   dist=list(step="gamma"),formula=~indCov,
                   Par0=list(step=c(1,10,1,2)),
                   beta0=rbind(mu,betaCov))

# fit fixed effects model
fixFit <- fitHMM(reData,nbStates=2,
                  dist=list(step="gamma"),formula=~0+ID,
                  Par0=list(step=c(1,10,1,2)),
                  beta0=beta0)

# fit random effect model
reFit <- randomEffects(fixFit)

# fit random effect model with individual covariate
```

```
reCovFit <- randomEffects(fixFit, Xformula=~indCov)

# compare by AICc
AIC(nullFit,covFit,fixFit,reFit,reCovFit, n=nrow(reData))

## End(Not run)
```

`setModelName`*Set modelName for a momentuHMM, miHMM, HMMfits, or miSum object***Description**

Set modelName for a momentuHMM, miHMM, HMMfits, or miSum object

Usage

```
setModelName(model, modelName)
```

Arguments

<code>model</code>	<code>momentuHMM</code> , <code>miHMM</code> , <code>HMMfits</code> , or <code>miSum</code> object
<code>modelName</code>	Character string providing a name for the model. See fitHMM and MIfitHMM .

Value

model object with new modelName field

Examples

```
m <- example$m
mName <- setModelName(m, modelName="example")
```

`setStateNames`*Set stateNames for a momentuHMM, miHMM, HMMfits, or miSum object***Description**

Set stateNames for a momentuHMM, miHMM, HMMfits, or miSum object

Usage

```
setStateNames(model, stateNames)
```

Arguments

model	<code>momentuHMM</code> , <code>miHMM</code> , <code>HMMfits</code> , or <code>miSum</code> object
stateNames	Character string providing state names for the model. See <code>fitHMM</code> and <code>MIfitHMM</code> .

Value

`model` object with new `stateNames` field

Examples

```
m <- example$m
mName <- setStateNames(m, stateNames=c("encamped","exploratory"))
```

simData

Simulation tool

Description

Simulates data from a (multivariate) hidden Markov model. Movement data are assumed to be in Cartesian coordinates (not longitude/latitude) and can be generated with or without observation error attributable to temporal irregularity or location measurement error.

Usage

```
simData(
  nbAnimals = 1,
  nbStates = 2,
  dist,
  Par,
  beta = NULL,
  delta = NULL,
  formula = ~1,
  formulaDelta = NULL,
  mixtures = 1,
  formulaPi = NULL,
  covs = NULL,
  nbCovs = 0,
  spatialCovs = NULL,
  zeroInflation = NULL,
  oneInflation = NULL,
  circularAngleMean = NULL,
  centers = NULL,
  centroids = NULL,
  angleCovs = NULL,
  obsPerAnimal = c(500, 1500),
  initialPosition = c(0, 0),
```

```
DM = NULL,
userBounds = NULL,
workBounds = NULL,
betaRef = NULL,
mvnCoords = NULL,
stateNames = NULL,
model = NULL,
states = FALSE,
retrySims = 0,
lambda = NULL,
errorEllipse = NULL,
ncores = 1
)

