Package ‘saemix’

June 27, 2023

**Title**  Stochastic Approximation Expectation Maximization (SAEM) Algorithm

**Version**  3.2

**Description**  The ‘saemix’ package implements the Stochastic Approximation EM algorithm for parameter estimation in (non)linear mixed effects models. The SAEM algorithm (i) computes the maximum likelihood estimator of the population parameters, without any approximation of the model (linearisation, quadrature approximation,...), using the Stochastic Approximation Expectation Maximization (SAEM) algorithm, (ii) provides standard errors for the maximum likelihood estimator (iii) estimates the conditional modes, the conditional means and the conditional standard deviations of the individual parameters, using the Hastings-Metropolis algorithm (see Comets et al. (2017) [doi:10.18637/jss.v080.i03]). Many applications of SAEM in agronomy, animal breeding and PKPD analysis have been published by members of the Monolix group. The full PDF documentation for the package including references about the algorithm and examples can be downloaded on the github of the IAME research institute for ‘saemix’: <https://github.com/iame-researchCenter/saemix/blob/7638e1b09cc01cdff173068e01c266e906f76eb/docsaem.pdf>.

**License**  GPL (>= 2)

**LazyLoad**  yes

**LazyData**  yes

**Imports**  graphics, stats, methods, gridExtra, ggplot2, grid, rlang, mclust, scales, MASS

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backward.procedure  
*Backward procedure for joint selection of covariates and random effects*

**Description**

Joint selection of covariates and random effects in a nonlinear mixed effects model by a backward-type algorithm based on two different versions of BIC for covariate selection and random effects selection respectively. Selection is made among the covariates as such specified in the SaemixData object. Only uncorrelated random effects structures are considered.

**Usage**

```r
backward.procedure(saemixObject, trace = TRUE)
```

**Arguments**

- `saemixObject`: An object returned by the `saemix` function
- `trace`: If TRUE, a table summarizing the steps of the algorithm is printed. Default "TRUE"

**Value**

An object of the SaemixObject class storing the covariate model and the covariance structure of random effects of the final model.

**Author(s)**

Maud Delattre

**References**


coef.saemix  
*Extract coefficients from a saemix fit*

**Description**

Extract coefficients from a saemix fit

**Usage**

```r
## S3 method for class 'SaemixObject'
coef(object, ...)
```
compare.saemix

Model comparison with information criteria (AIC, BIC).

Arguments

object  a SaemixObject

...  further arguments to be passed to or from other methods

Value

a list with 3 components:

fixed  fixed effects

population  a list of population parameters with two elements, a matrix containing the untransformed parameters psi and a matrix containing the transformed parameters phi

individual  a list of individual parameters with two elements, a matrix containing the untransformed parameters psi and a matrix containing the transformed parameters phi

Description

A specific penalty is used for BIC (BIC.cov) when the compared models have in common the structural model and the covariance structure for the random effects (see Delattre et al., 2014).

Usage

compare.saemix(..., method = c("is", "lin", "gq"))

Arguments

...  Two or more objects returned by the saemix function

method  The method used for computing the likelihood: "is" (Importance Sampling), "lin" (Linearisation) or "gq" (Gaussian quadrature). The default is to use importance sampling "is". If the requested likelihood is not available in all model objects, the method stops with a warning.

Details

Note that the comparison between two or more models will only be valid if they are fitted to the same dataset.

Value

A matrix of information criteria is returned, with at least two columns containing respectively AIC and BIC values for each of the compared models. When the models have in common the structural model and the covariance structure for the random effects, the matrix includes an additional column with BIC.cov values that are more appropriate when the comparison only concerns the covariates.
Author(s)

Maud Delattre

References


Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
name.group=c("Id"),name.predictors=c("Dose","Time"),
name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
units=list(x="hr",y="mg/L",covariates=c("kg","-"), name.X="Time")

# Definition of models to be compared
model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-(CL/V)
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}

# Model with one covariate
saemix.model1<-saemixModel(model=model1cpt,modeltype="structural",
description="One-compartment model, clearance dependent on weight",
psi0=matrix(c(1,20,0.5,0.1,0,-0.01),ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))),
transform.par=c(1,1,1),covariate.model=matrix(c(0,0,1,0,0,0),ncol=3,byrow=TRUE))

# Model with two covariates
saemix.model2<-saemixModel(model=model1cpt,modeltype="structural",
description="One-compartment model, clearance dependent on weight, volume dependent on sex",
psi0=matrix(c(1,20,0.5,0.1,0,-0.01),ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))),
transform.par=c(1,1,1),covariate.model=matrix(c(0,0,1,0,1,0),ncol=3,byrow=TRUE))

# Model with three covariates
saemix.model3<-saemixModel(model=model1cpt,modeltype="structural",
description="One-compartment model, clearance, absorption dependent on weight, volume dependent on sex",
psi0=matrix(c(1,20,0.5,0.1,0,-0.01),ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))),
transform.par=c(1,1,1),covariate.model=matrix(c(1,0,1,0,1,0),ncol=3,byrow=TRUE))

# Running the main algorithm to estimate the population parameters
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE)

saemix.fit1<-saemix(saemix.model1,saemix.data,saemix.options)

saemix.fit2<-saemix(saemix.model2,saemix.data,saemix.options)

saemix.fit3<-saemix(saemix.model3,saemix.data,saemix.options)

compare.saemix(saemix.fit1, saemix.fit2, saemix.fit3)


```r
compare.saemix(saemix.fit1, saemix.fit2, saemix.fit3, method = "lin")

# We need to explicitly run Gaussian Quadrature if we want to use it in
# the comparisons
saemix.fit1 <- llgq.saemix(saemix.fit1)
saemix.fit2 <- llgq.saemix(saemix.fit2)
saemix.fit3 <- llgq.saemix(saemix.fit3)
compare.saemix(saemix.fit1, saemix.fit2, saemix.fit3, method = "gq")
```

---

### conddist.saemix

**Estimate conditional mean and variance of individual parameters using the MCMC algorithm**

**Description**

When the parameters of the model have been estimated, we can estimate the individual parameters ($\psi_i$).

**Usage**

```r
conddist.saemix(saemixObject, nsamp = 1, max.iter = NULL, plot = FALSE, ...)
```

**Arguments**

- `saemixObject` an object returned by the `saemix` function
- `nsamp` Number of samples to be drawn in the conditional distribution for each subject. Defaults to 1
- `max.iter` Maximum number of iterations for the computation of the conditional estimates. Defaults to twice the total number of iterations (see above)
- `plot` a boolean indicating whether to display convergence plots (defaults to FALSE)
- `...` optional arguments passed to the plots. Plots will appear if the argument plot is set to TRUE

**Details**

Let $\hat{\theta}$ be the estimated value of $\theta$ computed with the SAEM algorithm and let $p(\phi_i | y_i; \hat{\theta})$ be the conditional distribution of $\phi_i$ for $1 \leq i \leq N$. We use the MCMC procedure used in the SAEM algorithm to estimate these conditional distributions. We empirically estimate the conditional mean $E(\phi_i | y_i; \hat{\theta})$ and the conditional standard deviation $sd(\phi_i | y_i; \hat{\theta})$.

See PDF documentation for details of the computation. Briefly, the MCMC algorithm is used to obtain samples from the individual conditional distributions of the parameters. The algorithm is initialised for each subject to the conditional estimate of the individual parameters obtained at the end of the SAEMIX fit. A convergence criterion is used to ensure convergence of the mean and variance of the conditional distributions. When $\text{nsamp}>1$, several chains of the MCMC algorithm
are run in parallel to obtain samples from the conditional distributions, and the convergence criterion must be achieved for all chains. When nsamp>1, the estimate of the conditional mean is obtained by averaging over the different samples, and the samples from the conditional distribution are output as an array of dimension N x nb of parameters x nsamp in arrays phi.samp for the sampled phi_i and psi.samp for the corresponding psi_i. The variance of the conditional phi_i for each sample is given in a corresponding array phi.samp.var (the variance of the conditional psi_i is not given but may be computed via the delta-method or by transforming the confidence interval for phi_i).

The shrinkage for any given parameter for the conditional estimate is obtained as
\[
Sh = 1 - \text{var}(\eta_i) / \omega(\eta)
\]
where var(\eta_i) is the empirical variance of the estimates of the individual random effects, and omega(\eta) is the estimated variance.

When the plot argument is set to TRUE, convergence graphs are produced. They can be used to assess whether the mean of the individual estimates (on the scale of the parameters) and the mean of the SD of the random effects have stabilised over the ipar.lmcmc (defaults to 50) iterations. The evolution of these variables for each parameter is shown as a continuous line while the shaded area represents the acceptable variation.

The function adds or modifies the following elements in the results:

- **cond.mean.phi** Conditional mean of the individual distribution of the parameters (obtained as the mean of the samples)
- **cond.var.phi** Conditional variance of the individual distribution of the parameters (obtained as the mean of the estimated variance of the samples)
- **cond.shrinkage** Estimate of the shrinkage for the conditional estimates
- **cond.mean.eta** Conditional mean of the individual distribution of the parameters (obtained as the mean of the samples)
- **phi.samp** An array with 3 dimensions, giving nsamp samples from the conditional distributions of the individual parameters
- **phi.samp.var** The estimated individual variances for the sampled parameters phi.samp

A warning is output if the maximum number of iterations is reached without convergence (the maximum number of iterations is the sum of the elements in saemix.options$nbiter.saemix).

**Author(s)**

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Audrey Lavenu
Marc Lavielle

**References**


See Also
SaemixData, SaemixModel, SaemixObject, saemixControl, saemix

Examples

# Not run (strict time constraints for CRAN)
data(theo.saemix)

saemix.data <- saemixData(name.data = theo.saemix, header = TRUE, sep = " ", na = NA,
                           name.group = c("Id"), name.predictors = c("Dose", "Time"),
                           name.response = c("Concentration"), name.covariates = c("Weight", "Sex"),
                           units = list(x = "hr", y = "mg/L", covariates = c("kg", "-")), name.X = "Time")

model1cpt <- function(psi, id, xidep) {
  dose <- xidep[, 1]
  tim <- xidep[, 2]
  ka <- psi[id, 1]
  V <- psi[id, 2]
  CL <- psi[id, 3]
  k <- CL/V
  ypred <- dose * ka / (V * (ka - k)) * (exp(-k * tim) - exp(-ka * tim))
  return(ypred)
}

saemix.model <- saemixModel(model = model1cpt,
                             description = "One-compartment model with first-order absorption",
                             psi0 = matrix(c(1., 20, 0.5, 0.1, 0.01), ncol = 3, byrow = TRUE,
                                           dimnames = list(NULL, c("ka", "V", "CL")), transform.par = c(1, 1, 1),
                                           covariate.model = matrix(c(0, 1, 0, 0, 0), ncol = 3, byrow = TRUE),
                                           fixed.estim = c(1, 1, 1),
                                           covariance.model = matrix(c(1, 0, 0, 0, 1, 0, 0, 1), ncol = 3, byrow = TRUE),
                                           omega.init = matrix(c(1, 0, 0, 0, 1, 0, 0, 1), ncol = 3, byrow = TRUE),
                                           error.model = "constant")
)

saemix.options <- list(seed = 632545, save = FALSE, save.graphs = FALSE)

saemix.fit <- saemix(saemix.model, saemix.data, saemix.options)

# First sample from the conditional distribution
# (a N (nb of subject) by nb.etas (nb of parameters) matrix)
print(head(saemix.fit[["results"]]["phi.samp"][, , 1]))

# Second sample
print(head(saemix.fit[["results"]]["phi.samp"][, , 2])))
Description

The cow.saemix data contains records of the weight of 560 cows on 9 or 10 occasions.

Usage

cow.saemix

Format

This data frame contains the following columns:

- **cow** the unique identifier for each cow
- **time** time (days)
- **weight** a numeric vector giving the weight of the cow (kg)
- **birthyear** year of birth (between 1988 and 1998)
- **twin** existence of a twin (no=1, yes=2)
- **birthrank** the rank of birth (between 3 and 7)

Details

An exponential model was assumed to describe the weight gain with time: \( y_{ij} = A_i \left(1 - B_i \exp(-K_i t_{ij})\right) + \epsilon_{ij} \)

References


Examples

data(cow.saemix)
saemix.data<-saemixData(name.data=cow.saemix,header=TRUE,name.group=c("cow"),
name.predictors=c("time"),name.response=c("weight"),
name.covariates=c("birthyear","twin","birthrank"),
units=list(x="days",y="kg",covariates=c("yr","-","-")))
growthcow<-function(psi,id,xidep) {
  x<-xidep[,1]
  a<-psi[id,1]
  b<-psi[id,2]
  k<-psi[id,3]
  f<-a*(1-b*exp(-k*x))
  return(f)
}
saemix.model<-saemixModel(model=growthcow,
description="Exponential growth model",
psi0=matrix(c(700,0.9,0.02,0,0,0),ncol=3,byrow=TRUE,
dinames=list(NULL,c("A","B","k"))),transform.par=c(1,1,1),fixed.estim=c(1,1,1),
covariate.model=matrix(c(0,0,0),ncol=3,byrow=TRUE),
createSaemixObject

createSaemixObject

Create saemix objects with only data filled in

Description

Create saemix objects either with empty results or with parameters set by the user. This is an internal function used as a preliminary step to obtain predictions for new data and is not intended to be used directly.

Usage

createSaemixObject.empty(model, data, control = list())

Arguments

model an saemixModel object
data an saemixData object
control a list of options (if empty, will be set to the default values by saemixControl)

Details

with createSaemixObject.empty, the data component is set to the data object, the model component is set to the model object, and the result component is empty

with createSaemixObject.initial, the data and model are set as with createSaemixObject.empty, but the population parameter estimates are used to initialise the result component as in the initialisation step of the algorithm (initialiseMainAlgo)

Value

an object of class "SaemixObject".

Examples

# TODO
dataGen.case  

**Bootstrap datasets**

Description

These functions create bootstrapped datasets using the bootstrap methods described in `saemix.bootstrap`.

Usage

dataGen.case(saemixObject)

dataGen.NP(saemixObject, nsamp = 0, eta.sampc = NULL, conditional = FALSE)

dataGen.Par(saemixObject)

Arguments

- **saemixObject**: an object returned by the `saemix` function
- **nsamp**: number of samples from the conditional distribution (for method="conditional")
- **eta.sampc**: if available, samples from the conditional distribution (otherwise, they are obtained within the function)
- **conditional**: if TRUE, sample from the conditional distribution, if FALSE, sample within the empirical Bayes estimates (EBE) as in the traditional non-parametric residual bootstrap

default.saemix.plots  

**Wrapper functions to produce certain sets of default plots**

Description

These functions produce default sets of plots, corresponding to diagnostic or individual fits.

Usage

default.saemix.plots(saemixObject, ...)

Arguments

- **saemixObject**: an object returned by the `saemix` function
- **...**: optional arguments passed to the plots

Details

These functions are wrapper functions designed to produce default sets of plots to help the user assess their model fits.
Value

Depending on the type argument, the following plots are produced:

- default.saemix.plots by default, the following plots are produced: a plot of the data, convergence plots, plot of the likelihood by importance sampling (if computed), plots of observations versus predictions, scatterplots and distribution of residuals, boxplot of the random effects, correlations between random effects, distribution of the parameters, VPC
- basic.gof basic goodness-of-fit plots: convergence plots, plot of the likelihood by importance sampling (if computed), plots of observations versus predictions
- advanced.gof advanced goodness-of-fit plots: scatterplots and distribution of residuals, VPC,...
- covariate.fits plots of all estimated parameters versus all covariates in the dataset
- individual.fits plots of individual predictions (line) overlayed on individual observations (dots) for all subjects in the dataset

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

saemix, saemix.plot.data, saemix.plot.setoptions, plot.saemix

Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
name.group=c("Id"),name.predictors=c("Dose","Time"),
name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
\begin{verbatim}
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}

saemix.model<-saemixModel(model=model1cpt,
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
    dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")

# Reducing the number of iterations due to time constraints for CRAN
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE,nbiter.saemix=c(100,100))

saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

default.saemix.plots(saemix.fit)

# Not run (time constraints for CRAN)
# basic.gof(saemix.fit)

# Not run (time constraints for CRAN)
# advanced.gof(saemix.fit)

individual.fits(saemix.fit)
\end{verbatim}

---

discreteVPC  VPC for non Gaussian data models

description

This function provides VPC plots for non Gaussian data models (work in progress)

usage

discreteVPC(object, outcome = "categorical", verbose = FALSE, ...)

arguments

object an saemixObject object returned by the \texttt{saemix} function. The object must include simulated data under the empirical design, using the model and estimated parameters from a fit, produced via the \texttt{simulateDiscreteSaemix} function.

outcome type of outcome (valid types are "TTE", "binary", "categorical", "count")

verbose whether to print messages (defaults to FALSE)

... additional arguments, used to pass graphical options (to be implemented, currently not available)
Details

This function is a very rough first attempt at automatically creating VPC plots for models defined through their log-likelihood (categorical, count, or TTE data). It makes use of the new element simulate.function in the model component of the object

- for TTE data, a KM-VPC plot will be produced
- for count, categorical and binary data, a plot showing the proportion of each score/category across time will be shown along with the corresponding prediction intervals from the model

These plots can be stratified over a covariate in the data set (currently only categorical covariates) by passing an argument which.cov='name’ to the call

Author(s)