simHierData(
  nbAnimals = 1,
  hierStates,
  hierDist,
  Par,
  hierBeta = NULL,
  hierDelta = NULL,
  hierFormula = NULL,
  hierFormulaDelta = NULL,
  mixtures = 1,
  formulaPi = NULL,
  covs = NULL,
  nbHierCovs = NULL,
  spatialCovs = NULL,
  zeroInflation = NULL,
  oneInflation = NULL,
  circularAngleMean = NULL,
  centers = NULL,
  centroids = NULL,
  angleCovs = NULL,
  obsPerLevel,
  initialPosition = c(0, 0),
  DM = NULL,
  userBounds = NULL,
  workBounds = NULL,
  mvnCoords = NULL,
  model = NULL,
  states = FALSE,
  retrySims = 0,
  lambda = NULL,
  errorEllipse = NULL,
  ncores = 1
)
```

Arguments

nbAnimals	Number of observed individuals to simulate.
nbStates	Number of behavioural states to simulate.
dist	A named list indicating the probability distributions of the data streams. Currently supported distributions are 'bern', 'beta', 'cat', 'exp', 'gamma', 'lnorm', 'logis', 'negbinom', 'norm', 'mvnrm2' (bivariate normal distribution), 'mvnrm3' (trivariate normal distribution), 'pois', 'rw_norm' (normal random walk), 'rw_mvnorm2' (bivariate normal random walk), 'rw_mvnorm3' (trivariate normal random walk), 'vm', 'vmConsensus', 'weibull', and 'wrpcalpha'. For example, <code>dist=list(step='gamma', angle='vm', dives='pois')</code> indicates 3 data streams ('step', 'angle', and 'dives') and their respective probability distributions ('gamma', 'vm', and 'pois').
Par	A named list containing vectors of initial state-dependent probability distribution parameters for each data stream specified in <code>dist</code> . The parameters should be in the order expected by the pdfs of <code>dist</code> , and any zero-mass and/or one-mass parameters should be the last (if both are present, then zero-mass parameters must precede one-mass parameters). If <code>DM</code> is not specified for a given data stream, then <code>Par</code> is on the natural (i.e., real) scale of the parameters. However, if <code>DM</code> is specified for a given data stream, then <code>Par</code> must be on the working (i.e., beta) scale of the parameters, and the length of <code>Par</code> must match the number of columns in the design matrix. See details below.
beta	Matrix of regression parameters for the transition probabilities (more information in "Details").
delta	Initial value for the initial distribution of the HMM. Default: <code>rep(1/nbStates, nbStates)</code> . If <code>formulaDelta</code> includes a formula, then <code>delta</code> must be specified as a $k \times (nbStates-1)$ matrix, where k is the number of covariates and the columns correspond to states $2:nbStates$. See details below.
formula	Regression formula for the transition probability covariates. Default: <code>~1</code> (no covariate effect). In addition to allowing standard functions in R formulas (e.g., <code>cos(cov)</code> , <code>cov1*cov2</code> , <code>I(cov^2)</code>), special functions include <code>cosinor(cov, period)</code> for modeling cyclical patterns, spline functions (<code>bs</code> , <code>ns</code> , <code>bSpline</code> , <code>cSpline</code> , <code>iSpline</code> , and <code>mSpline</code>), and state- or parameter-specific formulas (see details). Any formula terms that are not state- or parameter-specific are included on all of the transition probabilities.
formulaDelta	Regression formula for the initial distribution. Default: <code>NULL</code> (no covariate effects and <code>delta</code> is specified on the real scale). Standard functions in R formulas are allowed (e.g., <code>cos(cov)</code> , <code>cov1*cov2</code> , <code>I(cov^2)</code>). When any formula is provided, then <code>delta</code> must be specified on the working scale.
mixtures	Number of mixtures for the state transition probabilities (i.e. discrete random effects *sensu* DeRuiter et al. 2017). Default: <code>mixtures=1</code> .
formulaPi	Regression formula for the mixture distribution probabilities. Default: <code>NULL</code> (no covariate effects; both <code>beta\$pi</code> and <code>fixPar\$pi</code> are specified on the real scale). Standard functions in R formulas are allowed (e.g., <code>cos(cov)</code> , <code>cov1*cov2</code> , <code>I(cov^2)</code>). When any formula is provided, then both <code>beta\$pi</code> and <code>fixPar\$pi</code> are specified on the working scale. Note that only the covariate values corresponding to the first time step for each individual ID are used (i.e. time-varying covariates cannot be used for the mixture probabilities).

<code>covs</code>	Covariate values to include in the simulated data, as a data frame. The names of any covariates specified by <code>covs</code> can be included in <code>formula</code> and/or <code>DM</code> . Covariates can also be simulated according to a standard normal distribution, by setting <code>covs</code> to <code>NULL</code> (the default), and specifying <code>nbCovs > 0</code> .
<code>nbCovs</code>	Number of covariates to simulate (0 by default). Does not need to be specified if <code>covs</code> is specified. Simulated covariates are provided generic names (e.g., 'cov1' and 'cov2' for <code>nbCovs=2</code>) and can be included in <code>formula</code> and/or <code>DM</code> .
<code>spatialCovs</code>	List of <code>raster</code> objects for spatio-temporally referenced covariates. Covariates specified by <code>spatialCovs</code> are extracted from the raster layer(s) based on any simulated location data (and the z values for a raster <code>stack</code> or <code>brick</code>) for each time step. If an element of <code>spatialCovs</code> is a raster <code>stack</code> or <code>brick</code> , then z values must be set using <code>raster::setZ</code> and <code>covs</code> must include column(s) of the corresponding z value(s) for each observation (e.g., 'time'). The names of the raster layer(s) can be included in <code>formula</code> and/or <code>DM</code> . Note that <code>simData</code> usually takes longer to generate simulated data when <code>spatialCovs</code> is specified.
<code>zeroInflation</code>	A named list of logicals indicating whether the probability distributions of the data streams should be zero-inflated. If <code>zeroInflation</code> is <code>TRUE</code> for a given data stream, then values for the zero-mass parameters should be included in the corresponding element of <code>Par</code> .
<code>oneInflation</code>	A named list of logicals indicating whether the probability distributions of the data streams should be one-inflated. If <code>oneInflation</code> is <code>TRUE</code> for a given data stream, then values for the one-mass parameters should be included in the corresponding element of <code>Par</code> .
<code>circularAngleMean</code>	An optional named list indicating whether to use circular-linear (<code>FALSE</code>) or circular-circular (<code>TRUE</code>) regression on the mean of circular distributions ('vm' and 'wrpc Cauchy') for turning angles. For example, <code>circularAngleMean=list(angle=TRUE)</code> indicates the angle mean is to be estimated for 'angle' using circular-circular regression. Whenever circular-circular regression is used for an angular data stream, a corresponding design matrix (<code>DM</code>) must be specified for the data stream, and the previous movement direction (i.e., a turning angle of zero) is automatically used as the reference angle (i.e., the intercept). Default is <code>NULL</code> , which assumes circular-linear regression is used for any angular distributions. Any <code>circularAngleMean</code> elements corresponding to data streams that do not have angular distributions are ignored. <code>circularAngleMean</code> is also ignored for any 'vmConsensus' data streams (because the consensus model is a circular-circular regression model). Alternatively, <code>circularAngleMean</code> can be specified as a numeric scalar, where the value specifies the coefficient for the reference angle (i.e., directional persistence) term in the circular-circular regression model. For example, setting <code>circularAngleMean</code> to 0 specifies a circular-circular regression model with no directional persistence term (thus specifying a biased random walk instead of a biased correlated random walk). Setting <code>circularAngleMean</code> to 1 is equivalent to setting it to <code>TRUE</code> , i.e., a circular-circular regression model with a coefficient of 1 for the directional persistence reference angle.
<code>centers</code>	2-column matrix providing the x-coordinates (column 1) and y-coordinates (column 2) for any activity centers (e.g., potential centers of attraction or repul-

	sion) from which distance and angle covariates will be calculated based on the simulated location data. These distance and angle covariates can be included in formula and DM using the row names of centers. If no row names are provided, then generic names are generated for the distance and angle covariates (e.g., 'center1.dist', 'center1.angle', 'center2.dist', 'center2.angle'); otherwise the covariate names are derived from the row names of centers as <code>paste0(rep(rownames(centers),each=2),c(".dist",".angle"))</code> . Note that the angle covariates for each activity center are calculated relative to the previous movement direction instead of standard directions relative to the x-axis; this is to allow turning angles to be simulated as a function of these covariates using circular-circular regression.
centroids	List where each element is a data frame consisting of at least <code>max(unlist(obsPerAnimal))</code> rows that provides the x-coordinates ('x') and y-coordinates ('y) for centroids (i.e., dynamic activity centers where the coordinates can change for each time step) from which distance and angle covariates will be calculated based on the simulated location data. These distance and angle covariates can be included in formula and DM using the names of centroids. If no list names are provided, then generic names are generated for the distance and angle covariates (e.g., 'centroid1.dist', 'centroid1.angle', 'centroid2.dist', 'centroid2.angle'); otherwise the covariate names are derived from the list names of centroids as <code>paste0(rep(names(centroids),ea</code> . Note that the angle covariates for each centroid are calculated relative to the previous movement direction instead of standard directions relative to the x-axis; this is to allow turning angles to be simulated as a function of these covariates using circular-circular regression.
angleCovs	Character vector indicating the names of any circular-circular regression angular covariates in <code>covs</code> or <code>spatialCovs</code> that need conversion from standard direction (in radians relative to the x-axis) to turning angle (relative to previous movement direction) using <code>circAngles</code> .
obsPerAnimal	Either the number of observations per animal (if single value) or the bounds of the number of observations per animal (if vector of two values). In the latter case, the numbers of observations generated for each animal are uniformly picked from this interval. Alternatively, <code>obsPerAnimal</code> can be specified as a list of length <code>nbAnimals</code> with each element providing the number of observations (if single value) or the bounds (if vector of two values) for each individual. Default: <code>c(500, 1500)</code> .
initialPosition	2-vector providing the x- and y-coordinates of the initial position for all animals. Alternatively, <code>initialPosition</code> can be specified as a list of length <code>nbAnimals</code> with each element a 2-vector providing the x- and y-coordinates of the initial position for each individual. Default: <code>c(0, 0)</code> . If <code>mvnCoord</code> corresponds to a data stream with "mvnorm3" or "rw_mvnorm3" probability distributions, then <code>initialPosition</code> must be composed of 3-vector(s) for the x-, y-, and z-coordinates.
DM	An optional named list indicating the design matrices to be used for the probability distribution parameters of each data stream. Each element of <code>DM</code> can either be a named list of regression formulas or a "pseudo" design matrix. For example, for a 2-state model using the gamma distribution for a data stream named 'step',

`DM=list(step=list(mean=~cov1, sd=~1))` specifies the mean parameters as a function of the covariate 'cov1' for each state. This model could equivalently be specified as a 4x6 "pseudo" design matrix using character strings for the covariate: `DM=list(step=matrix(c(1,0,0,0, 'cov1', 0,0,0,0,1,0,0,0, 'cov1', 0,0,0,0,1,0,0,0,0,` where the 4 rows correspond to the state-dependent parameters (`mean_1,mean_2,sd_1,sd_2`) and the 6 columns correspond to the regression coefficients.

Design matrices specified using formulas allow standard functions in R formulas (e.g., `cos(cov)`, `cov1*cov2`, `I(cov^2)`). Special formula functions include `cosinor(cov, period)` for modeling cyclical patterns, spline functions (`bs`, `ns`, `bSpline`, `cSpline`, `iSpline`, and `mSpline`), `angleFormula(cov, strength, by)` for the angle mean of circular-circular regression models, and state-specific formulas (see details). Any formula terms that are not state-specific are included on the parameters for all `nbStates` states.

<code>userBounds</code>	An optional named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream. For example, for a 2-state model using the wrapped Cauchy ('wrpcauchy') distribution for a data stream named 'angle' with <code>estAngleMean\$angle=TRUE</code> , <code>userBounds=list(angle=matrix(c(-pi,-pi,-1,-1,pi,pi,1,1),4,2,dimnames=list(c("mean_1"</code> specifies (-1,1) bounds for the concentration parameters instead of the default [0,1) bounds.
<code>workBounds</code>	An optional named list of 2-column matrices specifying bounds on the working scale of the probability distribution, transition probability, and initial distribution parameters. For each matrix, the first column pertains to the lower bound and the second column the upper bound. For data streams, each element of <code>workBounds</code> should be a $k \times 2$ matrix with the same name of the corresponding element of <code>Par</code> , where k is the number of parameters. For transition probability parameters, the corresponding element of <code>workBounds</code> must be a $k \times 2$ matrix named "beta", where $k=\text{length}(\text{beta})$. For initial distribution parameters, the corresponding element of <code>workBounds</code> must be a $k \times 2$ matrix named "delta", where $k=\text{length}(\text{delta})$. <code>workBounds</code> is ignored for any given data stream unless <code>DM</code> is also specified.
<code>betaRef</code>	Numeric vector of length <code>nbStates</code> indicating the reference elements for the t.p.m. multinomial logit link. Default: NULL, in which case the diagonal elements of the t.p.m. are the reference. See fitHMM .
<code>mvnCoords</code>	Character string indicating the name of location data that are to be simulated using a multivariate normal distribution. For example, if <code>mu="rw_mvnorm2"</code> was included in <code>dist</code> and <code>(mu.x, mu.y)</code> are intended to be location data, then <code>mvnCoords="mu"</code> needs to be specified in order for these data to be treated as such.
<code>stateNames</code>	Optional character vector of length <code>nbStates</code> indicating state names.
<code>model</code>	A <code>momentuHMM</code> , <code>momentuHierHMM</code> , <code>miHMM</code> , or <code>miSum</code> object. This option can be used to simulate from a fitted model. Default: NULL. Note that, if this argument is specified, most other arguments will be ignored – except for <code>nbAnimals</code> , <code>obsPerAnimal</code> , <code>states</code> , <code>initialPosition</code> , <code>lambda</code> , <code>errorEllipse</code> , and, if covariate values different from those in the data should be specified, <code>covs</code> , <code>spatialCovs</code> , <code>centers</code> , and <code>centroids</code> . It is not appropriate to simulate move-

	ment data from a model that was fitted to latitude/longitude data (because <code>simData</code> assumes Cartesian coordinates).
<code>states</code>	TRUE if the simulated states should be returned, FALSE otherwise (default).
<code>retrySims</code>	Number of times to attempt to simulate data within the spatial extent of <code>spatialCovs</code> . If <code>retrySims=0</code> (the default), an error is returned if the simulated tracks(s) move beyond the extent(s) of the raster layer(s). Instead of relying on <code>retrySims</code> , in many cases it might be better to simply expand the extent of the raster layer(s) and/or adjust the step length and turning angle probability distributions. Ignored if <code>spatialCovs=NULL</code> .
<code>lambda</code>	Observation rate for location data. If NULL (the default), location data are obtained at regular intervals. Otherwise <code>lambda</code> is the rate parameter of the exponential distribution for the waiting times between successive location observations, i.e., $1/\lambda$ is the expected time between successive location observations. Only the 'step' and 'angle' data streams are subject to temporal irregularity; any other data streams are observed at temporally-regular intervals. Ignored unless a valid distribution for the 'step' data stream is specified.
<code>errorEllipse</code>	List providing the upper bound for the semi-major axis (<code>M</code> ; on scale of x- and y-coordinates), semi-minor axis (<code>m</code> ; on scale of x- and y-coordinates), and orientation (<code>r</code> ; in degrees) of location error ellipses. If NULL (the default), no location measurement error is simulated. If <code>errorEllipse</code> is specified, then each observed location is subject to bivariate normal errors as described in McClintock et al. (2015), where the components of the error ellipse for each location are randomly drawn from <code>runif(1,min(errorEllipse\$M),max(errorEllipse\$M))</code> , <code>runif(1,min(errorEllipse\$m),max(errorEllipse\$m))</code> , and <code>runif(1,min(errorEllipse\$r),max(errorEllipse\$r))</code> . If only a single value is provided for any of the error ellipse elements, then the corresponding component is fixed to this value for each location. Only the 'step' and 'angle' data streams are subject to location measurement error; any other data streams are observed without error. Ignored unless a valid distribution for the 'step' data stream is specified.
<code>ncores</code>	Number of cores to use for parallel processing. Default: 1 (no parallel processing).
<code>hierStates</code>	A hierarchical model structure Node for the states ('state'). See details.
<code>hierDist</code>	A hierarchical data structure Node for the data streams ('dist'). Currently supported distributions are 'bern', 'beta', 'exp', 'gamma', 'Inorm', 'norm', 'mvnrm2' (bivariate normal distribution), 'mvnrm3' (trivariate normal distribution), 'pois', 'rw_norm' (normal random walk), 'rw_mvnorm2' (bivariate normal random walk), 'rw_mvnorm3' (trivariate normal random walk), 'vm', 'vmConsensus', 'weibull', and 'wrpecauchy'. See details.
<code>hierBeta</code>	A hierarchical data structure Node for the matrix of initial values for the regression coefficients of the transition probabilities at each level of the hierarchy ('beta'). See fitHMM .
<code>hierDelta</code>	A hierarchical data structure Node for the matrix of initial values for the regression coefficients of the initial distribution at each level of the hierarchy ('delta'). See fitHMM .
<code>hierFormula</code>	A hierarchical formula structure for the transition probability covariates for each level of the hierarchy ('formula'). Default: NULL (only hierarchical-level effects,

with no covariate effects). Any formula terms that are not state- or parameter-specific are included on all of the transition probabilities within a given level of the hierarchy. See details.

hierFormulaDelta

A hierarchical formula structure for the initial distribution covariates for each level of the hierarchy ('formulaDelta'). Default: NULL (no covariate effects and fixPar\$delta is specified on the working scale).

nbHierCovs

A hierarchical data structure [Node](#) for the number of covariates ('nbCovs') to simulate for each level of the hierarchy (0 by default). Does not need to be specified if covs is specified. Simulated covariates are provided generic names (e.g., 'cov1.1' and 'cov1.2' for nbHierCovs\$level1\$nbCovs=2) and can be included in hierFormula and/or DM.

obsPerLevel