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References


Ron Keizer, tutorials on VPC TODO

See Also

SaemixObject, saemix, saemix.plot.vpc, simulateDiscreteSaemix

Examples

data(lung.saemix)

saemix.data<-saemixData(name.data=lung.saemix,header=TRUE,name.group=c("id"),
name.predictors=c("time","status","cens"),name.response=c("status"),
name.covariates=c("age","sex","ph.ecog","ph.karno","pat.karno","wt.loss","meal.cal"),
units=list(x="days",y="",covariates=c("yr","","-","%","%","cal","pounds")))

weibulltte.model<-function(psi,id,xidep) {
  T<-xidep[,1]
  y<-xidep[,2] # events (1=event, 0=no event)
  cens<-which(xidep[,3]==1) # censoring times (subject specific)
  init <- which(T==0)
  lambda <- psi[id,1] # Parameters of the Weibull model
  beta <- psi[id,2]
  Nj <- length(T)
  ind <- setdiff(1:Nj, append(init,cens)) # indices of events
  hazard <- (beta/lambda)*(T/lambda)^(beta-1) # H'
  H <- (T/lambda)^beta # H
  logpdf <- rep(0,Nj) # ln(ln(l(T=0))=0
  logpdf[cens] <- -H[cens] + H[cens-1] # ln(ln(l(T=censoring time)))
logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind]) # ln(l(T=event time))
return(logpdf)
}

simulateWeibullTTE <- function(psi, id, xidep) {
  T <- xidep[,1]
  y <- xidep[,2] # events (1=event, 0=no event)
  cens <- which(xidep[,3] == 1) # censoring times (subject specific)
  init <- which(T == 0)
  lambda <- psi[,1] # Parameters of the Weibull model
  beta <- psi[,2]
  tevent <- T
  Vj <- runif(dim(psi)[1])
  tsim <- lambda*(-log(Vj))^(1/beta) # nsuj events
  tevent[T > 0] <- tsim
  return(tevent)
}

saemix.model <- saemixModel(model=weibulltte.model, description="time model", modeltype="likelihood",
  simulate.function = simulateWeibullTTE,
  psi0 = matrix(c(1, 2), ncol=2, byrow=TRUE, dimnames=list(NULL, c("lambda", "beta"))),
  transform.par = c(1, 1), covariance.model = matrix(c(1, 0, 0, 0), ncol=2, byrow=TRUE))

saemix.options <- list(seed=632545, save=FALSE, save.graphs=FALSE, displayProgress=FALSE)

tte.fit <- saemix(saemix.model, saemix.data, saemix.options)
simtte.fit <- simulateDiscreteSaemix(tte.fit, nsim=100)
gpl <- discreteVPC(simtte.fit, outcome="TTE")
plot(gpl)

---

discreteVPCTTE  VPC for time-to-event models

Description
This function provides VPC plots for time-to-event data models (work in progress)

Usage

discreteVPCTTE(
  object,
  ngrid = 200,
  interpolation.method = "step",
  verbose = FALSE,
  ...
)
Arguments

- **object**: an saemixObject object returned by the `saemix` function. The object must include simulated data under the empirical design, using the model and estimated parameters from a fit, produced via the `simulateDiscreteSaemix` function.
- **ngrid**: number of grid points on the X-axis to extrapolate the KM-VPC.
- **interpolation.method**: method to use for the interpolation of the KM for the simulated datasets. Available methods are "step": the value of the survival function for a given grid point is set to the value of the last time "lin": a linear approximation is used between two consecutive times (defaults to "step")
- **verbose**: whether to print messages (defaults to FALSE).
- **...**: additional arguments, used to pass graphical options (to be implemented, currently not available).

Details

- add details on TTE VPC, RTTE VPC, etc...

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr

See Also

- `SaemixObject`, `saemix`, `saemix.plot.vpc`, `simulateDiscreteSaemix`
- Tutorials on TTE-VPC TODO

---

**epilepsy.saemix**

**Epilepsy count data**

Description

The epilepsy data from Thall and Vail (1990), available from the MASS package, records two-week seizure counts for 59 epileptics. The number of seizures was recorded for a baseline period of 8 weeks, and then patients were randomly assigned to a treatment group or a control group. Counts were then recorded for four successive two-week periods. The subject’s age is the only covariate. See the documentation for epil in the MASS package for details on the dataset.

Source

MASS package in R

References

Examples

# You need to have MASS installed to successfully run this example
if (requireNamespace("MASS")) {
  epilepsy<-MASS::epil
  saemix.data<-saemixData(name.data=epilepsy, name.group=c("subject"),
    name.predictors=c("period","y"),name.response=c("y"),
    name.covariates=c("trt","base", "age"), units=list(x="2-week",y="",covariates=c("","","yr")))
  ## Poisson model with one parameter
  countPoi< function(psi,id,xidep) {
    y<xidep[,2]
    lambda<psi[id,1]
    logp <- -lambda + y*log(lambda) - log(factorial(y))
    return(logp)
  }
  saemix.model<-saemixModel(model=countPoi,description="Count model Poisson",modeltype="likelihood",
    psi0=matrix(c(0.5),ncol=1,byrow=TRUE,dimnames=list(NULL, c("lambda"))), transform.par=c(1))
  saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE)
  poisson.fit<-saemix(saemix.model,saemix.data,saemix.options)
}

fim.saemix

Computes the Fisher Information Matrix by linearisation

Description

Estimate by linearisation the Fisher Information Matrix and the standard error of the estimated parameters.

Usage

fim.saemix(saemixObject)

Arguments

saemixObject an object returned by the saemix function

Details

The inverse of the Fisher Information Matrix provides an estimate of the variance of the estimated parameters theta. This matrix cannot be computed in closed-form for nonlinear mixed-effect models; instead, an approximation is obtained as the Fisher Information Matrix of the Gaussian model deduced from the nonlinear mixed effects model after linearisation of the function f around the conditional expectation of the individual Gaussian parameters. This matrix is a block matrix (no correlations between the estimated fixed effects and the estimated variances).
Value

The function returns an updated version of the object saemix.fit in which the following elements have been added:

- **se.fixed**: standard error of fixed effects, obtained as part of the diagonal of the inverse of the Fisher Information Matrix (only when fim.saemix has been run, or when saemix.options$algorithms[2] is 1)
- **se.omega**: standard error of the variance of random effects, obtained as part of the diagonal of the inverse of the Fisher Information Matrix (only when fim.saemix has been run, or when the saemix.options$algorithms[2] is 1)
- **se.res**: standard error of the parameters of the residual error model, obtained as part of the diagonal of the inverse of the Fisher Information Matrix (only when fim.saemix has been run, or when the saemix.options$algorithms[2] is 1)
- **fim**: Fisher Information Matrix
- **ll.lin**: likelihood calculated by linearisation

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

SaemixObject,saemix

Examples

```R
# Running the main algorithm to estimate the population parameters
data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep= " ",na=NA, name.group=c("Id"),name.predictors=c("Dose","Time"), name.response=c("Concentration"),name.covariates=c("Weight","Sex"), units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ...
}
```
ka<-ψ[1,1]
V<-ψ[1,2]
CL<-ψ[1,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
return(ypred)

saemix.model<-saemixModel(model=model1cpt,
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
    dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE), error.model="constant")

saemix.options<-list(algorithm=c(1,0,0),seed=632545,save=FALSE,save.graphs=FALSE)

# Not run (strict time constraints for CRAN)
# saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

# Estimating the Fisher Information Matrix using the result of saemix
# & returning the result in the same object
# fim.saemix(saemix.fit)

---

**fitted.saemix**  
*Extract Model Predictions*

**Description**

fitted is a generic function which extracts model predictions from objects returned by modelling functions.

**Usage**

```r
# S3 method for class 'SaemixRes'
fitted(object, type = c("ipred", "ypred", "ppred", "icpred"), ...)
```

```r
# S3 method for class 'SaemixObject'
fitted(object, type = c("ipred", "ypred", "ppred", "icpred"), ...)
```

**Arguments**

- **object**: an object of type SaemixRes or SaemixObject
- **type**: string determining which predictions are extracted. Possible values are: "ipred" (individual predictions obtained using the mode of the individual distribution for each subject, default), "ppred" (population predictions obtained using the mode of the population distribution, default), "ypred" (model predictions based on the estimated model parameters), "icpred" (individual predictions based on the estimated model parameters)
population parameters $f(E(\theta))$, "ypred" (mean of the population predictions $E(f(\theta)))$ and "icpred" (individual predictions obtained using the conditional mean of the individual distribution). See user guide for details.

... further arguments to be passed to or from other methods

**Value**

Model predictions

---

**Description**

Joint selection of covariates and random effects in a nonlinear mixed effects model by a forward-type algorithm based on two different versions of BIC for covariate selection and random effects selection respectively. Selection is made among the covariates as such specified in the SaemixData object. Only uncorrelated random effects structures are considered.

**Usage**

`forward.procedure(saemixObject, trace = TRUE)`

**Arguments**

- `saemixObject` An object returned by the `saemix` function
- `trace` If TRUE, a table summarizing the steps of the algorithm is printed. Default "TRUE"

**Value**

An object of the SaemixObject class storing the covariate model and the covariance structure of random effects of the final model.

**Author(s)**

Maud Delattre

**References**


Methods for Function initialize

Description

Constructor functions for Classes in the saemix package

Usage

```r
## S4 method for signature 'SaemixData'
initialize(
  .Object,
  name.data,
  header,
  sep,
  na,
  name.group,
  name.predictors,
  name.response,
  name.covariates,
  name.X,
  units,
  name.mdv,
  name.cens,
  name.occ,
  name.ytype,
  verbose = TRUE,
  automatic = TRUE
)

## S4 method for signature 'SaemixRepData'
initialize(.Object, data = NULL, nb.chains = 1)

## S4 method for signature 'SaemixSimData'
initialize(.Object, data = NULL, datasim = NULL)

## S4 method for signature 'SaemixModel'
initialize(
  .Object,
  model,
  description,
  modeltype,
  psi0,
  name.response,
  name.sigma,
  transform.par,
  fixed.estim,
```
initialize-methods

```r
initialize(
  .Object,
  status = "empty",
  modeltype,
  name.fixed,
  name.random,
  name.sigma,
  fixed.effects,
  fixed.psi,
  betaC,
  betas,
  omega,
  respar,
  cond.mean.phi,
  cond.var.phi,
  mean.phi,
  phi,
  phi.samp,
  parpop,
  allpar,
  MCOV
)

# S4 method for signature 'SaemixRes'
initialize(
  .Object,
  status = "empty",
  modeltype,
  name.fixed,
  name.random,
  name.sigma,
  fixed.effects,
  fixed.psi,
  betaC,
  betas,
  omega,
  respar,
  cond.mean.phi,
  cond.var.phi,
  mean.phi,
  phi,
  phi.samp,
  parpop,
  allpar,
  MCOV
)

# S4 method for signature 'SaemixObject'
initialize(.Object, data, model, options = list())
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>.Object</code></td>
<td>a SaemixObject, SaemixRes, SaemixData or SaemixModel object to initialise</td>
</tr>
<tr>
<td><code>name.data</code></td>
<td>name of the dataset (can be a character string giving the name of a file on disk or of a dataset in the R session, or the name of a dataset)</td>
</tr>
<tr>
<td><code>header</code></td>
<td>whether the dataset/file contains a header. Defaults to TRUE</td>
</tr>
<tr>
<td><code>sep</code></td>
<td>the field separator character. Defaults to any number of blank spaces (&quot;&quot;)</td>
</tr>
<tr>
<td><code>na</code></td>
<td>a character vector of the strings which are to be interpreted as NA values. Defaults to c(NA)</td>
</tr>
<tr>
<td><code>name.group</code></td>
<td>name (or number) of the column containing the subject id</td>
</tr>
</tbody>
</table>
name.predictors
  name (or number) of the column(s) containing the predictors (the algorithm requires at least one predictor x)

name.response
  name (or number) of the column containing the response variable y modelled by predictor(s) x

name.covariates
  name (or number) of the column(s) containing the covariates, if present (otherwise missing)

name.X
  name of the column containing the regression variable to be used on the X axis in the plots (defaults to the first predictor)

units
  list with up to three elements, x, y and optionally covariates, containing the units for the X and Y variables respectively, as well as the units for the different covariates (defaults to empty)

name.mdv
  name of the column containing the indicator for missing variable

name.cens
  name of the column containing the indicator for censoring

name.occ
  name of the column containing the occasion

name.ytype
  name of the column containing the index of the response

verbose
  a boolean indicating whether messages should be printed out during the creation of the object

automatic
  a boolean indicating whether to attempt automatic name recognition when some column names are missing or wrong (defaults to TRUE)

data
  an SaemixData object

nb.chains
  number of chains used in the algorithm

datasim
  dataframe containing the simulated data

model
  name of the function used to compute the structural model. The function should return a vector of predicted values given a matrix of individual parameters, a vector of indices specifying which records belong to a given individual, and a matrix of dependent variables (see example below).

description
  a character string, giving a brief description of the model or the analysis

modeltype
  a character string giving the model used for analysis

psi0
  a matrix with a number of columns equal to the number of parameters in the model, and one (when no covariates are available) or two (when covariates enter the model) giving the initial estimates for the fixed effects. The column names of the matrix should be the names of the parameters in the model, and will be used in the plots and the summaries. When only the estimates of the mean parameters are given, psi0 may be a named vector.

name.sigma
  a vector of character string giving the names of the residual error parameters (defaults to "a" and "b")

transform.par
  the distribution for each parameter (0=normal, 1=log-normal, 2=probit, 3=logit). Defaults to a vector of 1s (all parameters have a log-normal distribution)

fixed.estim
  whether parameters should be estimated (1) or fixed to their initial estimate (0). Defaults to a vector of 1s
initialize-methods

error.model type of residual error model (valid types are constant, proportional, combined and exponential). Defaults to constant

covariate.model a matrix giving the covariate model. Defaults to no covariate in the model

covariance.model a square matrix of size equal to the number of parameters in the model, giving the variance-covariance matrix of the model: 1s correspond to estimated variances (in the diagonal) or covariances (off-diagonal elements). Defaults to the identity matrix

omega.init a square matrix of size equal to the number of parameters in the model, giving the initial estimate for the variance-covariance matrix of the model. The current default is a diagonal matrix with ones for all transformed parameters as well as for all untransformed parameters with an absolute value smaller than one. For untransformed parameters greater or equal to one, their squared value is used as the corresponding diagonal element.

error.init a vector of size 2 giving the initial value of a and b in the error model. Defaults to 1 for each estimated parameter in the error model

name.modpar names of the model parameters, if they are not given as the column names (or names) of psi0

status string indicating whether a model has been run successfully; set to "empty" at initialisation, used to pass on error messages or fit status

name.fixed a character string giving the name of the fixed parameters

name.random a character string giving the name of the random parameters

fixed.effects vector with the estimates of h(mu) and betas in estimation order

fixed.psi vector with the estimates of h(mu)

betaC vector with the estimates of betas (estimated fixed effects for covariates)

betas vector with the estimates of mu

omega estimated variance-covariance matrix

respar vector with the estimates of the parameters of the residual error

cond.mean.phi matrix of size (number of subjects) x (nb of parameters) containing the conditional mean estimates of the, defined as the mean of the conditional distribution

cond.var.phi matrix of the variances on cond.mean.phi, defined as the variance of the conditional distribution

mean.phi matrix of size (number of subjects) x (nb of parameters) giving for each subject the estimates of the population parameters including covariate effects

phi matrix of size (number of subjects) x (nb of parameters) giving for each subject

phi.samp samples from the individual conditional distributions of the phi

parpop population parameters at each iteration

allpar all parameters (including covariate effects) at each iteration

MCOV design matrix C

options a list of options passed to the algorithm
Methods

- `list("signature(.Object = "SaemixData")")` create a SaemixData object. Please use the `saemixData` function.
- `list("signature(.Object = "SaemixModel")")` create a SaemixModel object Please use the `saemixModel` function.
- `list("signature(.Object = "SaemixObject")")` create a SaemixObject object. This object is obtained after a successful call to `saemix`
- `list("signature(.Object = "SaemixRepData")")` create a SaemixRepData object
- `list("signature(.Object = "SaemixRes")")` create a SaemixRes object
- `list("signature(.Object = "SaemixSimData")")` create a SaemixSimData object

---

knee.saemix  Knee pain data

Description

The knee.saemix data represents pain scores recorded in a clinical study in 127 patients with sport related injuries treated with two different therapies. After 3, 7 and 10 days of treatment the pain occurring during knee movement was observed.

Usage

knee.saemix

Format

This data frame contains the following columns:

- **id** subject index in file
- **time** time of measurement (in days)
- **y** knee pain (0=none to 4=severe)
- **Age** patient age (scaled and centered)
- **Sex** patient gender (0=male, 1=female)
- **RD** moderate knee pain (defined as pain score 2 or more)
- **treatment** treatment indicator (0=placebo, 1=treatment)
- **Age2** patient age, squared (Age^2)

#
Details

The data in the knee.saemix was reformatted from the knee dataset provided by the catdata package (see data(knee, package="catdata")). A time column was added representing the day of the measurement (with 0 being the baseline value) and each observation corresponds to a different line in the dataset. Treatment was recoded as 0/1 (placebo/treatment), gender as 0/1 (male/female) and Age^2 represents the squared of centered Age.

Please refer to the PDF documentation (chapter 4, section 4.6) for more details on the analysis, including examples of diagnostic plots.

Source

catdata package in R

References


#' @examples data(knee.saemix)
#' @keywords datasets

Description

Estimate the log-likelihood using Gaussian Quadrature (multidimensional grid)

Usage

llgq.saemix(saemixObject)

Arguments

saemixObject an object returned by the saemix function

Details

The likelihood of the observations is estimated using Gaussian Quadrature (see documentation).

Value

the log-likelihood estimated by Gaussian Quadrature

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.
References


See Also

SaemixObject, saemix, llis.saemix

Examples

# Running the main algorithm to estimate the population parameters
data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix, header=TRUE, sep=" ", na=NA, name.group=c("Id"), name.predictors=c("Dose", "Time"), name.response=c("Concentration"), name.covariates=c("Weight", "Sex"), units=list(x="hr", y="mg/L", covariates=c("kg", "-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
dose<-xidep[,1]
tim<-xidep[,2]
ka<-psi[id,1]
V<-psi[id,2]
CL<-psi[id,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
return(ypred)
}
saemix.model<-saemixModel(model=model1cpt,
description="One-compartment model with first-order absorption",
psi0=matrix(c(1,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
dimnames=list(NULL, c("ka", "V", "CL")), transform.par=c(1,1,1),
covariate.model=matrix(c(0,1,0,0,0),ncol=3, byrow=TRUE), fixed.estim=c(1,1,1),
covariance.model=matrix(c(1,0,0,1,0,0,1),ncol=3, byrow=TRUE),
omega.init=matrix(c(1,0,0,1,0,0,1),ncol=3, byrow=TRUE), error.model="constant")

saemix.options<-list(seed=632545, save=FALSE, save.graphs=FALSE)

# Not run (strict time constraints for CRAN)
# saemix.fit<-saemix(saemix.model, saemix.data, saemix.options)

# Estimating the likelihood by Gaussian Quadrature using the result of saemix
# & returning the result in the same object
# saemix.fit<-llgq.saemix(saemix.fit)
llis.saemix

Log-likelihood using Importance Sampling

Description

Estimate the log-likelihood using Importance Sampling

Usage

llis.saemix(saemixObject)

Arguments

saemixObject  an object returned by the saemix function

Details

The likelihood of the observations is estimated without any approximation using a Monte-Carlo approach (see documentation).

Value

the log-likelihood estimated by Importance Sampling

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

SaemixObject,saemix,llgq.saemix
Examples