A hierarchical data structure [Node](#) indicating the number of observations for each level of the hierarchy ('obs'). For each level, the 'obs' field can either be the number of observations per animal (if single value) or the bounds of the number of observations per animal (if vector of two values). In the latter case, the numbers of observations generated per level for each animal are uniformly picked from this interval. Alternatively, obsPerLevel can be specified as a list of length nbAnimals with each element providing the hierarchical data structure for the number of observations for each level of the hierarchy for each animal, where the 'obs' field can either be the number of observations (if single value) or the bounds of the number of observations (if vector of two values) for each individual.

Details

- `simHierData` is very similar to `simData` except that instead of simply specifying the number of states (`nbStates`), distributions (`dist`), observations (`obsPerAnimal`), covariates (`nbCovs`), and a single t.p.m. formula (`formula`), the `hierStates` argument specifies the hierarchical nature of the states, the `hierDist` argument specifies the hierarchical nature of the data streams, the `obsPerLevel` argument specifies the number of observations for each level of the hierarchy, the `nbHierCovs` argument specifies the number of covariates for each level of the hierarchy, and the `hierFormula` argument specifies a t.p.m. formula for each level of the hierarchy. All of the hierarchical arguments in `simHierData` are specified as [Node](#) objects from the `data.tree` package.
- x- and y-coordinate location data are generated only if valid 'step' and 'angle' data streams are specified. Valid distributions for 'step' include 'gamma', 'weibull', 'exp', and 'Inorm'. Valid distributions for 'angle' include 'vm' and 'wrpcalpha'. If only a valid 'step' data stream is specified, then only x-coordinates are generated.
- If DM is specified for a particular data stream, then the initial values are specified on the working (i.e., beta) scale of the parameters. The working scale of each parameter is determined by the link function used. The function `getParDM` is intended to help with obtaining initial values on the working scale when specifying a design matrix and other parameter constraints.
- Simulated data that are temporally regular (i.e., `lambda=NULL`) and without location measurement error (i.e., `errorEllipse=NULL`) are returned as a `momentuHMMData` (or `momentuHierHMMData`) object suitable for analysis using `fitHMM`.

- Simulated location data that are temporally-irregular (i.e., `lambda>0`) and/or with location measurement error (i.e., `errorEllipse!=NULL`) are returned as a data frame suitable for analysis using `crawlWrap`.
- The matrix `beta` of regression coefficients for the transition probabilities has one row for the intercept, plus one row for each covariate, and one column for each non-diagonal element of the transition probability matrix. For example, in a 3-state HMM with 2 formula covariates, the matrix `beta` has three rows (intercept + two covariates) and six columns (six non-diagonal elements in the 3x3 transition probability matrix - filled in row-wise). In a covariate-free model (default), `beta` has one row, for the intercept.
- State-specific formulas can be specified in DM using special formula functions. These special functions can take the names `paste0("state", 1:nbStates)` (where the integer indicates the state-specific formula). For example, `DM=list(step=list(mean=~cov1+state1(cov2), sd=~cov2+state2(cov1)))` includes `cov1` on the mean parameter for all states, `cov2` on the mean parameter for state 1, `cov2` on the `sd` parameter for all states, and `cov1` on the `sd` parameter for state 2.
- State- and parameter-specific formulas can be specified for transition probabilities in `formula` using special formula functions. These special functions can take the names `paste0("state", 1:nbStates)` (where the integer indicates the current state from which transitions occur), `paste0("toState", 1:nbStates)` (where the integer indicates the state to which transitions occur), or `paste0("betaCol", nbStates*(nbStates-1))` (where the integer indicates the column of the `beta` matrix). For example with `nbStates=3`, `formula=~cov1+betaCol1(cov2)+state3(cov3)+toState1(cov4)` includes `cov1` on all transition probability parameters, `cov2` on the beta column corresponding to the transition from state 1->2, `cov3` on transition probabilities from state 3 (i.e., beta columns corresponding to state transitions 3->1 and 3->2), and `cov4` on transition probabilities to state 1 (i.e., beta columns corresponding to state transitions 2->1 and 3->1).
- Cyclical relationships (e.g., hourly, monthly) may be simulated using the `cosinor(x, period)` special formula function for covariate `x` and sine curve period of time length `period`. For example, if the data are hourly, a 24-hour cycle can be simulated using `~cosinor(cov1, 24)`, where the covariate `cov1` is a repeating series of integers $0, 1, \dots, 23, 0, 1, \dots, 23, 0, 1, \dots$ (note that `simData` will not do this for you, the appropriate covariate must be specified using the `covs` argument; see example below). The `cosinor(x, period)` function converts `x` to 2 covariates `cosinorCos(x)=cos(2*pi*x/period)` and `cosinorSin(x)=sin(2*pi*x/period)` for inclusion in the model (i.e., 2 additional parameters per state). The amplitude of the sine wave is thus `sqrt(B_cos^2 + B_sin^2)`, where `B_cos` and `B_sin` are the working parameters corresponding to `cosinorCos(x)` and `cosinorSin(x)`, respectively (e.g., see Cornelissen 2014).

When the circular-circular regression model is used, the special function `angleFormula(cov, strength, by)` can be used in DM for the mean of angular distributions (i.e. '`vm`', '`vmConsensus`', and '`wr-pcauchy`'), where `cov` is an angle covariate (e.g. wind direction), `strength` is a positive real covariate (e.g. wind speed), and `by` is an optional factor variable for individual- or group-level effects (e.g. ID, sex). This allows angle covariates to be weighted based on their strength or importance at time step `t` as in Rivest et al. (2016).

- If the length of covariate values passed (either through '`covs`', or '`model`') is not the same as the number of observations suggested by '`nbAnimals`' and '`obsPerAnimal`' (or '`obsPerLevel`' for `simHierData`), then the series of covariates is either shortened (removing last values - if too long) or extended (starting over from the first values - if too short).
- For `simData`, when covariates are not included in `formulaDelta` (i.e. `formulaDelta=NULL`), then `delta` is specified as a vector of length `nbStates` that sums to 1. When covariates are in-

cluded in `formulaDelta`, then `delta` must be specified as a $k \times (\text{nbStates}-1)$ matrix of working parameters, where k is the number of regression coefficients and the columns correspond to states $2:\text{nbStates}$. For example, in a 3-state HMM with `formulaDelta=~cov1+cov2`, the matrix `delta` has three rows (intercept + two covariates) and 2 columns (corresponding to states 2 and 3). The initial distribution working parameters are transformed to the real scale as `exp(covsDelta*Delta)/rowSums(exp(covsDelta*Delta))`, where `covsDelta` is the $N \times k$ design matrix, `Delta=cbind(rep(0,k),delta)` is a $k \times \text{nbStates}$ matrix of working parameters, and $N=\text{length}(\text{unique}(\text{data}\$ID))$.

- For `simHierData`, `delta` must be specified as a $k \times (\text{nbStates}-1)$ matrix of working parameters, where k is the number of regression coefficients and the columns correspond to states $2:\text{nbStates}$.

Value

If the simulated data are temporally regular (i.e., `lambda=NULL`) with no measurement error (i.e., `errorEllipse=NULL`), an object `momentuHMMDData` (or `momentuHierHMMDData`), i.e., a data frame of:

<code>ID</code>	The ID(s) of the observed animal(s)
<code>...</code>	Data streams as specified by <code>dist</code> (or <code>hierDist</code>)
<code>x</code>	Either easting or longitude (if data streams include valid non-negative distribution for 'step')
<code>y</code>	Either norting or latitude (if data streams include valid non-negative distribution for 'step')
<code>...</code>	Covariates (if any)

If simulated location data are temporally irregular (i.e., `lambda>0`) and/or include measurement error (i.e., `errorEllipse!=NULL`), a data frame of:

<code>time</code>	Numeric time of each observed (and missing) observation
<code>ID</code>	The ID(s) of the observed animal(s)
<code>x</code>	Either easting or longitude observed location
<code>y</code>	Either norting or latitude observed location
<code>...</code>	Data streams that are not derived from location (if applicable)
<code>...</code>	Covariates at temporally-regular true (<code>mux,muy</code>) locations (if any)
<code>mux</code>	Either easting or longitude true location
<code>muy</code>	Either norting or latitude true location
<code>error_semimajor_axis</code>	error ellipse semi-major axis (if applicable)
<code>error_semiminor_axis</code>	error ellipse semi-minor axis (if applicable)
<code>error_ellipse_orientation</code>	error ellipse orientation (if applicable)
<code>ln.sd.x</code>	log of the square root of the x-variance of bivariate normal error (if applicable; required for error ellipse models in <code>crawlWrap</code>)

ln.sd.y	log of the square root of the y-variance of bivariate normal error (if applicable; required for error ellipse models in crawlWrap)
error.corr	correlation term of bivariate normal error (if applicable; required for error ellipse models in crawlWrap)

References

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- McClintock BT, London JM, Cameron MF, Boveng PL. 2015. Modelling animal movement using the Argos satellite telemetry location error ellipse. *Methods in Ecology and Evolution* 6(3):266-277.
- Rivest, LP, Duchesne, T, Nicosia, A, Fortin, D, 2016. A general angular regression model for the analysis of data on animal movement in ecology. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, 65(3):445-463.
- Leos-Barajas, V., Gangloff, E.J., Adam, T., Langrock, R., van Beest, F.M., Nabe-Nielsen, J. and Morales, J.M. 2017. Multi-scale modeling of animal movement and general behavior data using hidden Markov models with hierarchical structures. *Journal of Agricultural, Biological and Environmental Statistics*, 22 (3), 232-248.

See Also

[prepData](#), [simObsData](#)

Examples

```
# 1. Pass a fitted model to simulate from
# (m is a momentuHMM object - as returned by fitHMM - automatically loaded with the package)
# We keep the default nbAnimals=1.
m <- example$m
obsPerAnimal=c(50,100)
data <- simData(model=m,obsPerAnimal=obsPerAnimal)

## Not run:
# 2. Pass the parameters of the model to simulate from
stepPar <- c(1,10,1,5,0.2,0.3) # mean_1, mean_2, sd_1, sd_2, zeromass_1, zeromass_2
anglePar <- c(pi,0,0.5,2) # mean_1, mean_2, concentration_1, concentration_2
omegaPar <- c(1,10,10,1) # shape1_1, shape1_2, shape2_1, shape2_2
stepDist <- "gamma"
angleDist <- "vm"
omegaDist <- "beta"
data <- simData(nbAnimals=4,nbStates=2,dist=list(step=stepDist,angle=angleDist,omega=omegaDist),
               Par=list(step=stepPar,angle=anglePar,omega=omegaPar),nbCovs=2,
               zeroInflation=list(step=TRUE),
               obsPerAnimal=obsPerAnimal)

# 3. Include covariates
# (note that it is useless to specify "nbCovs", which are overruled
# by the number of columns of "cov")
cov <- data.frame(temp=log(rnorm(500,20,5)))
stepPar <- c(log(10),0.1,log(100),-0.1,log(5),log(25)) # working scale parameters for step DM
```

```

anglePar <- c(pi,0,0.5,2) # mean_1, mean_2, concentration_1, concentration_2
stepDist <- "gamma"
angleDist <- "vm"
data <- simData(nbAnimals=2,nbStates=2,dist=list(step=stepDist,angle=angleDist),
                Par=list(step=stepPar,angle=anglePar),
                DM=list(step=list(mean=~temp,sd=~1)),
                covs=cov,
                obsPerAnimal=obsPerAnimal)