```r
# Running the main algorithm to estimate the population parameters
data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA, 
  name.group=c("Id"),name.predictors=c("Dose","Time"), 
  name.response=c("Concentration"),name.covariates=c("Weight","Sex"), 
  units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}
saemix.model<-saemixModel(model=model1cpt, 
  description="One-compartment model with first-order absorption", modeltype="structural", 
  psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE, 
    dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")
saemix.options<-list(algorithm=c(1,0,0),seed=632545,save=FALSE,save.graphs=FALSE)

# Not run (strict time constraints for CRAN)
# saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

# Estimating the likelihood by importance sampling using the result of saemix 
# & returning the result in the same object
# saemix.fit<-llis.saemix(saemix.fit)
```

---

**logLik**

Extract likelihood from a saemixObject resulting from a call to saemix

**Description**

The likelihood in saemix can be computed by one of three methods: linearisation (linearisation of the model), importance sampling (stochastic integration) and gaussian quadrature (numerical integration). The linearised likelihood is obtained as a byproduct of the computation of the Fisher Information Matrix (argument FIM=TRUE in the options given to the saemix function). If no method argument is given, this function will attempt to extract the likelihood computed by importance sampling (method="is"), unless the object contains the likelihood computed by linearisation, in which case the function will extract this component instead. If the requested likelihood is not present in the object, it will be computed and added to the object before returning.
Usage

## S3 method for class 'SaemixObject'
logLik(object, method = c("is", "lin", "gq"), ...)

## S3 method for class 'SaemixObject'
AIC(object, method = c("is", "lin", "gq"), ..., k = 2)

## S3 method for class 'SaemixObject'
BIC(object, method = c("is", "lin", "gq"), ...)

## S3 method for class 'covariate'
BIC(object, method = c("is", "lin", "gq"), ...)

Arguments

- **object**: name of an SaemixObject object
- **method**: character string, one of c("is","lin","gq"), to select one of the available approximations to the log-likelihood (is: Importance Sampling; lin: linearisation and gq: Gaussian Quadrature). See documentation for details
- **...**: additional arguments
- **k**: numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC

Details

BIC.covariate implements the computation of the BIC from Delattre et al. 2014.

Value

Returns the selected statistical criterion (log-likelihood, AIC, BIC) extracted from the SaemixObject, computed with the 'method' argument if given (defaults to IS).

Author(s)

Emmanuelle Comets <emmanuelle.comets@inserm.fr>
Audrey Lavenu
Marc Lavielle
Maud Delattre

References


See Also

AIC, BIC, saemixControl, saemix

lung.saemix  NCCTG Lung Cancer Data, in SAEM format

Description

The lung.saemix contains survival data in patients with advanced lung cancer from the North Central Cancer Treatment Group. Performance scores rate how well the patient can perform usual daily activities. This data is available in the survival library for R and has been reformatted here for use in saemix (see details).

Usage

lung.saemix

Format

This data frame contains the following columns:

<table>
<thead>
<tr>
<th>Column</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>id</td>
<td>subject index in file</td>
</tr>
<tr>
<td>inst</td>
<td>institution code</td>
</tr>
<tr>
<td>time</td>
<td>observation time since the beginning of follow-up</td>
</tr>
<tr>
<td>status</td>
<td>0=alive, 1=dead</td>
</tr>
<tr>
<td>cens</td>
<td>0=observed, 1=censored</td>
</tr>
<tr>
<td>sex</td>
<td>patient gender (0=male, 1=female)</td>
</tr>
<tr>
<td>age</td>
<td>age in years</td>
</tr>
<tr>
<td>ph.ecog</td>
<td>ECOG performance score as rated by the physician.</td>
</tr>
<tr>
<td></td>
<td>0=asymptomatic, 1= symptomatic but completely ambulatory, 2= in bed &lt;50% of the day, 3= in bed &gt; 50% of the day but not bedbound, 4 = bedbound</td>
</tr>
<tr>
<td>ph.karno</td>
<td>Karnofsky performance score (bad=0-good=100) rated by physician (%)</td>
</tr>
<tr>
<td>pat.karno</td>
<td>Karnofsky performance score (bad=0-good=100) rated by patient (%)</td>
</tr>
<tr>
<td>meal.cal</td>
<td>calories consumed at meals (cal)</td>
</tr>
<tr>
<td>wt.loss</td>
<td>weight loss in last six months (pounds)</td>
</tr>
</tbody>
</table>

Details

The data in the lung.saemix was reformatted from the lung cancer dataset (see data(cancer, package="survival"). Patients with missing age, sex, institution or physician assessments were removed from the dataset. Status was recoded as 1 for death and 0 for a censored event, and a censoring column was added to denote whether the patient was dead or alive at the time of the last observation. For saemix, a line at time=0 was added for all subjects. Finally, subjects were numbered consecutively from 0 to 1.
Source

Terry Therneau from the survival package in R

References


Examples

data(lung.saemix)

saemix.data<-saemixData(name.data=lung.saemix,header=TRUE,name.group=c("id"),
name.predictors=c("time","status","cens"),name.response=c("status"),
name.covariates=c("age","sex","ph.ecog","ph.karno","pat.karno","wt.loss","meal.cal"),
units=list(x="days",y="",covariates=c("yr","","%","%","","cal","pounds"))

weibulltte.model<-function(psi,id,xidep) {
  T<-xidep[,1] # events (1=event, 0=no event)
cens<-which(xidep[,3]==1) # censoring times (subject specific)
init <- which(T==0)
lambda <- psi[id,1] # Parameters of the Weibull model
beta <- psi[id,2]
Nj <- length(T)
ind <- setdiff(1:Nj, append(init,cens)) # indices of events
hazard <- (beta/lambda)*(T/lambda)^(beta-1) # H'
H <- (T/lambda)^beta # H
logpdf <- rep(0,Nj) # ln(l(T=0))=0
logpdf[cens] <- -H[cens] + H[cens-1] # ln(l(T=censoring time))
logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind]) # ln(l(T=event time))
return(logpdf)
}

saemix.model<-saemixModel(model=weibulltte.model,description="time model",modeltype="likelihood",
psi0=matrix(c(1,2),ncol=2,byrow=TRUE,dimnames=list(NULL, c("lambda","beta"))),
transform.par=c(1,1),covariance.model=matrix(c(1,0,0,0),ncol=2, byrow=TRUE)

saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE)

tte.fit<-saemix(saemix.model,saemix.data,saemix.options)

# The fit from saemix using the above Weibull model may be compared
# to the non-parametric KM estimate
## Not run:
library(survival)
lung.surv<-lung.saemix[lung.saemix$time>0,]
lung.surv$status<-lung.surv$status+1
Surv(lung.surv$time, lung.surv$status) # 1=censored, 2=dead
f1 <- survfit(Surv(time, status) ~ 1, data = lung.surv)
xtim<-seq(0,max(lung.saemix$time), length.out=200)
estpar<-tte.fit$results@fixed.effects
ypred<-exp(-(-xtim/estpar[1])^(estpar[2]))
plot(f1, xlab = "Days", ylab = "Overall survival probability")
map.saemix

lines(xtim, ypred, col="red", lwd=2)

## End(Not run)

map.saemix

Estimates of the individual parameters (conditional mode)

Description

Compute the estimates of the individual parameters PSI_i (conditional mode - Maximum A Posteriori)

Usage

map.saemix(saemixObject)

Arguments

saemixObject an object returned by the saemix function

Details

The MCMC procedure is used to estimate the conditional mode (or Maximum A Posteriori) m(phi_i | y_i; hattheta) = Argmax_phi_i p(phi_i | y_i; hattheta)

Value

saemixObject: returns the object with the estimates of the MAP parameters (see example for usage)

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

SaemixObject, saemix
Examples

```r
data(theo.saemix)

saemix.data <- saemixData(name.data = theo.saemix, header = TRUE, sep = " ", na = NA,
  name.group = c("Id"), name.predictors = c("Dose", "Time"),
  name.response = c("Concentration"), name.covariates = c("Weight", "Sex"),
  units = list(x = "hr", y = "mg/L", covariates = c("kg", "-")), name.X = "Time")

model1cpt <- function(psi, id, xidep) {
  dose <- xidep[, 1]
  tim <- xidep[, 2]
  ka <- psi[id, 1]
  V <- psi[id, 2]
  CL <- psi[id, 3]
  k <- CL / V
  ypred <- dose * ka / (V * (ka - k) * (exp(-k * tim) - exp(-ka * tim))
  return(ypred)
}

saemix.model <- saemixModel(model = model1cpt,
  description = "One-compartment model with first-order absorption",
  psi0 = matrix(c(1., 20, 0.5, 0.1, 0, -0.01), ncol = 3, byrow = TRUE,
    dimnames = list(NULL, c("ka", "V", "CL"))),
  transform.par = c(1, 1, 1),
  covariate.model = matrix(c(0, 1, 0, 0, 0, 0), ncol = 3, byrow = TRUE),
  fixed.estim = c(1, 1, 1),
  covariance.model = matrix(c(1, 0, 0, 1, 0, 0, 1, 0, 0, 1, 0, 0, 1), ncol = 3, byrow = TRUE),
  omega.init = matrix(c(1, 0, 0, 1, 0, 0, 1), ncol = 3, byrow = TRUE),
  error.model = "constant")

saemix.options <- list(algorithm = c(1, 0, 0), seed = 632545,
  save = FALSE, save.graphs = FALSE)

# Not run (strict time constraints for CRAN)
# saemix.fit <- saemix(saemix.model, saemix.data, saemix.options)

# Estimating the individual parameters using the result of saemix
# & returning the result in the same object
# saemix.fit <- map.saemix(saemix.fit)
```

---

**mydiag**

*Matrix diagonal*

**Description**

Extract or replace the diagonal of a matrix, or construct a diagonal matrix (replace `diag` function from R-base)

**Usage**

```r
mydiag(x = 1, nrow, ncol)
```
Arguments

- **x**: a matrix, vector or 1D array, or missing.
- **nrow**: Optional number of rows for the result when x is not a matrix.
- **ncol**: Optional number of columns for the result when x is not a matrix.

Value

If x is a matrix then diag(x) returns the diagonal of x. The resulting vector will have names if the matrix x has matching column and rownames.

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

See Also

diag

Examples

```
mydiag(1)
mydiag(c(1,2))
```

---

**npdeSaemix**

Create an npdeObject from an saemixObject

Description

This function uses the npde library to compute normalised prediction distribution errors (npde) and normalised prediction discrepancies (npd). The simulations can also be used to plot VPCs. As of version 3.0, the plot functions for diagnostics involving VPC, npde or npd are deprecated and the user will be redirected to the current proposed method (creating an NpdeObject and using the plot functions from the library npde).

Usage

```
npdeSaemix(saemixObject, nsim = 1000)
```

Arguments

- **saemixObject**: a fitted object resulting from a call to saemix()
- **nsim**: the number of simulations used to compute npde (1000 by default, we suggest increasing it for large datasets)
Details

Since version 3.0, the saemix package depends on the npde package, which computes the npd/npde and produces graphs. See the documentation for npde for details on the computation methods. See the PDF documentation and the bookdown https://iame-researchcenter.github.io/npde_bookdown/ for details on the different plots available.

Value

An object of class NpdeObject

Author(s)

Emmanuelle Comets emmanuelle.comets@bichat.inserm.fr

References


See Also

npde.graphs, gof.test

NpdeObject npde.plot.select autonpde

npde.plot.scatterplot npde.plot.dist

Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
  name.group=c("Id"),name.predictors=c("Dose","Time"),
  name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
  units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
oxboys.saemix <- saemixModel(model=model1cpt, 
  description="One-compartment model with first-order absorption", 
  psi0=matrix(c(1,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE, 
  dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1), 
  covariate.model=matrix(c(0,1,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1), 
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE), 
  omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")

saemix.options<-list(algorithm=c(1,0,0),seed=632545,save=FALSE,save.graphs=FALSE, 
  displayProgress=FALSE)

# Not run
# Works interactively but not in the contained environment of CRAN (it looks for a datafile
# instead of finding the dataset in the environment)
# saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)
# npde.obj<-npdeSaemix(saemix.fit)
# plot(npde.obj)
# plot(npde.obj, plot.type="vpc")
# plot(npde.obj, plot.type="covariates")
# plot(npde.obj, plot.type="cov.x.scatter")
# plot(npde.obj, plot.type="cov.ecdf")

oxboys.saemix 

Heights of Boys in Oxford

Description

The oxboys.saemix data collects the height and age of students in Oxford measured over time. 
The data frame has 234 rows and 4 columns.

Usage

oxboys.saemix

Format

This data frame contains the following columns:

Subject an ordered factor giving a unique identifier for each boy in the experiment
age a numeric vector giving the standardized age (dimensionless)
height a numeric vector giving the height of the boy (cm)
Occasion an ordered factor - the result of converting 'age' from a continuous variable to a count
so these slightly unbalanced data can be analyzed as balanced
Details

These data are described in Goldstein (1987) as data on the height of a selection of boys from Oxford, England versus a standardized age. The dataset can be found in the package nlme. We use an linear model for this data: \( y_{ij} = Base_i + slope_i x_{ij} + \epsilon_{ij} \)

References


Examples

data(oxboys.saemix)
saemix.data<-saemixData(name.data=oxboys.saemix,header=TRUE, name.group=c("Subject"),name.predictors=c("age"),name.response=c("height"), units=list(x="yr",y="cm"))

# plot the data
plot(saemix.data)

growth.linear<-function(psi,id,xidep) {
  x<-xidep[,1]
  base<-psi[id,1]
  slope<-psi[id,2]
  f<-base+slope*x
  return(f)
}
saemix.model<-saemixModel(model=growth.linear,description="Linear model", psi0=matrix(c(140,1),ncol=2,byrow=TRUE,dimnames=list(NULL,c("base","slope"))), transform.par=c(1,0),covariance.model=matrix(c(1,1,1,1),ncol=2,byrow=TRUE), error.model="constant")

saemix.options<-list(algorithms=c(1,1,1),nb.chains=1,seed=201004, save=FALSE,save.graphs=FALSE,displayProgress=FALSE)

saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

---

**PD1.saemix**

*Data simulated according to an Emax response model, in SAEM format*

Description

The PD1.saemix and PD2.saemix data frames were simulated according to an Emax dose-response model.
PD1.saemix

PD2.saemix

Usage

PD1.saemix

PD2.saemix

Format

This data frame contains the following columns:

- **subject**: an variable identifying the subject on whom the observation was made. The ordering is by the dose at which the observation was made.
- **dose**: simulated dose.
- **response**: simulated response.
- **gender**: gender (0 for male, 1 for female).

Details

These examples were used by P. Girard and F. Mentre for the symposium dedicated to Comparison of Algorithms Using Simulated Data Sets and Blind Analysis, that took place in Lyon, France, September 2004. The datasets contain 100 individuals, each receiving 3 different doses: (0, 10, 90), (5, 25, 65) or (0, 20, 30). It was assumed that doses were given in a cross-over study with sufficient wash out period to avoid carry over. Responses \( y_{ij} \) were simulated with the following pharmacodynamic model:

\[
y_{ij} = E0_i + D_{ij} Emax_i/(D_{ij} + ED50_i) + \epsilon_{ij}
\]

The individual parameters were simulated according to

\[
\begin{align*}
\log(E0_i) &= \log(E0) + \eta_{i1} \\
\log(Emax_i) &= \log(Emax) + \eta_{i2} \\
\log(ED50_i) &= \log(ED50) + \beta w_i + \eta_{i3}
\end{align*}
\]

PD1.saemix contains the data simulated with a gender effect, \( \beta = 0.3 \). PD2.saemix contains the data simulated without a gender effect, \( \beta = 0 \).

References


Examples

data(PD1.saemix)
saemix.data<-saemixData(name.data=PD1.saemix,header=TRUE,name.group=c("subject"),
                        name.predictors=c("dose"),name.response=c("response"),
                        name.covariates=c("gender"), units=list(x="mg",y="-",covariates=c("-"))

modelemax<-function(psi,id,xidep) {
  # input:
  # psi : matrix of parameters (3 columns, E0, Emax, EC50)
  # id : vector of indices
  # xidep : dependent variables (same nb of rows as length of id)
  # returns:
  # a vector of predictions of length equal to length of id
  dose<-xidep[,1]
e0<-psi[id,1]
```r
emax <- psi[id, 2]
e50 <- psi[id, 3]
f <- e0 + emax * dose / (e50 + dose)
return(f)
```