# 4. Include example 'forest' spatial covariate raster layer
# nbAnimals and obsPerAnimal kept small to reduce example run time
spatialCov<-list(forest=forest)
data <- simData(nbAnimals=1,nbStates=2,dist=list(step=stepDist,angle=angleDist),
                Par=list(step=c(100,1000,50,100),angle=c(0,0,0.1,5)),
                beta=matrix(c(5,-10,-25,50),nrow=2,ncol=2,byrow=TRUE),
                formula=~forest,spatialCovs=spatialCov,
                obsPerAnimal=250,states=TRUE,
                retrySims=100)

# 5. Specify design matrix for 'omega' data stream
# natural scale parameters for step and angle
stepPar <- c(1,10,1,5) # shape_1, shape_2, scale_1, scale_2
anglePar <- c(pi,0,0.5,0.7) # mean_1, mean_2, concentration_1, concentration_2

# working scale parameters for omega DM
omegaPar <- c(log(1),0.1,log(10),-0.1,log(10),-0.1,log(1),0.1)

stepDist <- "weibull"
angleDist <- "wrcpcauchy"
omegaDist <- "beta"

data <- simData(nbStates=2,dist=list(step=stepDist,angle=angleDist,omega=omegaDist),
                Par=list(step=stepPar,angle=anglePar,omega=omegaPar),nbCovs=2,
                DM=list(omega=list(shape1=~cov1,shape2=~cov2)),
                obsPerAnimal=obsPerAnimal,states=TRUE)

# 6. Include temporal irregularity and location measurement error
lambda <- 2 # expect 2 observations per time step
errorEllipse <- list(M=50,m=25,r=180)
obsData <- simData(model=m,obsPerAnimal=obsPerAnimal,
                    lambda=lambda, errorEllipse=errorEllipse)

# 7. Cosinor and state-dependent formulas
nbStates<-2
dist<-list(step="gamma")
Par<-list(step=c(100,1000,50,100))

# include 24-hour cycle on all transition probabilities
# include 12-hour cycle on transitions from state 2
formula=~cosinor(hour24,24)+state2(cosinor(hour12,12))

# specify appropriate covariates
covs<-data.frame(hour24=0:23,hour12=0:11)

```

```

beta<-matrix(c(-1.5,1,1,NA,NA,-1.5,-1,-1,1,1),5,2)
# row names for beta not required but can be helpful
rownames(beta)<-c("Intercept",
                   "cosinorCos(hour24, 24)",
                   "cosinorSin(hour24, 24)",
                   "cosinorCos(hour12, 12)",
                   "cosinorSin(hour12, 12)")
data.cos<-simData(nbStates=nbStates,dist=dist,Par=Par,
                  beta=beta,formula=formula,covs=covs)

# 8. Piecewise constant B-spline on step length mean and angle concentration
nObs <- 1000 # length of simulated track
cov <- data.frame(time=1:nObs) # time covariate for splines
dist <- list(step="gamma",angle="vm")
stepDM <- list(mean=~splines2::bSpline(time,df=2,degree=0),sd=~1)
angleDM <- list(mean=~1,concentration=~splines2::bSpline(time,df=2,degree=0))
DM <- list(step=stepDM,angle=angleDM)
Par <- list(step=c(log(1000),1,-1,log(100)),angle=c(0,log(10),2,-5))

data.spline<-simData(obsPerAnimal=nObs,nbStates=1,dist=dist,Par=Par,DM=DM,covs=cov)

# 9. Initial state (delta) based on covariate
nObs <- 100
dist <- list(step="gamma",angle="vm")
Par <- list(step=c(100,1000,50,100),angle=c(0,0,0.01,0.75))

# create sex covariate
cov <- data.frame(sex=factor(rep(c("F","M"),each=nObs))) # sex covariate
formulaDelta <- ~ sex + 0

# Female begins in state 1, male begins in state 2
delta <- matrix(c(-100,100),2,1,dimnames=list(c("sexF","sexM"),"state 2"))

data.delta<-simData(nbAnimals=2,obsPerAnimal=nObs,nbStates=2,dist=dist,Par=Par,
                     delta=delta,formulaDelta=formulaDelta,covs=cov,
                     beta=matrix(-1.5,1,2),states=TRUE)

## End(Not run)

```

Description

Simulates observed location data subject to temporal irregularity and/or location measurement error

Usage

```
simObsData(data, lambda, errorEllipse, ...)
```

```
## S3 method for class 'momentuHMMData'
simObsData(data, lambda, errorEllipse, ...)

## S3 method for class 'momentuHierHMMData'
simObsData(data, lambda, errorEllipse, coordLevel, ...)
```

Arguments

<code>data</code>	A <code>momentuHMMData</code> or <code>momentuHierHMMData</code> object with necessary fields 'x' (easting/longitudinal coordinates) and 'y' (northing/latitudinal coordinates)
<code>lambda</code>	Observation rate for location data. If NULL, location data are kept at temporally-regular intervals. Otherwise <code>lambda</code> is the rate parameter of the exponential distribution for the waiting times between successive location observations, i.e., $1/\lambda$ is the expected time between successive location observations. Only the 'step' and 'angle' data streams (or multivariate normal data streams identified by <code>mvnCoords</code>) are subject to temporal irregularity; any other data streams are kept at temporally-regular intervals. Ignored unless a valid distribution for the 'step' (or 'mvnCoord') data stream has been specified.
<code>errorEllipse</code>	List providing the bounds for the semi-major axis (<code>M</code> ; on scale of x- and y-coordinates), semi-minor axis (<code>m</code> ; on scale of x- and y-coordinates), and orientation (<code>r</code> ; in degrees) of location error ellipses. If NULL, no location measurement error is simulated. If <code>errorEllipse</code> is specified, then each observed location is subject to bivariate normal errors as described in McClintock et al. (2015), where the components of the error ellipse for each location are randomly drawn from <code>runif(1,min(errorEllipse\$M),max(errorEllipse\$M))</code> , <code>runif(1,min(errorEllipse\$m),max(errorEllipse\$m))</code> , and <code>runif(1,min(errorEllipse\$r),max(errorEllipse\$r))</code> . If only a single value is provided for any of the error ellipse elements, then the corresponding component is fixed to this value for each location. Only the 'step' and 'angle' data streams are subject to location measurement error; any other data streams are observed without error. Ignored unless a valid distribution for the 'step' data stream is specified.
<code>...</code>	further arguments passed to or from other methods
<code>coordLevel</code>	Level of the hierarchy in which the location data are obtained

Details

Simulated location data that are temporally-irregular (i.e., $\lambda > 0$) and/or with location measurement error (i.e., `errorEllipse`!=NULL) are returned as a data frame suitable for analysis using `crawlWrap`.

Value

A dataframe of:

<code>time</code>	Numeric time of each observed (and missing) observation
<code>ID</code>	The ID(s) of the observed animal(s)
<code>x</code>	Either easting or longitude observed location

y	Either norting or latitude observed location
...	Data streams that are not derived from location (if applicable)
...	Covariates at temporally-regular true (<code>mux,muy</code>) locations (if any)
<code>mux</code>	Either easting or longitude true location
<code>muy</code>	Either norting or latitude true location
<code>error_semimajor_axis</code>	error ellipse semi-major axis (if applicable)
<code>error_seminor_axis</code>	error ellipse semi-minor axis (if applicable)
<code>error_ellipse_orientation</code>	error ellipse orientation (if applicable)
<code>ln.sd.x</code>	log of the square root of the x-variance of bivariate normal error (if applicable; required for error ellipse models in <code>crawlWrap</code>)
<code>ln.sd.y</code>	log of the square root of the y-variance of bivariate normal error (if applicable; required for error ellipse models in <code>crawlWrap</code>)
<code>error.corr</code>	correlation term of bivariate normal error (if applicable; required for error ellipse models in <code>crawlWrap</code>)

References

McClintock BT, London JM, Cameron MF, Boveng PL. 2015. Modelling animal movement using the Argos satellite telemetry location error ellipse. Methods in Ecology and Evolution 6(3):266-277.

See Also

[crawlWrap](#), [prepData](#), [simData](#)
[simHierData](#)

Examples