```r
# Plotting the data
plot(saemix.data, main = "Simulated data PD1")

# Compare models with and without covariates with LL by Importance Sampling
model1 <- saemixModel(model = modelemax, description = "Emax growth model",
psi0 = matrix(c(20, 300, 20, 0, 0, 0), ncol = 3, byrow = TRUE, dimnames = list(NULL, c("E0", "Emax", "EC50"))), transform.par = c(1, 1, 1),
covariate.model = matrix(c(0, 0, 0), ncol = 3, byrow = TRUE), fixed.estim = c(1, 1, 1))

model2 <- saemixModel(model = modelemax, description = "Emax growth model",
psi0 = matrix(c(20, 300, 20, 0, 0, 0), ncol = 3, byrow = TRUE, dimnames = list(NULL, c("E0", "Emax", "EC50"))), transform.par = c(1, 1, 1),
covariate.model = matrix(c(0, 0, 1), ncol = 3, byrow = TRUE), fixed.estim = c(1, 1, 1))

# SE not computed as not needed for the test
saemix.options <- list(algorithms = c(0, 1, 1), nb.chains = 3, seed = 765754,
nbiter.saemix = c(500, 300), save = FALSE, save.graphs = FALSE, displayProgress = FALSE)

fit1 <- saemix(model1, saemix.data, saemix.options)

fit2 <- saemix(model2, saemix.data, saemix.options)

wstat <- (-2) * (fit1["results"]["ll.is"] - fit2["results"]["ll.is"])

cat("LRT test for covariate effect on EC50: p-value =", 1 - pchisq(wstat, 1), "\n")
```

---

**plot.SaemixModel,ANY-method**

*Plot model predictions using an SaemixModel object*

**Description**

This function will plot predictions obtained from an SaemixModel object over a given range of X. Additional predictors may be passed on to the function using the predictors argument.

**Usage**

```r
## S4 method for signature 'SaemixModel,ANY'
plot(x, y, range = c(0, 1), psi = NULL, predictors = NULL, ...)
```

**Arguments**

- **x** an SaemixData object or an SaemixSimData object
- **y** unused, present for compatibility with base plot function
range
range of X over which the model is to be plotted. Important note: the first predictor will be used for the X-axis, the other predictors when present need to be passed sequentially in the predictors argument, in the order in which they appear in the model. Less important note: please use explicitly range=XXX where XXX is of the form c(a,b) to pass the plotting range on the X-axis.

psi
parameters of the model

predictors
additional predictors needed to pass on to the model

... additional arguments to be passed on to plot (titles, legends, ...). Use verbose=TRUE to print some messages concerning the characteristics of the plot.

Examples

# Note that we have written the PK model so that time is the first predictor (xidep[,1])
# and dose the second
model1cpt<-function(psi,id,xidep) {
  tim<-xidep[,1]
dose<-xidep[,2]
  ka<psi[id,1]
  V<psi[id,2]
  CL<psi[id,3]
  k<-CL/V
  ypred<-(dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}
x<-saemixModel(model=model1cpt,description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.5,30,1), ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))))
# Plot the model over 0-24h, using the parameters given in psi0 and a dose of 300
plot(x, range=c(0,24), predictors=300, verbose=TRUE)
# Plot the model over 0-24h, using another set of parameters and a dose of 350
plot(x, range=c(0,24), psi=c(1.5,20,2), predictors=350, verbose=TRUE)
Arguments

- `x` an SaemixModel object
- `y` an SaemixData object
- `...` additional arguments. Passing psi=X where X is a vector or a dataframe will allow changing the parameters for which predictions are to be computed (defaults to the population parameters defined by the psi element of x) (see details)

Details

The function uses the model slot of the SaemixModel object to obtain predictions, using the dataset contained in the SaemixData object. The user is responsible for making sure data and model match.

If psi is not given, the predictions will be computed for the population parameters (first line of the psi0 slot) of the object. If psi is given, the number of columns in psi (or the number of elements of psi, if psi is given as a vector) should match the number of parameters in the model, otherwise an error message will be shown and the function will return empty. If psi is a dataframe, each line will be used for a separate subject of the smx.data object. Elements of psi will be recycled if psi has less lines than the number of subjects in the dataset.

Currently this function only works for models defined as `structural`.

Value

- a ggplot object

Examples

```r
data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
  name.group=c("Id"),name.predictors=c("Dose","Time"),
  name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
  units=list(x="hr",y="mg/L", covariates=c("kg","-") ), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}

saemix.model<-saemixModel(model=model1cpt,modeltype="structural",
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.0,20.0,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
  dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,1,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")
```
plot(saemix.model, saemix.data)
plot(saemix.model, saemix.data, psi=c(2, 40, 3))
indpsi<-data.frame(ka=2, V=seq(25,47,2), CL=seq(2.5,4.7, 0.2))
plot(saemix.model, saemix.data, psi=indpsi)

---

**plot, SaemixObject, ANY-method**

*General plot function from SAEM*

**Description**

Several plots (selectable by the type argument) are currently available: convergence plot, individual plots, predictions versus observations, distribution plots, VPC, residual plots, mirror.

**Usage**

```r
## S4 method for signature 'SaemixObject,ANY'
plot(x, y, ...)
```

**Arguments**

- **x**: an object returned by the `saemix` function
- **y**: empty
- **...**: optional arguments passed to the plots

**Details**

This is the generic plot function for an SaemixObject object, which implements different graphs related to the algorithm (convergence plots, likelihood estimation) as well as diagnostic graphs. A description is provided in the PDF documentation. Arguments such as main, xlab, etc... that can be given to the generic plot function may be used, and will be interpreted according to the type of plot that is to be drawn.

A special argument `plot.type` can be set to determine the type of plot; it can be one of:

- **data**: A spaghetti plot of the data, displaying the observed data `y` as a function of the regression variable (time for a PK application)
- **convergence**: For each parameter in the model, this plot shows the evolution of the parameter estimate versus the iteration number
- **likelihood**: Graph showing the evolution of the log-likelihood during the estimation by importance sampling
- **observations.vs.predictions**: Plot of the predictions computed with the population parameters versus the observations (left), and plot of the predictions computed with the individual parameters versus the observations (right)
**residuals.scatter:** Scatterplot of the residuals versus the predictor (top) and versus predictions (bottom), for weighted residuals (population residuals, left), individual weighted residuals (middle) and npde (right).

**residuals.distribution:** Distribution of the residuals, plotted as histogram (top) and as a QQ-plot (bottom), for weighted residuals (population residuals, left), individual weighted residuals (middle) and npde (right).

**individual.fit:** Individual fits are obtained using the individual parameters with the individual covariates

**population.fit:** Population fits are obtained using the population parameters with the individual covariates

**both.fit:** Individual fits, superposing fits obtained using the population parameters with the individual covariates (red) and using the individual parameters with the individual covariates (green)

**mirror:** Mirror plots assessing the compatibility of simulated data compared to the original

**marginal.distribution:** Distribution of the parameters (conditional on covariates when some are included in the model). A histogram of individual parameter estimates can be overlayed on the plot, but it should be noted that the histogram does not make sense when there are covariates influencing the parameters and a warning will be displayed

**random.effects:** Boxplot of the random effects

**correlations:** Correlation between the random effects

**parameters.vs.covariates:** Plots of the estimates of the individual parameters versus the covariates, using scatterplot for continuous covariates, boxplot for categorical covariates

**randeff.vs.covariates:** Plots of the estimates of the random effects versus the covariates, using scatterplot for continuous covariates, boxplot for categorical covariates

**npde:** Plots 4 graphs to evaluate the shape of the distribution of the normalised prediction distribution errors (npde)

**vpc:** Visual Predictive Check, with options to include the prediction intervals around the boundaries of the selected interval as well as around the median (50th percentile of the simulated data).

In addition, the following values for plot.type produce a series of plots:

**reduced:** produces the following plots: plot of the data, convergence plots, plot of the likelihood by importance sampling (if computed), plots of observations versus predictions. This is the default behaviour of the plot function applied to an SaemixObject object

**full:** produces the following plots: plot of the data, convergence plots, plot of the likelihood by importance sampling (if computed), plots of observations versus predictions, scatterplots and distribution of residuals, VPC, npde, boxplot of the random effects, distribution of the parameters, correlations between random effects, plots of the relationships between individually estimated parameters and covariates, plots of the relationships between individually estimated random effects and covariates

Each plot can be customised by modifying options, either through a list of options set by the `saemix.plot.setoptions` function, or on the fly by passing an option in the call to the plot (see examples).

**Value**

None
Author(s)
Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References

See Also
SaemixObject, saemix, saemix.plot.setoptions, saemix.plot.select, saemix.plot.data

Examples

data(theo.saemix)

saemix.data <- saemixData(name.data=theo.saemix, header=TRUE, sep=" ", na=NA, name.group=c("Id"), name.predictors=c("Dose", "Time"), name.response=c("Concentration"), name.covariates=c("Weight", "Sex"), units=list(x="hr", y="mg/L", covariates=c("kg", "-")), name.X="Time")

model1cpt <- function(psi, id, xidep) {
doese <- xidep[, 1]
dose <- dose[, 1]
tim <- xidep[, 2]
ka <- psi[id, 1]
V <- psi[id, 2]
CL <- psi[id, 3]
k <- CL/V
ypred <- dose * ka / (V * (ka - k)) * (exp(-k * tim) - exp(-ka * tim))
return(ypred)
}

saemix.model <- saemixModel(model=model1cpt,
description="One-compartment model with first-order absorption",
 psi0 = matrix(c(1, 20, 0.5, 0.1, 0, -0.01), ncol=3, byrow=TRUE,
dimnames=list(NULL, c("ka", "V", "CL"))),
transform.par = c(1, 1, 1),
covariate.model = matrix(c(0, 1, 0, 0, 0), ncol=3, byrow=TRUE),
fixed.estim = c(1, 1, 1),
covariance.model = matrix(c(1, 0, 0, 0, 1, 0, 0, 0, 1), ncol=3, byrow=TRUE),
omega.init = matrix(c(1, 0, 0, 1, 0, 0, 0, 1, 0), ncol=3, byrow=TRUE),
error.model = "constant")

saemix.options <- list(seed=632545, save=FALSE, save.graphs=FALSE)

# Not run (strict time constraints for CRAN)
# saemix.fit <- saemix(saemix.model, saemix.data, saemix.options)
plot-methods

# Set of default plots
# plot(saemix.fit)

# Data
# plot(saemix.fit, plot.type="data")

# Convergence
# plot(saemix.fit, plot.type="convergence")

# Individual plot for subject 1, smoothed
# plot(saemix.fit, plot.type="individual.fit", ilist=1, smooth=TRUE)

# Individual plot for subject 1 to 12, with ask set to TRUE
# (the system will pause before a new graph is produced)
# plot(saemix.fit, plot.type="individual.fit", ilist=c(1:12), ask=TRUE)

# Diagnostic plot: observations versus population predictions
# par(mfrow=c(1,1))
# plot(saemix.fit, plot.type="observations.vs.predictions", level=0, new=FALSE)

# LL by Importance Sampling
# plot(saemix.fit, plot.type="likelihood")

# Scatter plot of residuals
# Data will be simulated to compute weighted residuals and npde
# the results shall be silently added to the object saemix.fit
# plot(saemix.fit, plot.type="residuals.scatter")

# Boxplot of random effects
# plot(saemix.fit, plot.type="random.effects")

# Relationships between parameters and covariates
# plot(saemix.fit, plot.type="parameters.vs.covariates")

# Relationships between parameters and covariates, on the same page
# par(mfrow=c(3,2))
# plot(saemix.fit, plot.type="parameters.vs.covariates", new=FALSE)

# VPC
# Not run (time constraints for CRAN)
# plot(saemix.fit, plot.type="vpc")

plot-methods

Methods for Function plot

Description

Methods for function plot
Methods

- `signature(x = "ANY")` default plot function
- `signature(x = "SaemixData")` Plots the data. Defaults to a spaghetti plot of response versus predictor, with lines joining the data for one individual.
- `signature(x = "SaemixModel")` Plots prediction of the model
- `signature(x = "SaemixObject")` This method gives access to a number of plots that can be performed on a SaemixObject
- `signature(x = "SaemixSimData")` Plots simulated datasets

---

plot.SaemixData  
Plot of longitudinal data

Description

This function will plot a longitudinal dataframe contained in a SaemixData object. By default it produces a spaghetti plot, but arguments can be passed on to modify this behaviour.

Usage

```r
## S3 method for class 'SaemixData'
plot(x, y, ...)

## S3 method for class 'SaemixSimData'
plot(x, y, irep = -1, ...)
```

Arguments

- `x` an SaemixData object or an SaemixSimData object
- `y` unused, present for compatibility with base plot function
- `...` additional arguments to be passed on to plot (titles, legends, ...)
- `irep` which replicate datasets to use in the mirror plot (defaults to -1, causing a random simulated dataset to be sampled from the nsim simulated datasets)
**plotDiscreteData**

*Plot non Gaussian data*

**Description**

This function provides exploration plots for non Gaussian longitudinal data (work in progress)

**Usage**

```r
plotDiscreteData(object, outcome = "continuous", verbose = FALSE, ...)
```

**Arguments**

- **object**
  - an SaemixData object returned by the `saemixData` function.
- **outcome**
  - type of outcome (valid types are "TTE", "binary", "categorical", "count")
- **verbose**
  - whether to print messages (defaults to FALSE)
- **...**
  - additional arguments, used to pass graphical options (to be implemented, currently not available)

**Details**

This function is a very rough first attempt at automatically creating plots to explore discrete longitudinal data.

- for TTE data, a KM plot will be produced
- for count, categorical and binary data, a plot showing the proportion of each score/category across time will be shown. These plots can be stratified over a covariate in the data set (currently only categorical covariates) by passing an argument which.cov='name' to the call

**Author(s)**

Emmanuelle Comets emmanuelle.comets@inserm.fr

**References**


Ron Keizer, tutorials on VPC TODO

**See Also**

`SaemixObject`, `saemix`, `saemix.plot.vpc`, `simulateDiscreteSaemix`
Examples

# Time-to-event data
data(lung.saemix)

saemix.data<-saemixData(name.data=lung.saemix,header=TRUE,name.group=c("id"),
  name.predictors=c("time","status","cens"),name.response=c("status"),
  name.covariates=c("age","sex","ph.ecog","ph.karno","pat.karno","wt.loss","meal.cal"),
  units=list(x="days",y="",covariates=c("yr","%","%","cal","pounds")))

# Plots a KM survival plot
plotDiscreteData(saemix.data, outcome="TTE")
# Plots a KM survival plot, stratified by sex
plotDiscreteData(saemix.data, outcome="TTE", which.cov="sex")

# Count data
data(rapi.saemix)
saemix.data<-saemixData(name.data=rapi.saemix, name.group=c("id"),
  name.predictors=c("time","rapi"),name.response=c("rapi"),
  name.covariates=c("gender"),units=list(x="months",y="",covariates=c("")))

# Plots a histogram of the counts
plotDiscreteData(saemix.data, outcome="count")

predict-methods

Methods for Function predict

Description

Methods for function predict

Usage

## S4 method for signature 'SaemixObject'
predict(
  object,
  newdata = NULL,
  type = c("ipred", "ypred", "ppred", "icpred"),
  se.fit = FALSE,
  ...
)

Arguments

  object       an SaemixObject
  newdata      an optional dataframe for which predictions are desired. If newdata is given, it must contain the predictors needed for the model in object
predict.SaemixModel

<table>
<thead>
<tr>
<th>type</th>
<th>the type of predictions (ipred= individual, ppred=population predictions obtained with the population estimates, ypred=mean of the population predictions, icpred=conditional predictions). With newdata, individual parameters can be estimated if the new data contains observations; otherwise, predictions correspond to the population predictions ppred, and type is ignored.</th>
</tr>
</thead>
<tbody>
<tr>
<td>se.fit</td>
<td>whether the SE are to be taken into account in the model predictions</td>
</tr>
<tr>
<td>...</td>
<td>additional arguments passed on to fitted()</td>
</tr>
</tbody>
</table>

**Value**

a vector or a dataframe (if more than one type) with the corresponding predictions for each observation in the dataframe

**Methods**

- list("signature(object = \"ANY\")")  Default predict functions
- list("signature(object = \"SaemixObject\")")  Computes predictions using the results of an SAEM fit

**predict.SaemixModel**  
Predictions for a new dataset

**Description**

Predictions for a new dataset

**Usage**

```r
## S3 method for class 'SaemixModel'
predict(object, predictors, psi = c(), id = c(), ...)
```

**Arguments**

- `object` an SaemixModel object
- `predictors` a dataframe with the predictors for the model (must correspond to the predictors used by the model function)
- `psi` a vector or a dataframe giving the parameters for which predictions are to be computed (defaults to empty). The number of columns in psi (or the number of elements of psi, if psi is given as a vector) should match the number of parameters in the model, otherwise an error message will be shown and the function will return empty. If psi is NA, the predictions are computed for the population parameters in the model (first line of the psi0 slot). If psi is a dataframe, each line will be used for a separate 'subject' in the predictors dataframe, as indicated by the id argument; if id is not given, only the first line of psi will be used.
id

id is a vector of indices of length equal to the number of lines in predictors, matching each line of predictors to the corresponding line in psi, i.e., the parameters for this predictors (defaults to empty). If id is given, the unique values in id must be equal to the number of lines in psi, otherwise id will be set to 1. If id is given and its values do not take the consecutive values 1:N, the indices will be matched to 1:N to follow the lines in psi.

... unused argument, for consistency with the generic

Details

The function uses the model slot of the SaemixModel object to obtain predictions, using the predictors object. The user is responsible for giving all the predictors needed by the model function. If psi is not given, the predictions will be computed for the population parameters (first line of the psi0 slot) of the object.

The predictions correspond to the structure of the model; for models defined in terms of their likelihood, the predictions are the log-pdf of the model (see documentation for details).

Warning: this function is currently under development and the output may change in future versions of the package to conform to the usual predict functions.

Value

a list with two components

- param a dataframe with the estimated parameters
- predictions a dataframe with the population predictions

Examples

data(theo.saemix)
xpred<-theo.saemix[,c("Dose","Time")]

model1cpt<--function(psi,id,xidep) {
dose<-xidep[,1]
tim<-xidep[,2]
ka<-psi[id,1]
V<-psi[id,2]
CL<-psi[id,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
return(ypred)
}

saemix.model<--saemixModel(model=model1cpt,modeltype="structural",
description="One-compartment model with first-order absorption",
psi0=matrix(c(1,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
covariance.model=matrix(c(1,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
omega.init=matrix(c(1,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")
```r
head(predict(saemix.model, xpred)$predictions)
head(predict(saemix.model, xpred, psi=c(2, 40, 0.5))$predictions)
indpsi<-data.frame(ka=2, V=seq(25,47,2), CL=seq(2.5,4.7, 0.2))
head(predict(saemix.model, xpred, psi=indpsi)$predictions)
```

### print-methods

#### Methods for Function `print`

**Description**

Prints a summary of an object

**Usage**

```r
## S4 method for signature 'SaemixData'
print(x, nlines = 10, ...)

## S4 method for signature 'SaemixModel'
print(x, ...)

## S4 method for signature 'SaemixRes'
print(x, digits = 2, map = FALSE, ...)

## S4 method for signature 'SaemixObject'
print(x, nlines = 10, ...)
```

#### Arguments

- `x`:
  - an object of type `SaemixData`, `SaemixModel`, `SaemixRes` or `SaemixObject`
- `nlines`:
  - maximum number of lines of data to print (defaults to 10)
- `...`:
  - additional arguments passed on the print function
- `digits`:
  - number of digits to use for pretty printing
- `map`:
  - when `map` is TRUE the individual parameter estimates are shown (defaults to FALSE)

#### Methods

- `list("signature(x ="ANY")")`  Default print function
- `list("signature(x ="SaemixData")")`  Prints a summary of a `SaemixData` object
- `list("signature(x ="SaemixModel")")`  Prints a summary of a `SaemixModel` object
- `list("signature(x ="SaemixObject")")`  Prints a summary of the results from a SAEMIX fit
- `list("signature(x ="SaemixRes")")`  Not user-level
Description

These three functions are used to access the estimates of individual parameters and random effects.

Usage

\[ \text{psi}(\text{object}, \text{type} = \text{c("mode", "mean")}) \]
\[ \text{phi}(\text{object}, \text{type} = \text{c("mode", "mean")}) \]
\[ \text{eta}(\text{object}, \text{type} = \text{c("mode", "mean")}) \]

## S4 method for signature 'SaemixObject'
\[ \text{psi}(\text{object}, \text{type} = \text{c("mode", "mean")}) \]
## S4 method for signature 'SaemixObject'
\[ \text{phi}(\text{object}, \text{type} = \text{c("mode", "mean")}) \]
## S4 method for signature 'SaemixObject'
\[ \text{eta}(\text{object}, \text{type} = \text{c("mode", "mean")}) \]

Arguments

- \text{object} an SaemixObject object returned by the \text{saemix} function
- \text{type} a string specifying whether to use the MAP (type="mode") or the mean (type="mean") of the conditional distribution of the individual parameters. Defaults to mode

Details

The psi_i represent the individual parameter estimates. In the SAEM algorithm, these parameters are assumed to be a transformation of a Gaussian random vector phi_i, where the phi_i can be written as a function of the individual random effects (eta_i), the covariate matrix (C_i) and the vector of fixed effects (mu):
\[ \text{phi_i} = C_i \mu + \text{eta}_i \]

More details can be found in the PDF documentation.

Value

a matrix with the individual parameters (psi/phi) or the random effects (eta). These functions are used to access and output the estimates of parameters and random effects. When the object passed to the function does not contain these estimates, they are automatically computed. The object is then returned (invisibly) with these estimates added to the results.
Methods

```r
list("signature(object = "SaemixObject")")
```
please refer to the PDF documentation for the models

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

`SaemixData, SaemixModel, SaemixObject, saemixControl, plot.saemix`

Examples

data(theo.saemix)

```r
saemix.data<-saemixData(name.data=theo.saemix, header=TRUE, sep=" ", na=NA, name.group=c("Id"), name.predictors=c("Dose","Time"), name.response=c("Concentration"), name.covariates=c("Weight","Sex"), units=list(x="hr", y="mg/L", covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}

saemix.model<-saemixModel(model=model1cpt, description="One-compartment model with first-order absorption", psi0=matrix(c(1.,20,0.5,0.1,0,-0.01), ncol=3, byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))), transform.par=c(1,1,1), covariate.model=matrix(c(0,1,0,0,0), ncol=3, byrow=TRUE), fixed.estim=c(1,1,1), covariance.model=matrix(c(1,0,0,0,1,0,0,0,1), ncol=3, byrow=TRUE), omega.init=matrix(c(1,0,0,0,1,0,0,0,1), ncol=3, byrow=TRUE), error.model="constant")
```
saemix.options<-list(algorithm=c(1,0,0),seed=632545,save=FALSE,save.graphs=FALSE,displayProgress=FALSE)

# Not run (strict time constraints for CRAN)

saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)
psi(saemix.fit)
phi(saemix.fit)
eta(saemix.fit,type="mean")

---

rapi.saemix

**Rutgers Alcohol Problem Index**

**Description**

The RAPI data studies gender differences across two years in alcohol-related problems, as measured by the Rutgers Alcohol Problem Index (RAPI; White & Labouvie, 1989). The dataset includes 3,616 repeated measures of counts representing the number of alcohol problems reported over six months period, across five time points from 818 individuals.

**Format**

This data frame contains the following columns:

- **id**: subject identification number
- **time**: time since the beginning of the study (months)
- **rapi**: the number of reported alcohol problems during a six-months period
- **gender**: gender (0=Men, 1=Women)

**Source**

David Atkins, University of Washington

**References**


Examples

data(rapi.saemix)
saemix.data<-saemixData(name.data=rapi.saemix, name.group=c("id"),
 name.predictors=c("time","rapi"),name.response=c("rapi"),
 name.covariates=c("gender"),units=list(x="months",y="",covariates=c("")))
hist(rapi.saemix$rapi, main="", xlab="RAPI score", breaks=30)

# Fitting a Poisson model
count.poisson<-function(psi,id,xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  intercept<-psi[id,1]
  slope<-psi[id,2]
  lambda<- exp(intercept + slope*time)
  logp <- -lambda + y*log(lambda) - log(factorial(y))
  return(logp)
}
countsimulate.poisson<-function(psi, id, xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  ymax<-max(y)
  intercept<-psi[id,1]
  slope<-psi[id,2]
  lambda<- exp(intercept + slope*time)
  y<-rpois(length(time), lambda=lambda)
  y[y>ymax]<-ymax+1 # truncate to maximum observed value to avoid simulating aberrant values
  return(y)
}

# Gender effect on intercept and slope
rapimod.poisson<-saemixModel(model=count.poisson, simulate.function=countsimulate.poisson,
 description="Count model Poisson",modeltype="likelihood",
 psi=matrix(c(log(5),0.01),ncol=2,byrow=TRUE,dimnames=list(NULL, c("intercept","slope")),
 transform.par=c(0,0), omega.init=diag(c(0.5, 0.5)),
 covariance.model =matrix(data=1, ncol=2, nrow=2),
 covariate.model=matrix(c(1,1), ncol=2, byrow=TRUE))
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE, fim=FALSE)
poisson.fit<-saemix(rapimod.poisson,saemix.data,saemix.options)

# Fitting a ZIP model
count.poissonzip<-function(psi, id, xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  intercept<-psi[id,1]
  slope<-psi[id,2]
  p0<-psi[id,3] # Probability of zero's
  lambda<- exp(intercept + slope*time)
  logp <- -log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson
  logp0 <- -log(p0+(1-p0)*exp(-lambda)) # Zeroes
  logp[y==0]<-logp0[y==0]
  return(logp)
}
counts\_simulate.poissonzip<-function(psi, id, xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  ymax<-max(y)
  intercept<-psi[id,1]
  slope<-psi[id,2]
  p0<-psi[id,3] # Probability of zero's
  lambda<- exp(intercept + slope*time)
  prob0<-rbinom(length(time), size=1, prob=p0)
  y<-rpois(length(time), lambda=lambda)
  y[prob0==1]<-0
  y[y>ymax]<-ymax+1 # truncate to maximum observed value to avoid simulating aberrant values
  return(y)
}

rapimod.zip<-saemixModel(model=count.poissonzip, simulate.function=counts\_simulate.poissonzip, 
  description="count model ZIP", modeltype="likelihood", 
  psi0=matrix(c(1.5, 0.01, 0.2),ncol=3,byrow=TRUE, 
  dimnames=list(NULL, c("intercept", "slope","p0"))), 
  transform.par=c(0,0,3), covariance.model=diag(c(1,1,0)), omega.init=diag(c(0.5,0.3,0)), 
  covariate.model = matrix(c(1,1,0),ncol=3, byrow=TRUE))

zippoisson.fit<-saemix(rapimod.zip,saemix.data,saemix.options)

# Using simulations to compare the predicted proportion of 0's in the two models
nsim<-100
yfit1<-simulateDiscreteSaemix(poisson.fit, nsim=nsim)
yfit2<-simulateDiscreteSaemix(zippoisson.fit, nsim=100)

nobssim=length(yfit1@sim.data@datasim$ysim)
cat("Observed proportion of 0's",
  length(yfit1@data@data$rapi[yfit1@data@data$rapi==0])/yfit1@data@ntot.obs,"\n")
cat(" Poisson model, p=",
  length(yfit1@sim.data@datasim$ysim[yfit1@sim.data@datasim$ysim==0])/nobssim,"\n")
cat(" ZIP model, p=",
  length(yfit2@sim.data@datasim$ysim[yfit2@sim.data@datasim$ysim==0])/nobssim,"\n")

---

readSaemix, SaemixData-method

Create a longitudinal data structure from a file or a dataframe
Helper function not intended to be called by the user

Description

Create a longitudinal data structure from a file or a dataframe
Helper function not intended to be called by the user
Usage

```r
## S4 method for signature 'SaemixData'
readSaemix(object, dat = NULL)
```

Arguments

- `object`: an SaemixData object
- `dat`: the name of a dataframe in the R environment, defaults to NULL; if NULL, the function will attempt to read the file defined by the slot name.data.

Description

Returns an SaemixObject object where the data object has been replaced by the data provided in a dataframe.

Usage

```r
replaceData.saemixObject(saemixObject, newdata)
```

Arguments

- `saemixObject`: an SaemixObject object
- `newdata`: a dataframe containing data

Value

An object of class "SaemixObject". The population parameters are retained but all the predictions, individual parameters and statistical criteria are removed. The function attempts to extract the elements entering the statistical model (subject id, predictors, covariates and response).

Examples

```r
# TODO
```
resid.saemix  

**Extract Model Residuals**

**Description**

residuals is a generic function which extracts model residuals from objects returned by modelling functions. The abbreviated form resid is an alias for residuals.

**Usage**

```r
## S3 method for class 'SaemixRes'
resid(object, type = c("ires", "wres", "npde", "pd", "iwres", "icwres"), ...)

## S3 method for class 'SaemixObject'
resid(object, type = c("ires", "wres", "npde", "pd", "iwres", "icwres"), ...)
```

**Arguments**

- **object**: an SaemixRes or an SaemixObject object
- **type**: string determining which residuals are extracted. Possible values are: "ires" (individual residuals, default), "wres" (weighted population residuals), "npde" (normalised prediction distribution errors), "pd" (prediction discrepancies), "iwres" (individual weighted residuals) and "icwres" (conditional individual weighted residuals). See user guide for details.
- **...**: further arguments to be passed to or from other methods

**Value**

By default, individual residuals are extracted from the model object.

---

**saemix**  

**Stochastic Approximation Expectation Maximization (SAEM) algorithm**

**Description**

SAEM algorithm perform parameter estimation for nonlinear mixed effects models without any approximation of the model (linearization, quadrature approximation, ...)

**Usage**