```
# extract momentuHMMData example
data <- example$m$data
lambda <- 2 # expect 2 observations per time step
errorEllipse <- list(M=c(0,50),m=c(0,50),r=c(0,180))
obsData1 <- simObsData(data,lambda=lambda,errorEllipse=errorEllipse)

errorEllipse <- list(M=50,m=50,r=180)
obsData2 <- simObsData(data,lambda=lambda,errorEllipse=errorEllipse)
```

stateProbs

*State probabilities***Description**

For a given model, computes the probability of the process being in the different states at each time point.

Usage

```
stateProbs(m, hierarchical = FALSE)
```

Arguments

- | | |
|---------------------------|--|
| <code>m</code> | A <code>momentuHMM</code> or <code>momentuHierHMM</code> object. |
| <code>hierarchical</code> | Logical indicating whether or not to return a list of state probabilities for each level of a hierarchical HMM. Ignored unless <code>m</code> is a <code>momentuHierHMM</code> object. |

Value

The matrix of state probabilities, with element [i,j] the probability of being in state j in observation i.

References

Zucchini, W. and MacDonald, I.L. 2009. Hidden Markov Models for Time Series: An Introduction Using R. Chapman & Hall (London).

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

sp <- stateProbs(m)
```

stationary

*Stationary state probabilities***Description**

Calculates the stationary probabilities of each state based on covariate values.

Usage

```
stationary(model, covs, covIndex)
```

Arguments

model	<code>momentuHMM</code> , <code>miHMM</code> , or <code>miSum</code> object
covs	Either a data frame or a design matrix of covariates. If <code>covs</code> is not provided, then the stationary probabilities are calculated based on the covariate data for each time step.
covIndex	Integer vector indicating specific rows of the data to be used in the calculations. This can be useful for reducing unnecessarily long computation times, e.g., when <code>formula</code> includes factor covariates (such as ID) but no temporal covariates. Ignored unless <code>covs</code> is missing.

Value

A list of length `model$conditions$mixtures` where each element is a matrix of stationary state probabilities for each mixture. For each matrix, each row corresponds to a row of `covs`, and each column corresponds to a state.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

# data frame of covariates
stationary(m, covs = data.frame(cov1 = 0, cov2 = 0))

# design matrix (each column corresponds to row of m$mle$beta)
stationary(m, covs = matrix(c(1,0,cos(0)),1,3))

# get stationary distribution for first 3 observations
stationary(m, covIndex = c(1,2,3))
```

```
summary.momentuHMMData
Summary momentuHMMData
```

Description

Summary `momentuHMMData`

Usage

```
## S3 method for class 'momentuHMMData'
summary(object, dataNames = c("step", "angle"), animals = NULL, ...)

## S3 method for class 'momentuHierHMMData'
summary(object, dataNames = c("step", "angle", "level"), animals = NULL, ...)
```

Arguments

object	A <code>momentuHMMData</code> or <code>momentuHierHMMData</code> object.
dataNames	Names of the variables to summarize. Default is <code>dataNames=c("step", "angle")</code> .
animals	Vector of indices or IDs of animals for which data will be summarized. Default: <code>NULL</code> ; data for all animals are summarized.
...	Currently unused. For compatibility with generic method.

Examples

```
# data is a momentuHMMData object (as returned by prepData), automatically loaded with the package
data <- example$m$data

summary(data,dataNames=c("step","angle","cov1","cov2"))
```

timeInStates

Calculate proportion of time steps assigned to each state (i.e. “activity budgets”)

Description

Calculate proportion of time steps assigned to each state (i.e. “activity budgets”)

Usage

```
timeInStates(m, by = NULL, alpha = 0.95, ncores = 1)

## S3 method for class 'momentuHMM'
timeInStates(m, by = NULL, alpha = 0.95, ncores = 1)

## S3 method for class 'HMMfits'
timeInStates(m, by = NULL, alpha = 0.95, ncores = 1)

## S3 method for class 'miHMM'
timeInStates(m, by = NULL, alpha = 0.95, ncores = 1)
```

Arguments

m	A <code>momentuHMM</code> , <code>miHMM</code> , or <code>HMMfits</code> object.
by	A character vector indicating any groupings by which to calculate the proportions, such as individual (“ID”) or group-level (e.g. sex or age class) covariates. Default is <code>NULL</code> (no groupings are used).
alpha	Significance level for calculating confidence intervals of pooled estimates. Default: 0.95. Ignored unless m is a <code>miHMM</code> or <code>HMMfits</code> object.
ncores	Number of cores to use for parallel processing. Default: 1 (no parallel processing). Ignored unless m is a <code>miHMM</code> or <code>HMMfits</code> object.

Value

If m is a `momentuHMM` object, a data frame containing the estimated activity budgets for each state (grouped according to by). If m is a `miHMM` or `HMMfits` object, a list containing the activity budget estimates, standard errors, lower bounds, and upper bounds across all imputations.

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m
timeInStates(m)
timeInStates(m, by = "ID")
```

trMatrix_rcpp	<i>Transition probability matrix</i>
---------------	--------------------------------------

Description

Computation of the transition probability matrix, as a function of the covariates and the regression parameters. Written in C++. Used in `viterbi`.

Usage

```
trMatrix_rcpp(nbStates, beta, covs, betaRef)
```

Arguments

nbStates	Number of states
beta	Matrix of regression parameters
covs	Matrix of covariate values
betaRef	Indices of reference elements for t.p.m. multinomial logit link.

Value

Three dimensional array `trMat`, such that `trMat[, , t]` is the transition matrix at time t .

turnAngle*Turning angle***Description**

Used in [prepData](#) and [simData](#).

Usage

```
turnAngle(x, y, z, type = "UTM", angleCov = FALSE)
```

Arguments

x	First point
y	Second point
z	Third point
type	'UTM' if easting/northing provided (the default), 'LL' if longitude/latitude. If type='LL' then the geosphere package must be installed.
angleCov	logical indicating to not return NA when x=y or y=z. Default: FALSE (i.e. NA is returned if x=y or y=z).

Value

The angle between vectors (x,y) and (y,z).

If type='LL' then turning angle is calculated based on initial bearings using [bearing](#).

Examples

```
## Not run:
x <- c(0,0)
y <- c(4,6)
z <- c(10,7)
momentuHMM:::turnAngle(x,y,z)

## End(Not run)
```

viterbi*Viterbi algorithm*

Description

For a given model, reconstructs the most probable states sequence, using the Viterbi algorithm.

Usage

```
viterbi(m, hierarchical = FALSE)
```

Arguments

- m An object `momentuHMM` or `momentuHierHMM`
- hierarchical Logical indicating whether or not to return a list of Viterbi-decoded states for each level of a hierarchical HMM. Ignored unless m is a `momentuHierHMM` object.

Value

The sequence of most probable states. If `hierarchical` is TRUE, then a list of the most probable states for each level of the hierarchy is returned.

References

Zucchini, W. and MacDonald, I.L. 2009. Hidden Markov Models for Time Series: An Introduction Using R. Chapman & Hall (London).

Examples

```
# m is a momentuHMM object (as returned by fitHMM), automatically loaded with the package
m <- example$m

# reconstruction of states sequence
states <- viterbi(m)
```

w2n

Scaling function: working to natural parameters

Description

Scales each parameter from the set of real numbers, back to its natural interval. Used during the optimization of the log-likelihood.

Usage

```
w2n(
  wpar,
  bounds,
  parSize,
  nbStates,
  nbCovs,
  estAngleMean,
  circularAngleMean,
  consensus,
  stationary,
  fullDM,
  DMind,
  nbObs,
  dist,
  Bndind,
  nc,
  meanind,
  covsDelta,
  workBounds,
  covsPi
)
```

Arguments

wpar	Vector of working parameters.
bounds	Named list of 2-column matrices specifying bounds on the natural (i.e, real) scale of the probability distribution parameters for each data stream.
parSize	Named list indicating the number of natural parameters of the data stream probability distributions
nbStates	The number of states of the HMM.
nbCovs	The number of beta covariates.
estAngleMean	Named list indicating whether or not to estimate the angle mean for data streams with angular distributions ('vm' and 'wrpcauchy').
circularAngleMean	Named list indicating whether to use circular-linear or circular-circular regression on the mean of circular distributions ('vm' and 'wrpcauchy') for turning angles. See fitHMM .
consensus	Named list indicating whether to use the circular-circular regression consensus model
stationary	FALSE if there are time-varying covariates in <code>formula</code> or any covariates in <code>formulaDelta</code> . If TRUE, the initial distribution is considered equal to the stationary distribution. Default: FALSE.
fullDM	Named list containing the full (i.e. not shorthand) design matrix for each data stream.

DMind	Named list indicating whether fullDM includes individual- and/or temporal-covariates for each data stream specifies (-1,1) bounds for the concentration parameters instead of the default [0,1) bounds.
nbObs	Number of observations in the data.
dist	Named list indicating the probability distributions of the data streams.
Bndind	Named list indicating whether DM is NULL with default parameter bounds for each data stream.
nc	indicator for zeros in fullDM
meanind	index for circular-circular regression mean angles with at least one non-zero entry in fullDM
covsDelta	data frame containing the delta model covariates
workBounds	named list of 2-column matrices specifying bounds on the working scale of the probability distribution, transition probability, and initial distribution parameters
covsPi	data frame containing the pi model covariates

Value

A list of:

...	Matrices containing the natural parameters for each data stream (e.g., 'step', 'angle', etc.)
beta	Matrix of regression coefficients of the transition probabilities
delta	Initial distribution

Examples

```
## Not run:
m<-example$m
nbStates <- 2
nbCovs <- 2
parSize <- list(step=2,angle=2)
par <- list(step=c(t(m$mle$step)),angle=c(t(m$mle$angle)))
bounds <- m$conditions$bounds
beta <- matrix(rnorm(6),ncol=2,nrow=3)
delta <- c(0.6,0.4)

#working parameters
wpar <- momentuHMM:::n2w(par,bounds,list(beta=beta),log(delta[-1]/delta[1]),nbStates,
m$conditions$estAngleMean,NULL,m$conditions$Bndind,
m$conditions$dist)

#natural parameter
p <- momentuHMM:::w2n(wpar,bounds,parSize,nbStates,nbCovs,m$conditions$estAngleMean,
m$conditions$circularAngleMean,lapply(m$conditions$dist,function(x) x=="vmConsensus"),
m$conditions$stationary,m$conditions$fullDM,
m$conditions$DMind,1,m$conditions$dist,m$conditions$Bndind,
matrix(1,nrow=length(unique(m$data$ID)),ncol=1),covsDelta=m$covsDelta,
workBounds=m$conditions$workBounds)
```

```
## End(Not run)
```

XBloop_rcpp

Get XB

Description

Loop for computation of design matrix (X) times the working scale parameters (B). Written in C++. Used in [w2n](#).

Usage

```
XBloop_rcpp(
  DM,
  Xvec,
  nb0bs,
  nr,
  nc,
  circularAngleMean,
  consensus,
  rindex,
  cindex,
  nbStates,
  refCoeff = 1
)
```

Arguments

DM	design matrix
Xvec	working parameters
nb0bs	number of observations
nr	number of rows in design matrix
nc	number of column in design matrix
circularAngleMean	indicator for whether or not circular-circular regression model
consensus	indicator for whether or not circular-circular regression consensus model
rindex	row index for design matrix
cindex	column index for design matrix
nbStates	number of states
refCoeff	intercept coefficient for circular-circular regression model

Value

XB matrix

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