```r
saemix(model, data, control = list())
```
Arguments

model  an object of class SaemixModel, created by a call to the function saemixModel

data  an object of class SaemixData, created by a call to the function saemixData

control  a list of options, see saemixControl

Details

The SAEM algorithm is a stochastic approximation version of the standard EM algorithm proposed by Kuhn and Lavielle (see reference). Details of the algorithm can be found in the pdf file accompanying the package.

Value

An object of class SaemixObject containing the results of the fit of the data by the non-linear mixed effect model. A summary of the results is printed out to the terminal, and, provided the appropriate options have not been changed, numerical and graphical outputs are saved in a directory.

Author(s)

Emmanuelle Comets <emmanuelle.comets@inserm.fr>
Audrey Lavenu
Marc Lavielle

References


See Also

SaemixData, SaemixModel, SaemixObject, saemixControl, plot.saemix

Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
name.group=c("Id"),name.predictors=c("Dose","Time"),
name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
units=list(x="hr",y="mg/L", covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {


dose<-xidep[,1]
tim<-xidep[,2]
ka<-psi[id,1]
V<-psi[id,2]
CL<-psi[id,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
return(ypred)
}
saemix.model<-saemixModel(model=model1cpt,modeltype="structural",
description="One-compartment model with first-order absorption",
psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
covariance.model=matrix(c(1,0,0,1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
omega.init=matrix(c(1,0,0,1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")

# Not run (strict time constraints for CRAN)
saemix.fit<-saemix(saemix.model,saemix.data,list(seed=632545,directory="newtheo",
save=FALSE,save.graphs=FALSE, print=FALSE))

# Prints a summary of the results
print(saemix.fit)

# Outputs the estimates of individual parameters
psi(saemix.fit)

# Shows some diagnostic plots to evaluate the fit
plot(saemix.fit)

saemix.bootstrap Bootstrap for saemix fits

Description

This function provides bootstrap estimates for a saemixObject run. Different bootstrap approaches have been implemented (see below for details), with the default method being the conditional non-parametric bootstrap ()

Usage

saemix.bootstrap(
saemixObject,
method = "conditional",
nboot = 200,
nsamp = 100,
saemix.options = NULL
)
Arguments

saemixObject    an object returned by the saemix function
method          the name of the bootstrap algorithm to use (one of: "case", "residual", "parametric" or "conditional") (defaults to "conditional")
nboot           number of bootstrap samples
nsamp           number of samples from the conditional distribution (for method="conditional")

saemix.options  list of options to run the saemix algorithm. Defaults to the options in the object, but suppressing the estimation of individual parameters (map=FALSE) and likelihood (ll.is=FALSE), and the options to print out intermediate results and graphs (displayProgress=FALSE,save.graphs=FALSE,print=FALSE)

Details

Different bootstrap algorithms have been proposed for non-linear mixed effect models, to account for the hierarchical nature of these models (see review inThai et al. 2013, 2014) In saemix, we implemented the following bootstrap algorithms, which can be selected using the "method" argument:

- case refers to case bootstrap, where resampling is performed at the level of the individual and the bootstrap sample consists in sampling with replacement from the observed individuals
- residual refers to non-parametric residual bootstrap, where for each individual the residuals for the random effects and for the residual error are resampled from the individual estimates after the fit
- parametric refers to parametric residual bootstrap, where these residuals are sampled from their theoretical distributions. In saemix, random effects are drawn from a normal distribution using the estimated variance in saemixObject, and transformed to the distribution specified by the transform.par component of the model, and the residual error is drawn from a normal distribution using the estimated residual variance.
- conditional refers to the conditional non-parametric bootstrap, where the random effects are resampled from the individual conditional distributions as in (Comets et al. 2021) and the residual errors resampled from the corresponding residuals

Important note: for discrete data models, all residual-based bootstraps (residual, parametric and conditional) need a simulate.function slot to be included in the model object, as the algorithm will need to generate predictions with the resampled individual parameters in order to generate bootstrap samples. See SaemixModel and the examples provided as notebooks for details.

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr

References


Examples

# Bootstrap for the theophylline data
data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
name.group=c("Id"),name.predictors=c("Dose","Time"),
name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
dose<-xidep[,1]
tim<-xidep[,2]
ka<-psi[id,1]
V<-psi[id,2]
CL<-psi[id,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
return(ypred)
}

saemix.model<-saemixModel(model=model1cpt,
description="One-compartment model with first-order absorption",
psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")

saemix.options<-list(algorithm=c(1,0,0),seed=632545,save=FALSE,save.graphs=FALSE,
displayProgress=FALSE)

# Not run (strict time constraints for CRAN)

saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

# Only 10 bootstrap samples here for speed, please increase to at least 200
theo.case <- saemix.bootstrap(saemix.fit, method="case", nboot=10)
theo.cond <- saemix.bootstrap(saemix.fit, nboot=10)

# Bootstrap for the toenail data
data(toenail.saemix)

saemix.data<-saemixData(name.data=toenail.saemix,name.group=c("id"), name.predictors=c("time","y"),
name.response="y", name.covariates=c("treatment"),name.X=c("time"))

binary.model<-function(psi,id,xidep) {
tim<-xidep[,1]
y<-xidep[,2]
inter<-psi[id,1]
slopes <- psi[,2]
logit <- inter + slopes * tim
pevent <- exp(logit) / (1 + exp(logit))
pobs = (y == 0) * (1 - pevent) + (y == 1) * pevent
logpdf <- log(pobs)
return(logpdf)

simulBinary <- function(psi, id, xidep) {
  tim <- xidep[, 1]
y <- xidep[, 2]
inter <- psi[id, 1]
slope <- psi[id, 2]
logit <- inter + slopes * tim
pevent <- 1 / (1 + exp(-logit))
ysim <- rbinom(length(tim), size = 1, prob = pevent)
return(ysim)
}

saemix.model <- saemixModel(model = binary.model, description = "Binary model",
modeltype = "likelihood", simulate.function = simulBinary,
psi0 = matrix(c(-5, -0.1, 0, 0), ncol = 2, byrow = TRUE, dimnames = list(NULL, c("inter", "slope"))),
transform.par = c(0, 0))

saemix.options <- list(seed = 1234567, save = FALSE, save.graphs = FALSE, displayProgress = FALSE,
nb.chains = 10, fim = FALSE)
binary.fit <- saemix(saemix.model, saemix.data, saemix.options)
# Only 10 bootstrap samples here for speed, please increase to at least 200
toenail.case <- saemix.bootstrap(binary.fit, method = "case", nboot = 10)
toenail.cond <- saemix.bootstrap(binary.fit, nboot = 10)

saemix.plot.data  

Functions implementing each type of plot in SAEM

Description

Several plots (selectable by the type argument) are currently available: convergence plot, individual plots, predictions versus observations, distribution plots, VPC, residual plots, and mirror plots.

Usage

saemix.plot.data(saemixObject, ...)

Arguments

saemixObject an object returned by the saemix function
... optional arguments passed to the plots
Details

These functions implement plots different graphs related to the algorithm (convergence plots, likelihood estimation) as well as diagnostic graphs. A description is provided in the PDF documentation. saemix.plot.parcov.aux, compute.sres and compute.eta.map are helper functions, not intended to be called by the user directly.

By default, the following plots are produced:

**saemix.plot.data:** A spaghetti plot of the data, displaying the observed data y as a function of the regression variable (time for a PK application)

**saemix.plot.convergence:** For each parameter in the model, this plot shows the evolution of the parameter estimate versus the iteration number

**saemix.plot.llis:** Graph showing the evolution of the log-likelihood during the estimation by importance sampling

**saemix.plot.obsvspred:** Plot of the predictions computed with the population parameters versus the observations (left), and plot of the predictions computed with the individual parameters versus the observations (right)

**saemix.plot.scatterresiduals:** Scatterplot of the residuals versus the predictor (top) and versus predictions (bottom), for weighted residuals (population residuals, left), individual weighted residuals (middle) and npde (right).

**saemix.plot.distribresiduals:** Distribution of the residuals, plotted as histogram (top) and as a QQ-plot (bottom), for weighted residuals (population residuals, left), individual weighted residuals (middle) and npde (right).

**saemix.plot.fits:** Model fits. Individual fits are obtained using the individual parameters with the individual covariates. Population fits are obtained using the population parameters with the individual covariates (red) and the individual parameters with the individual covariates (green). By default the individual plots are displayed.

**saemix.plot.distpsi:** Distribution of the parameters (conditional on covariates when some are included in the model). A histogram of individual parameter estimates can be overlayed on the plot, but it should be noted that the histogram does not make sense when there are covariates influencing the parameters and a warning will be displayed

**saemix.plot.randeff:** Boxplot of the random effects

**saemix.plot.correlations:** Correlation between the random effects

**saemix.plot.parcov:** Plots of the estimates of the individual parameters versus the covariates, using scatterplot for continuous covariates, boxplot for categorical covariates

**saemix.plot.randeffcov:** Plots of the estimates of the random effects versus the covariates, using scatterplot for continuous covariates, boxplot for categorical covariates

**saemix.plot.npde:** Plots 4 graphs to evaluate the shape of the distribution of the normalised prediction distribution errors (npde)

**saemix.plot.vpc:** Visual Predictive Check, with options to include the prediction intervals around the boundaries of the selected interval as well as around the median (50th percentile of the simulated data). Several methods are available to define binning on the X-axis (see methods in the PDF guide).
**saemix.plot.data**

*saemix.plot.data* When simulated data is available in the object (component sim.dat, which can be filled by a call to `simulate`), this function plots the original data as spaghetti plot and compares it to several simulated datasets under the fitted model.

Each plot can be customised by modifying options, either through a list of options set by the `saemix.plot.setoptions` function, or on the fly by passing an option in the call to the plot (see examples).

**Value**

None

**Author(s)**

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

**References**


**See Also**

`SaemixObject,saemix,saemix.plot.setoptions,saemix.plot.select,plot.saemix`

**Examples**

data(theo.saemix)

```r
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA, name.group=c("Id"),name.predictors=c("Dose","Time"), name.response=c("Concentration"),name.covariates=c("Weight","Sex"), units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

doellcpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}
```
saemix.model <- saemixModel(model=model1cpt,
  description = "One-compartment model with first-order absorption",
  psi0 = matrix(c(1.,20.,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
    dimnames = list(NULL, c("ka","V","CL"))),
  transform.par = c(1,1,1),
  covariate.model = matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),
  fixed.estim = c(1,1,1),
  covariance.model = matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init = matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  error.model = "constant")

saemix.options <- list(seed=632545,save=FALSE,save.graphs=FALSE)

# Not run (strict time constraints for CRAN)
saemix.fit <- saemix(saemix.model,saemix.data,saemix.options)

# Simulate data and compute weighted residuals and npde
saemix.fit <- compute.sres(saemix.fit)

# Data
saemix.plot.data(saemix.fit)

# Convergence
saemix.plot.convergence(saemix.fit)

# Individual plot for subject 1, smoothed
saemix.plot.fits(saemix.fit,ilist=1,smooth=TRUE)

# Individual plot for subject 1 to 12, with ask set to TRUE
# (the system will pause before a new graph is produced)
saemix.plot.fits(saemix.fit,ilist=c(1:12),ask=TRUE)

# Mirror plots (plots of simulated data compared to the original)
saemix.plot.mirror(saemix.fit)

# Diagnostic plot: observations versus population predictions
par(mfrow=c(1,1))
saemix.plot.obsvspred(saemix.fit,level=0,new=FALSE)

# LL by Importance Sampling
saemix.plot.llis(saemix.fit)

# Boxplot of random effects
saemix.plot.randeff(saemix.fit)

# Relationships between parameters and covariates
saemix.plot.parcov(saemix.fit)

# Relationships between parameters and covariates, on the same page
par(mfrow=c(3,2))
saemix.plot.parcov(saemix.fit,new=FALSE)

# Scatter plot of residuals
# Not run
# Works interactively but not in the contained environment of CRAN (it looks for a datafile
# instead of finding the dataset in the environment)
# saemix.plot.select

Plots of the results obtained by SAEM

**Description**

Several plots (selectable by the type argument) are currently available: convergence plot, individual plots, predictions versus observations, distribution plots, residual plots, VPC.

**Usage**

```r
saemix.plot.select(
  saemixObject,
  data = FALSE,
  convergence = FALSE,
  likelihood = FALSE,
  individual.fit = FALSE,
  population.fit = FALSE,
  both.fit = FALSE,
  observations.vs.predictions = FALSE,
  residuals.scatter = FALSE,
  residuals.distribution = FALSE,
  random.effects = FALSE,
  correlations = FALSE,
  parameters.vs.covariates = FALSE,
  randeff.vs.covariates = FALSE,
  marginal.distribution = FALSE,
  vpc = FALSE,
  npde = FALSE,
  ...
)
```

**Arguments**

- `saemixObject`: an object returned by the `saemix` function
- `data`: if TRUE, produce a plot of the data. Defaults to FALSE
convergence if TRUE, produce a convergence plot. Defaults to FALSE
likelihood if TRUE, produce a plot of the estimation of the LL by importance sampling. Defaults to FALSE
individual.fit if TRUE, produce individual fits with individual estimates. Defaults to FALSE
population.fit if TRUE, produce individual fits with population estimates. Defaults to FALSE
both.fit if TRUE, produce individual fits with both individual and population estimates. Defaults to FALSE
observations.vs.predictions if TRUE, produce a plot of observations versus predictions. Defaults to FALSE
residuals.scatter if TRUE, produce scatterplots of residuals versus predictor and predictions. Defaults to FALSE
residuals.distribution if TRUE, produce plots of the distribution of residuals. Defaults to FALSE
random.effects if TRUE, produce boxplots of the random effects. Defaults to FALSE
correlations if TRUE, produce a matrix plot showing the correlation between random effects. Defaults to FALSE
parameters.vs.covariates if TRUE, produce plots of the relationships between parameters and covariates, using the Empirical Bayes Estimates of individual parameters. Defaults to FALSE
randeff.vs.covariates if TRUE, produce plots of the relationships between random effects and covariates, using the Empirical Bayes Estimates of individual random effects. Defaults to FALSE
marginal.distribution if TRUE, produce plots of the marginal distribution of the random effects. Defaults to FALSE
vpc if TRUE, produce Visual Predictive Check plots. Defaults to FALSE (we suggest to use npdeSaemix instead)
npde if TRUE, produce plots of the npde. Defaults to FALSE (deprecated in 3.0, please use npdeSaemix instead)
... optional arguments passed to the plots

Details

This function plots different graphs related to the algorithm (convergence plots, likelihood estimation) as well as diagnostic graphs. A description is provided in the PDF documentation.

data A spaghetti plot of the data, displaying the observed data y as a function of the regression variable (eg time for a PK application)

convergence For each parameter in the model, this plot shows the evolution of the parameter estimate versus the iteration number

likelihood Estimation of the likelihood estimated by importance sampling, as a function of the number of MCMC samples
individual.fit  Individual fits, using the individual parameters with the individual covariates
population.fit  Individual fits, using the population parameters with the individual covariates
both.fit  Individual fits, using the population parameters with the individual covariates and the individual parameters with the individual covariates
observations.vs.predictions  Plot of the predictions computed with the population parameters versus the observations (left), and plot of the predictions computed with the individual parameters versus the observations (right)
residuals.scatter  Scatterplot of standardised residuals versus the X predictor and versus predictions. These plots are shown for individual and population residuals, as well as for npde when they are available
residuals.distribution  Distribution of standardised residuals, using histograms and QQ-plot. These plots are shown for individual and population residuals, as well as for npde when they are available
random.effects  Boxplot of the random effects
correlations  Correlation between the random effects, with a smoothing spline
parameters-versus.covariates  Plots of the estimate of the individual parameters versus the covariates, using scatterplot for continuous covariates, boxplot for categorical covariates
randeff-versus.covariates  Plots of the estimate of the individual random effects versus the covariates, using scatterplot for continuous covariates, boxplot for categorical covariates
marginal.distribution  Distribution of each parameter in the model (conditional on covariates when some are included in the model)
npde  Plot of npde as in package npde (deprecated in 3.0, please use npdeSaemix instead)
vpc  Visual Predictive Check (we suggest to use npdeSaemix instead)

Value
None

Author(s)
Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References
saemix.plot.setoptions

Function setting the default options for the plots in SAEM

See Also

SaemixObject, saemix.default, saemix.plots, saemix.plot.setoptions, saemix.plot.data, saemix.plot.convergence, saemix.plot.llis, saemix.plot.randeff, saemix.plot.obsvspred, saemix.plot.fits, saemix.plot.parcov, saemix.plot.randeffcov, saemix.plot.distpsi, npdeSaemix saemix.plot.scatterresiduals, saemix.plot.distribresiduals, saemix.plot.vpc

Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix, header=TRUE, sep=" ", na=NA, name.group=c("Id"), name.predictors=c("Dose","Time"), name.response=c("Concentration"), name.covariates=c("Weight","Sex"), units=list(x="hr", y="mg/L", covariates=c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k)) * (exp(-k*tim) - exp(-ka*tim))
  return(ypred)
}

saemix.model<-saemixModel(model=model1cpt,
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.,20,0.5,0.1,0.5,0.1),ncol=3, byrow=TRUE,
  dimnames=list(NULL, c("ka","V","CL"))), transform.par=c(1,1,1),
  covariate.model=matrix(c(0,1,0,0,0,0),ncol=3,byrow=TRUE),
  fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  error.model="constant")

saemix.options<-list(seed=632545, save=FALSE, save.graphs=FALSE)

# Not run (strict time constraints for CRAN)
# saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

# saemix.plot.select(saemix.fit,data=TRUE,main="Spaghetti plot of data")

# Putting several graphs on the same plot
# par(mfrow=c(2,2))
# saemix.plot.select(saemix.fit,data=TRUE,vpc=TRUE,observations.vs.predictions=TRUE, new=FALSE)
Description

- ablinecol Color of the lines added to the plots (default: "DarkRed")
- ablinelty Type of the lines added to the plots. Defaults to 2 (dashed line)
- ablinelwd Width of the lines added to the plots (default: 2)
- ask A logical value. If TRUE, users will be prompted before each new plot. Defaults to FALSE
- cex A numerical value giving the amount by which plotting text and symbols should be magnified relative to the default. Defaults to 1 (no magnification)
- cex.axis Magnification to be used for axis annotation relative to the current setting of 'cex'. Defaults to 1 (no magnification)
- cex.main Magnification to be used for main titles relative to the current setting of 'cex'. Defaults to 1 (no magnification)
- cex.lab Magnification to be used for x and y labels relative to the current setting of 'cex'. Defaults to 1 (no magnification)
- col.fillmed For the VPC plots: color filling the prediction interval for the median. Defaults to "pink"
- col.fillpi For the VPC plots: color filling the prediction interval for the limits of the prediction interval. Defaults to "slategray1"
- col.lmed For the VPC plots: color of the line showing the median of the simulated data. Defaults to "indianred4"
- col.lobs For the VPC plots: color of the lines showing the median, 2.5 and 97.5th percentiles (for a 95% prediction interval). Defaults to "steelblue4"
- col.lpi For the VPC plots: color of the line showing the boundaries of the prediction intervals. Defaults to "slategray4"
- col.obs For the VPC plots: color used to plot the observations. Defaults to "steelblue4"
- cov.name Name of the covariate to be used in the plots. Defaults to the first covariate in the model
- cov.value Value of the covariate to be used in the plots. Defaults to NA, indicating that the median value of the covariate (for continuous covariates) or the reference category (for categorical covariates) will be used
- ilist List of indices of subjects to be included in the individual plots (defaults to all subjects)
- indiv.par a string, giving the type of the individual estimates ("map"= conditional mode, "eap"=conditional mean). Defaults to conditional mode
- lcol Main line color (default: black)
- line.smooth Type of smoothing when a smoothed line is used in the plot ("m": mean value, "l": linear regression; "s": natural splines). Several options may be combined, for instance "ls" will add both a linear regression line and a line representing the fit of a natural spline. Defaults to "s"
- lty Line type. Defaults to 1, corresponding to a straight line
- lty.lmed For the VPC plots: type of the line showing the median of the simulated data. Defaults to 2 (dashed)
• lty.obs For the VPC plots: type of the line showing the observed data. Defaults to 1
• lty.lpi For the VPC plots: type of the line showing the boundaries of the simulated data. Defaults to 2 (dashed)
• lwd Line width (default: 1)
• lwd.lmed For the VPC plots: thickness of the line showing the median of the simulated data. Defaults to 2
• lwd.obs For the VPC plots: thickness of the line showing the median and boundaries of the observed data. Defaults to 2
• lwd.lpi For the VPC plots: thickness of the line showing the boundaries of the simulated data. Defaults to 1
• par.name Name of the parameter to be used in the plots. Defaults to the first parameter in the model
• pch Symbol type. Defaults to 20, corresponding to small dots
• pcol Main symbol color (default: black)
• range Range (expressed in number of SD) over which to plot the marginal distribution. Defaults to 4, so that the random effects for the marginal distribution is taken over the range [-4 SD; 4 SD]
• res.plot Type of residual plot ("res.vs.x": scatterplot versus X, "res.vs.pred": scatterplot versus predictions, "hist": histogram, "qqplot": QQ-plot) (default: "res.vs.x")
• smooth When TRUE, smoothed lines are added in the plots of predictions versus observations (default: FALSE)
• tit Title of the graph (default: none)
• type Type of the plot (as in the R plot function. Defaults to "b", so that both lines and symbols are shown
• units Name of the predictor used in the plots (X). Defaults to the name of the first predictor in the model (saemix.data$names$predictors[1])
• vpc.bin Number of binning intervals when plotting the VPC (the (vpc.bin-1) breakpoints are taken as the empirical quantiles of the X data). Defaults to 10
• vpc.interval Size of the prediction intervals. Defaults to 0.95 for the 95\%
• vpc.obs Should the observations be overlayed on the VPC plot. Defaults to TRUE
• vpc.pi Should prediction bands be computed around the median and the bounds of the prediction intervals for the VPC. Defaults to TRUE
• xlab Label for the X-axis. Defaults to the name of the X predictor followed by the unit in bracket (eg "Time (hr)")
• xlim Range for the X-axis. Defaults to NA, indicating that the range is to be set by the plot function
• xlog A logical value. If TRUE, a logarithmic scale is in use. Defaults to FALSE
• xname Name of the predictor used in the plots (X)
• ylab Label for the Y-axis. Defaults to the name of the response followed by the unit in bracket (eg "Concentration (mg/L)" (Default: none)
• ylim Range for the Y-axis. Defaults to NA, indicating that the range is to be set by the plot function
• ylog  A logical value. If TRUE, a logarithmic scale is in use. Defaults to FALSE

Plotting a SaemixData object also allows the following options:

**individual**  if TRUE, plots separate plots for each individual, otherwise plots a spaghetti plot of all the data. Defaults to FALSE

**limit** for individual plots, plots only a limited number of subjects (nmax). Defaults to TRUE

**nmax** for individual plots, when limit is TRUE, the maximum number of plots to produce. Defaults to 12

**sample** for individual plots, if TRUE, randomly samples nmax different subjects to plot. Defaults to FALSE (the first nmax subjects are used in the plots)

Usage

```r
saemix.plot.setoptions(saemixObject)
```

Arguments

- **saemixObject** an object returned by the *saemix* function

Details

This function can be used to create a list containing the default options and arguments used by the plot functions.

A more detailed description of the options set via these lists is provided in the PDF documentation. The "replace" functions are helper functions used within the plot functions. saemix.plot.setoptions has more available options than saemix.data.setoptions since it applies to all possible plots while the latter only applies to data.

Value

A list containing the options set at their default value. This list can be stored in an object and its elements modified to provide suitable graphs.

Author(s)

Emmanuelle Comets <emmanuelle.comets@inserm.fr>, Audrey Lavenu, Marc Lavielle.

References


saemix.predict

Compute model predictions after an saemix fit

Description

In nonlinear mixed effect models, different types of predictions may be obtained, including individual predictions and population predictions. This function takes an SaemixObject and adds any missing predictions for maximum a posteriori and conditional mean estimations of the individual parameters, and for the different types of individual and population predictions for the response variable.

Usage

saemix.predict(object, type = c("ipred", "ypred", "ppred", "icpred"))

Arguments

object an SaemixObject object
type the type of predictions (ipred= individual, ppred=population predictions obtained with the population estimates, ypred=mean of the population predictions, icpred=conditional predictions). By default, computes all the predictions and residuals, along with the corresponding parameter estimates

Details

This function is used internally by saemix to automatically compute a number of elements needed for diagnostic plots. It is normally executed directly during a call to saemix() but can be called to add residuals

Value

an updated SaemixObject object
saemixControl

**List of options for running the algorithm SAEM**

**Description**

List containing the variables relative to the optimisation algorithm. All these elements are optional and will be set to default values when running the algorithm if they are not specified by the user.

**Usage**

```r
saemixControl(
  map = TRUE,
  fim = TRUE,
  ll.is = TRUE,
  ll.gq = FALSE,
  nbiter.saemix = c(300, 100),
  nbiter.sa = NA,
  nb.chains = 1,
  fix.seed = TRUE,
  seed = 23456,
  nmc.is = 5000,
  nu.is = 4,
  print.is = FALSE,
  nbdisplay = 100,
  displayProgress = FALSE,
  nbiter.burn = 5,
  nbiter.map = 5,
  nbiter.mcmc = c(2, 2, 2, 0),
  proba.mcmc = 0.4,
  stepsize.rw = 0.4,
  rw.init = 0.5,
  alpha.sa = 0.97,
  nnodes.gq = 12,
  nsd.gq = 4,
  maxim.maxiter = 100,
  nb.sim = 1000,
  nb.simpred = 100,
  ipar.lmcmc = 50,
  ipar.rmcnc = 0.05,
  print = TRUE,
  save = TRUE,
  save.graphs = TRUE,
  directory = "newdir",
  warnings = FALSE
)
```
Arguments

map a boolean specifying whether to estimate the individual parameters (MAP estimates). Defaults to TRUE

fim a boolean specifying whether to estimate the Fisher Information Matrix and derive the estimation errors for the parameters. Defaults to TRUE. The linearised approximation to the log-likelihood is also computed in the process

ll.is a boolean specifying whether to estimate the log-likelihood by importance sampling. Defaults to TRUE

ll.gq a boolean specifying whether to estimate the log-likelihood by Gaussian quadrature. Defaults to FALSE

nbiter.saemix nb of iterations in each step (a vector containing 2 elements, nbiter.saemix[1] for the exploration phase of the algorithm (K1) and nbiter.saemix[2] for the smoothing phase (K2))

nbiter.sa nb of iterations subject to simulated annealing (defaults to nbiter.saemix[1]/2, will be cut down to K1=nbiter.saemix[1] if greater than that value). We recommend to stop simulated annealing before the end of the exploration phase (nbiter.saemix[1]).

nb.chains nb of chains to be run in parallel in the MCMC algorithm (defaults to 1)

fix.seed TRUE (default) to use a fixed seed for the random number generator. When FALSE, the random number generator is initialised using a new seed, created from the current time. Hence, different sessions started at (sufficiently) different times will give different simulation results. The seed is stored in the element seed of the options list.

seed seed for the random number generator. Defaults to 123456

nmc.is nb of samples used when computing the likelihood through importance sampling

nu.is number of degrees of freedom of the Student distribution used for the estimation of the log-likelihood by Importance Sampling. Defaults to 4

print.is when TRUE, a plot of the likelihood as a function of the number of MCMC samples when computing the likelihood through importance sampling is produced and updated every 500 samples. Defaults to FALSE

nbdisplay nb of iterations after which to display progress

displayProgress when TRUE, the convergence plots are plotted after every nbdisplay iteration, and a dot is written in the terminal window to indicate progress. When FALSE, plots are not shown and the algorithm runs silently. Defaults to FALSE

nbiter.burn nb of iterations for burning

nbiter.map nb of iterations of the MAP kernel (4th kernel) to run at the beginning of the estimation process (defaults to nbiter.saemix[1]/10 if nbiter.mcmc[4] is more than 0) (EXPERIMENTAL, see Karimi et al. 2019 for details)

nbiter.mcmc nb of iterations in each kernel during the MCMC step

proba.mcmc probability of acceptance

stepsize.rw stepsize for kernels q2 and q3 (defaults to 0.4)
### saemixControl

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>rw.init</code></td>
<td>initial variance parameters for kernels (defaults to 0.5)</td>
</tr>
<tr>
<td><code>alpha.sa</code></td>
<td>parameter controlling cooling in the Simulated Annealing algorithm (defaults to 0.97)</td>
</tr>
<tr>
<td><code>nnodes.gq</code></td>
<td>number of nodes to use for the Gaussian quadrature when computing the likelihood with this method (defaults to 12)</td>
</tr>
<tr>
<td><code>nsd.gq</code></td>
<td>span (in SD) over which to integrate when computing the likelihood by Gaussian quadrature. Defaults to 4 (eg 4 times the SD)</td>
</tr>
<tr>
<td><code>maxim.maxiter</code></td>
<td>Maximum number of iterations to use when maximising the fixed effects in the algorithm. Defaults to 100</td>
</tr>
<tr>
<td><code>nb.sim</code></td>
<td>number of simulations to perform to produce the VPC plots or compute npde. Defaults to 1000</td>
</tr>
<tr>
<td><code>nb.simpred</code></td>
<td>number of simulations used to compute mean predictions (ypred element), taken as a random sample within the nb.sim simulations used for npde</td>
</tr>
<tr>
<td><code>ipar.lmcmc</code></td>
<td>number of iterations required to assume convergence for the conditional estimates. Defaults to 50</td>
</tr>
<tr>
<td><code>ipar.rmcmc</code></td>
<td>confidence interval for the conditional mean and variance. Defaults to 0.95</td>
</tr>
<tr>
<td><code>print</code></td>
<td>whether the results of the fit should be printed out. Defaults to TRUE</td>
</tr>
<tr>
<td><code>save</code></td>
<td>whether the results of the fit should be saved to a file. Defaults to TRUE</td>
</tr>
<tr>
<td><code>save.graphs</code></td>
<td>whether diagnostic graphs and individual graphs should be saved to files. Defaults to TRUE</td>
</tr>
<tr>
<td><code>directory</code></td>
<td>the directory in which to save the results. Defaults to &quot;newdir&quot; in the current directory</td>
</tr>
<tr>
<td><code>warnings</code></td>
<td>whether warnings should be output during the fit. Defaults to FALSE</td>
</tr>
</tbody>
</table>

### Details

All the variables are optional and will be set to their default value when running `saemix`.

The function `saemix` returns an object with an element `options` containing the options used for the algorithm, with defaults set for elements which have not been specified by the user.

These elements are used in subsequent functions and are not meant to be used directly.

### Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

### References


See Also

SaemixData, SaemixModel, SaemixObject, saemix

Examples

```r
# All default options
saemix.options<--saemixControl()

# All default options, changing seed
saemix.options<--saemixControl(seed=632545)
```

---

**saemixData**  
*Function to create a SaemixData object*

**Description**

This function creates a SaemixData object. The only mandatory argument is the name of the dataset. If the dataset has a header (or named columns), the program will attempt to detect which column correspond to ID, predictor(s) and response. Warning messages will be printed during the object creation and should be read for details.

**Usage**

```r
saemixData(
  name.data,
  header,
  sep,
  na,
  name.group,
  name.predictors,
  name.response,
  name.X,
  name.covariates = c(),
  name.genetic.covariates = c(),
  name.mdv = "",
  name.cens = "",
  name.occ = "",
  name.ytype = "",
  units = list(x = "", y = "", covariates = c()),
  verbose = TRUE,
  automatic = TRUE
)
```
Arguments

name.data name of the dataset (can be a character string giving the name of a file on disk or of a dataset in the R session, or the name of a dataset

header whether the dataset/file contains a header. Defaults to TRUE

sep the field separator character. Defaults to any number of blank spaces ("")

na a character vector of the strings which are to be interpreted as NA values. Defaults to c(NA)

name.group name (or number) of the column containing the subject id

name.predictors name (or number) of the column(s) containing the predictors (the algorithm requires at least one predictor x)

name.response name (or number) of the column containing the response variable y modelled by predictor(s) x

name.X name of the column containing the regression variable to be used on the X axis in the plots (defaults to the first predictor)

name.covariates name (or number) of the column(s) containing the covariates, if present (otherwise missing)

name.genetic.covariates name (or number) of the column(s) containing the covariates, if present (otherwise missing)

name.mdv name of the column containing the indicator for missing variable

name.cens name of the column containing the indicator for censoring

name.occ name of the column containing the occasion

name.ytype name of the column containing the index of the response

units list with up to three elements, x, y and optionally covariates, containing the units for the X and Y variables respectively, as well as the units for the different covariates (defaults to empty)

verbose a boolean indicating whether messages should be printed out during the creation of the object

automatic a boolean indicating whether to attempt automatic name recognition when some column names are missing or wrong (defaults to TRUE)

Details

This function is the user-friendly constructor for the SaemixData object class. The read.saemixData is a helper function, used to read the dataset, and is not intended to be called directly.

This function is the user-friendly constructor for the SaemixData object class. The read is a helper function, used to read the dataset, and is not intended to be called directly.

Value

A SaemixData object (see saemixData).
Author(s)
Emmanuelle Comets <emmanuelle.comets@inserm.fr>, Audrey Lavenu, Marc Lavielle.

References

See Also
SaemixData, SaemixModel, saemixControl, saemix

Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
      name.group=c("Id"),name.predictors=c("Dose","Time"),
      name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
      units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

print(saemix.data)

plot(saemix.data)

SaemixData-class  Class "SaemixData"

Description
An object of the SaemixData class, representing a longitudinal data structure, used by the SAEM algorithm.

Slots
name.data  Object of class "character": name of the dataset
header  Object of class "logical": whether the dataset/file contains a header. Defaults to TRUE
sep  Object of class "character": the field separator character
na  Object of class "character": a character vector of the strings which are to be interpreted as NA values
messages Object of class "logical": if TRUE, the program will display information about the creation of the data object
automatic Object of class "logical": if TRUE, automatic name recognition is on (used at the creation of the object)
name.group Object of class "character": name of the column containing the subject id
name.predictors Object of class "character": name of the column(s) containing the predictors
name.response Object of class "character": name of the column containing the response variable y modelled by predictor(s) x
name.covariates Object of class "character": name of the column(s) containing the covariates, if present (otherwise empty)
name.X Object of class "character": name of the column containing the regression variable to be used on the X axis in the plots
name.mdv Object of class "character": name of the column containing the indicator variable denoting missing data
name.cens Object of class "character": name of the column containing the indicator variable denoting censored data (the value in the name.response column will be taken as the censoring value)
name.occ Object of class "character": name of the column containing the value of the occasion
name.ytype Object of class "character": name of the column containing the response number
trans.cov Object of class "list": the list of transformation applied to the covariates (currently unused, TODO)
units Object of class "list": list with up to three elements, x, y and optionally covariates, containing the units for the X and Y variables respectively, as well as the units for the different covariates
data Object of class "data.frame": dataframe containing the data, with columns for id (name.group), predictors (name.predictors), response (name.response), and covariates if present in the dataset (name.covariates). A column "index" contains the subject index (used to map the subject id). The column names, except for the additional column index, correspond to the names in the original dataset.
N Object of class "numeric": number of subjects
yorig Object of class "numeric": response data, on the original scale. Used when the error model is exponential
ocov Object of class "data.frame": original covariate data (before transformation in the algorithm)
ind.gen Object of class "logical": indicator for genetic covariates (internal)
ntot.obs Object of class "numeric": total number of observations
nind.obs Object of class "numeric": vector containing the number of observations for each subject

**Objects from the Class**

An object of the SaemixData class can be created by using the function `saemixData` and contain the following slots:
Methods

[<- signature(x = "SaemixData"): replace elements of object

[ signature(x = "SaemixData"): access elements of object

initialize signature(.Object = "SaemixData"): internal function to initialise object, not to be used

plot signature(x = "SaemixData"): plot the data

print signature(x = "SaemixData"): prints details about the object (more extensive than show)

read signature(object = "SaemixData"): internal function, not to be used

showall signature(object = "SaemixData"): shows all the elements in the object

show signature(object = "SaemixData"): prints details about the object

summary signature(object = "SaemixData"): summary of the data. Returns a list with a number of elements extracted from the dataset (N: the number of subjects; nobs: the total number of observations; nind.obs: a vector giving the number of observations for each subject; id: subject ID; x: predictors; y: response, and, if present in the data, covariates: the covariates (as many lines as observations) and ind.covariates: the individual covariates (one line per individual).

subset signature(object = "SaemixData"): extract part of the data; this function will operate on the rows of the dataset (it can be used for instance to extract the data corresponding to the first ten subjects)

Author(s)

Emmanuelle Comets <emmanuelle.comets@inserm.fr>

Audrey Lavenu

Marc Lavielle.

References


See Also

saemixData SaemixModel saemixControl saemix
Examples

```r
showClass("SaemixData")

# Specifying column names
data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
  name.group=c("Id"),name.predictors=c("Dose","Time"),
  name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
  units=list(x="hr",y="mg/L",covariates=c("kg","-"), name.X="Time")

# Specifying column numbers
data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
  name.group=1,name.predictors=c(2,3),name.response=c(4), name.covariates=5:6,
  units=list(x="hr",y="mg/L",covariates=c("kg","-"), name.X="Time")

# No column names specified, using automatic recognition of column names
data(PD1.saemix)
saemix.data<-saemixData(name.data=PD1.saemix,header=TRUE,
  name.covariates=c("gender"),units=list(x="mg",y="-",covariates=c("-")))
```

---

### saemixModel

**Function to create a SaemixModel object**

**Description**

This function creates a SaemixModel object. The two mandatory arguments are the name of a R function computing the model in the SAEMIX format (see details and examples) and a matrix psi0 giving the initial estimates of the fixed parameters in the model, with one row for the population mean parameters and one row for the covariate effects (see documentation).

**Usage**

```r
saemixModel(model, psi0, description = "", modeltype = "structural",
  name.response = "", name.sigma = character(),
  error.model = character(),
  transform.par = numeric(),
  fixed.estim = numeric(),
  covariate.model = matrix(nrow = 0, ncol = 0),
  covariance.model = matrix(nrow = 0, ncol = 0),
  omega.init = matrix(nrow = 0, ncol = 0),
  error.init = numeric(),
```
Arguments

model  name of the function used to compute the structural model. The function should return a vector of predicted values given a matrix of individual parameters, a vector of indices specifying which records belong to a given individual, and a matrix of dependent variables (see example below).

psi0  a matrix with a number of columns equal to the number of parameters in the model, and one (when no covariates are available) or two (when covariates enter the model) giving the initial estimates for the fixed effects. The column names of the matrix should be the names of the parameters in the model, and will be used in the plots and the summaries. When only the estimates of the mean parameters are given, psi0 may be a named vector.

description  a character string, giving a brief description of the model or the analysis

description  a character string, giving model type (structural or likelihood)

name.response  the name of the dependent variable

name.sigma  a vector of character string giving the names of the residual error parameters

error.model  type of residual error model (valid types are constant, proportional, combined and exponential). Defaults to constant

transform.par  the distribution for each parameter (0=normal, 1=log-normal, 2=probit, 3=logit). Defaults to a vector of 1s (all parameters have a log-normal distribution)

fixed.estim  whether parameters should be estimated (1) or fixed to their initial estimate (0). Defaults to a vector of 1s

covariate.model  a matrix giving the covariate model. Defaults to no covariate in the model

covariance.model  a square matrix of size equal to the number of parameters in the model, giving the variance-covariance matrix of the model: 1s correspond to estimated variances (in the diagonal) or covariances (off-diagonal elements). Defaults to the identity matrix

omega.init  a square matrix of size equal to the number of parameters in the model, giving the initial estimate for the variance-covariance matrix of the model.

error.init  a vector of size 2 giving the initial value of a and b in the error model. Defaults to 1 for each estimated parameter in the error model

name.modpar  names of the model parameters, if they are not given as the column names (or names) of psi0

simulate.function  for non-Gaussian data models, defined as modeltype='likelihood', the name of the function used to simulate from the structural model. The function should have the same header as the model function, and should return a vector of simulated values given a matrix of individual parameters, a vector of indices specifying which records belong to a given individual, and a matrix of dependent variables (see example in the documentation, section discrete data examples)
verbose is a boolean, controlling whether information about the created should be printed out. Defaults to TRUE.

Details

This function is the user-friendly constructor for the SaemixModel object class.

Value

A SaemixModel object (see saemixModel).

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

SaemixData, SaemixModel, saemixControl, saemix

Examples

```r
model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}

saemix.model<-saemixModel(model=model1cpt,
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,
  dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,1,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,1,0,0,0,1,0),ncol=3,byrow=TRUE),error.model="constant")
```
Description

An object of the SaemixModel class, representing a nonlinear mixed-effect model structure, used by the SAEM algorithm.

Objects from the Class

An object of the SaemixModel class can be created by using the function `saemixModel` and contain the following slots:

- **model**: Object of class "function": name of the function used to get predictions from the model (see the User Guide and the online examples for the format and what this function should return).
- **description**: Object of class "character": an optional text description of the model
- **psi0**: Object of class "matrix": a matrix with named columns containing the initial estimates for the parameters in the model (first line) and for the covariate effects (second and subsequent lines, optional). The number of columns should be equal to the number of parameters in the model.
- **simulate.function**: Object of class "function": for non-Gaussian data models, name of the function used to simulate from the model.
- **transform.par**: Object of class "numeric": vector giving the distribution for each model parameter (0: normal, 1: log-normal, 2: logit, 3: probit). Its length should be equal to the number of parameters in the model.
- **fixed.estim**: Object of class "numeric": for each parameter, 0 if the parameter is fixed and 1 if it should be estimated. Defaults to a vector of 1 (all parameters are estimated). Its length should be equal to the number of parameters in the model.
- **error.model**: Object of class "character": name of the error model. Valid choices are "constant" (default), "proportional" and "combined" (see equations in User Guide, except for combined which was changed to $y = f + \sqrt{a^2 + b^2 f^2} \cdot e$)
- **covariate.model**: Object of class "matrix": a matrix of 0's and 1's, with a 1 indicating that a parameter-covariate relationship is included in the model (and an associated fixed effect will be estimated). The number of columns should be equal to the number of parameters in the model and the number of rows to the number of covariates.
- **covariance.model**: Object of class "matrix": a matrix f 0's and 1's giving the structure of the variance-covariance matrix. Defaults to the Identity matrix (diagonal IV, no correlations between parameters)
- **omega.init**: Object of class "matrix": a matrix giving the initial estimate for the variance-covariance matrix
- **error.init**: Object of class "numeric": a vector giving the initial estimate for the parameters of the residual error

Additional elements are added to the model object after a call to `saemix` and are used in the algorithm.
SaemixModel-class

Methods

[- signature(x = "SaemixModel"): replace elements of object
[ signature(x = "SaemixModel"): access elements of object
initialize signature(.Object = "SaemixModel"): internal function to initialise object, not to be used
plot signature(x = "SaemixModel"): plot predictions from the model
print signature(x = "SaemixModel"): prints details about the object (more extensive than show)
showall signature(object = "SaemixModel"): shows all the elements in the object
show signature(object = "SaemixModel"): prints details about the object

Author(s)

Emmanuelle Comets <emmanuelle.comets@inserm.fr>
Audrey Lavenu
Marc Lavielle.

References


See Also

SaemixData SaemixObject saemixControl saemix plot.saemix

Examples

showClass("SaemixModel")

# Model function for continuous data
## structural model: a one-compartment model with oral absorption
model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
tim<-xidep[,2]
ka<-psi[id,1]
V<-psi[id,2]
CL<-psi[id,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}
# Corresponding SaemixModel, assuming starting parameters ka=1, V=20, CL=0.5
# and log-normal distributions for the parameters
model1 <- saemixModel(model=model1cpt,
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1,20,0.5),ncol=3, byrow=TRUE,
  dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1))

# Model function for discrete data
## logistic regression for the probability of the observed outcome
binary.model<-function(psi,id,xidep) {
  tim<-xidep[,1]
  y<-xidep[,2]
  inter<-psi[id,1]
  slope<-psi[id,2]
  logit<-inter+slope*tim
  pevent<-exp(logit)/(1+exp(logit))
  pobs = (y==0)*(1-pevent)+(y==1)*pevent
  logpdf <- log(pobs)
  return(logpdf)
}

## Corresponding SaemixModel, assuming starting parameters inter=-5, slope=-1
# and normal distributions for both parameters
# note that the modeltype argument is set to likelihood
saemix.model<-saemixModel(model=binary.model,description="Binary model",
  modeltype="likelihood",
  psi0=matrix(c(-5,-.1,0,0),ncol=2,byrow=TRUE,dimnames=list(NULL,c("inter","slope"))),
  transform.par=c(0,0))

## saemix cannot infer the distribution of the outcome directly from the model
## Here we therefore define a simulation function, needed for diagnostics
### Note the similarity and differences with the model function
simulBinary<-function(psi,id,xidep) {
  tim<-xidep[,1]
  y<-xidep[,2]
  inter<-psi[id,1]
  slope<-psi[id,2]
  logit<-inter+slope*tim
  pevent<-1/(1+exp(-logit))
  ysim<-rbinom(length(tim),size=1, prob=pevent)
  return(ysim)
}

saemix.model<-saemixModel(model=binary.model,description="Binary model",
  modeltype="likelihood", simulate.function=simulBinary,
  psi0=matrix(c(-5,-.1,0,0),ncol=2,byrow=TRUE,dimnames=list(NULL,c("inter","slope"))),
  transform.par=c(0,0))
SaemixObject-class

Description

An object of the SaemixObject class, storing the input to saemix, and the results obtained by a call to the SAEM algorithm

Details

Details of the algorithm can be found in the pdf file accompanying the package.

Objects from the Class

An object of the SaemixObject class is created after a call to saemix and contain the following slots:

- data: Object of class "SaemixData": saemix dataset, created by a call to saemixData
- model: Object of class "SaemixModel": saemix model, created by a call to saemixModel
- results: Object of class "SaemixData": saemix dataset, created by a call to saemixData
- rep.data: Object of class "SaemixRepData": (internal) replicated saemix dataset, used the execution of the algorithm
- sim.data: Object of class "SaemixSimData": simulated saemix dataset
- options: Object of class "list": list of settings for the algorithm
- prefs: Object of class "list": list of graphical options for the graphs

Methods

- [<- signature(x = "SaemixObject"): replace elements of object
- [ signature(x = "SaemixObject"): access elements of object
- initialize signature(.Object = "SaemixObject"): internal function to initialise object, not to be used
- plot signature(x = "SaemixObject"): plot the data
- print signature(x = "SaemixObject"): prints details about the object (more extensive than show)
- showall signature(object = "SaemixObject"): shows all the elements in the object
- show signature(object = "SaemixObject"): prints details about the object
- summary signature(object = "SaemixObject"): summary of the object. Returns a list with a number of elements extracted from the object.

Author(s)

Emmanuelle Comets <emmanuelle.comets@inserm.fr>
Audrey Lavenu
Marc Lavielle.
References


See Also

SaemixData SaemixModel saemixControl saemix plot.saemix

Examples

showClass("SaemixObject")

saemixPredictNewdata Predictions for a new dataset

Description

Predictions for a new dataset

Usage

saemixPredictNewdata(
  saemixObject,
  newdata,
  type = c("ipred", "ypred", "ppred", "icpred"),
  nsamp = 1
)

Arguments

saemixObject an SaemixObject from a fitted run
newdata a dataframe containing the new data. The dataframe must contain the same information as the original dataset (column names, etc...)
type one or several of "ipred" (individual predictions using the MAP estimates), "ppred" (population predictions obtained using the population parameters f(E(theta))), "ypred" (mean of the population predictions (E(f(theta)))), "icpred" (individual predictions using the conditional mean estimates). Defaults to "ppred".
nsamp an integer, ignored for other types than icpred; if icpred, returns both the mean of the conditional distribution and nsamp samples, with the corresponding predictions. Defaults to 1.
**Details**

This function is the workhorse behind the predict method for SaemixObject. It computes predictions for a new dataframe based on the results of an saemix fit. Since the predict function only returns predicted values, this function is provided so that users can access other elements, for example the different types of parameter estimates (such as individual or population parameters) associated with the predictions. For other purposes such as simply obtaining model predictions, we suggest using the predict() method with or without newdata.

The function uses estimateMeanParametersNewdata() to set the population estimates for the individual parameters taking into account the individual covariates and doses, and estimateIndividualParametersNewdata() to derive individual estimates by computing the mean of the conditional distributions (type="icpred") or the MAP estimate (type="ipred")

**Value**

- a list with three components (five if type includes "icpred" and nsamp>1)
  - **param** a dataframe with the estimated parameters. The columns in the dataframe depend on which type of predictions were requested (argument type)
  - **predictions** a dataframe with the predictions. The columns in the dataframe depend on which type of predictions were requested (argument type)
  - **saemixObject** the SaemixObject with the data slot replaced by the new data. The elements of the results slot pertaining to individual (including population individual parameters) predictions and likelihood will have been removed, and only the elements computed within the function will have been replaced (eg individual estimated parameters and predictions for the new data)
  - **parSample** a dataframe with parameters sampled from the conditional distribution of the individual parameters (only present if type includes ‘icpred’ and nsamp>1)
  - **predSample** a dataframe with the predictions corresponding to the parameters sampled from the conditional distribution of the individual parameters (only present if type includes ‘icpred’ and nsamp>1)

**Examples**

# TODO

---

**SaemixRes-class**  
*Class "SaemixRes"*

**Description**

An object of the SaemixRes class, representing the results of a fit through the SAEM algorithm.
Slots

- `modeltype` - string giving the type of model used for analysis
- `status` - string indicating whether a model has been run successfully; set to "empty" at initialisation, used to pass on error messages or fit status
- `name.fixed` - a vector containing the names of the fixed parameters in the model
- `name.random` - a vector containing the names of the random parameters in the model
- `name.sigma` - a vector containing the names of the parameters of the residual error model
- `npar.est` - the number of parameters estimated (fixed, random and residual)
- `nbeta.random` - the number of estimated fixed effects for the random parameters in the model
- `nbeta.fixed` - the number of estimated fixed effects for the non random parameters in the model
- `fixed.effects` - a vector giving the estimated h(mu) and betas
- `fixed.psi` - a vector giving the estimated h(mu)
- `betas` - a vector giving the estimated mu
- `betaC` - a vector with the estimates of the fixed effects for covariates
- `omega` - the estimated variance-covariance matrix
- `respar` - the estimated parameters of the residual error model
- `fim` - the Fisher information matrix
- `se.fixed` - a vector giving the estimated standard errors of estimation for the fixed effect parameters
- `se.omega` - a vector giving the estimated standard errors of estimation for Omega
- `se.cov` - a matrix giving the estimated SE for each term of the covariance matrix (diagonal elements represent the SE on the variances of the random effects and off-diagonal elements represent the SE on the covariance terms)
- `se.respar` - a vector giving the estimated standard errors of estimation for the parameters of the residual variability
- `conf.int` - a dataframe containing the estimated parameters, their estimation error (SE), coefficient of variation (CV), and the associated confidence intervals; the variabilities for the random effects are presented first as estimated (variances) then converted to standard deviations (SD), and the correlations are computed. For SD and correlations, the SE are estimated via the delta-method
- `parpop` - a matrix tracking the estimates of the population parameters at each iteration
- `allpar` - a matrix tracking the estimates of all the parameters (including covariate effects) at each iteration
- `indx.fix` - the index of the fixed parameters (used in the estimation algorithm)
- `indx.cov` - the index of the covariance parameters (used in the estimation algorithm)
- `indx.omega` - the index of the random effect parameters (used in the estimation algorithm)
- `indx.res` - the index of the residual error model parameters (used in the estimation algorithm)
- `MCOV` - a matrix of covariates (used in the estimation algorithm)
- `cond.mean.phi` - a matrix giving the conditional mean estimates of phi (estimated as the mean of the conditional distribution)
cond.mean.psi a matrix giving the conditional mean estimates of psi (h(cond.mean.phi))
cond.var.phi a matrix giving the variance on the conditional mean estimates of phi (estimated as the variance of the conditional distribution)
cond.mean.eta a matrix giving the conditional mean estimates of the random effect eta
cond.shrinkage a vector giving the shrinkage on the conditional mean estimates of eta
mean.phi a matrix giving the population estimate (Ci*mu) including covariate effects, for each subject
map.psi a dataframe giving the MAP estimates of individual parameters
map.phi a dataframe giving the MAP estimates of individual phi
map.eta a matrix giving the individual estimates of the random effects corresponding to the MAP estimates
map.shrinkage a vector giving the shrinkage on the MAP estimates of eta
phi individual parameters, estimated at the end of the estimation process as the average over the chains of the individual parameters sampled during the successive E-steps
psi.samp a three-dimensional array with samples of psi from the conditional distribution
phi.samp a three-dimensional array with samples of phi from the conditional distribution
phi.samp.var a three-dimensional array with the variance of phi
ll.lin log-likelihood computed by lineariation
aic.lin Akaike Information Criterion computed by linearisation
bic.lin Bayesian Information Criterion computed by linearisation
bic.covariate.lin Specific Bayesian Information Criterion for covariate selection computed by linearisation
ll.is log-likelihood computed by Importance Sampling
aic.is Akaike Information Criterion computed by Importance Sampling
bic.is Bayesian Information Criterion computed by Importance Sampling
bic.covariate.is Specific Bayesian Information Criterion for covariate selection computed by Importance Sampling
LL a vector giving the conditional log-likelihood at each iteration of the algorithm
ll.gq log-likelihood computed by Gaussian Quadrature
aic.gq Akaike Information Criterion computed by Gaussian Quadrature
bic.gq Bayesian Information Criterion computed by Gaussian Quadrature
bic.covariate.gq Specific Bayesian Information Criterion for covariate selection computed by Gaussian Quadrature
predictions a data frame containing all the predictions and residuals in a table format
ppred a vector giving the population predictions obtained with the population estimates
ypred a vector giving the mean population predictions
ipred a vector giving the individual predictions obtained with the MAP estimates
ipred a vector giving the individual predictions obtained with the conditional estimates
ires a vector giving the individual residuals obtained with the MAP estimates
iwrres a vector giving the individual weighted residuals obtained with the MAP estimates
icwres a vector giving the individual weighted residuals obtained with the conditional estimates
wres a vector giving the population weighted residuals
npde a vector giving the normalised prediction distribution errors
pd a vector giving the prediction discrepancies

Objects from the Class
An object of the SaemixData class can be created by using the function `saemixData` and contain the following slots:

Methods

\[
\text{<- signature(x = "SaemixRes")}: \text{replace elements of object}
\]

\[
\text{[ signature(x = "SaemixRes")}: \text{access elements of object}
\]

\[
\text{initialize signature(.Object = "SaemixRes")}: \text{internal function to initialise object, not to be used}
\]

\[
\text{print signature(x = "SaemixRes")}: \text{prints details about the object (more extensive than show)}
\]

\[
\text{read signature(object = "SaemixRes")}: \text{internal function, not to be used}
\]

\[
\text{showall signature(object = "SaemixRes")}: \text{shows all the elements in the object}
\]

\[
\text{show signature(object = "SaemixRes")}: \text{prints details about the object}
\]

\[
\text{summary signature(object = "SaemixRes")}: \text{summary of the results. Returns a list with a number of elements extracted from the results ()}
\]

Author(s)
Emmanuelle Comets <emmanuelle.comets@inserm.fr>
Audrey Lavenu
Marc Lavielle.

References


See Also

saemixData SaemixModel saemixControl saemix
show-methods

Examples

methods(class="SaemixRes")
showClass("SaemixRes")

Description

Prints a short summary of an object

Usage

## S4 method for signature 'SaemixData'
show(object)

## S4 method for signature 'SaemixRepData'
show(object)

## S4 method for signature 'SaemixSimData'
show(object)

## S4 method for signature 'SaemixModel'
show(object)

## S4 method for signature 'SaemixRes'
show(object)

## S4 method for signature 'SaemixObject'
show(object)

Arguments

object an object of type SaemixData, SaemixModel, SaemixRes or SaemixObject

Methods

list("signature(x = "ANY")") Default show function
list("signature(x = "SaemixData")") Prints a short summary of a SaemixData object
list("signature(x = "SaemixModel")") Prints a short summary of a SaemixModel object
list("signature(x = "SaemixObject")") Prints a short summary of the results from a SAEMIX fit
list("signature(x = "SaemixRes")") Not user-level
The `showall` function is used to visualise the majority of the elements of an object.

### Usage

```r
showall(object)
```

### Arguments

- `object` showall methods are available for objects of type `SaemixData`, `SaemixModel` and `SaemixObject`

### Value

None

### Methods

- `list("signature(x = \"SaemixData\")")` Prints an extensive summary of a `SaemixData` object
- `list("signature(x = \"SaemixModel\")")` Prints an extensive summary of a `SaemixModel` object
- `list("signature(x = \"SaemixObject\")")` Prints an extensive summary of the results from a SAEMIX fit
- `list("signature(x = \"SaemixRes\")")` Not user-level
**simulate.SaemixObject**

Perform simulations under the model for an saemixObject object

**Description**

This function is used to simulate data under the empirical design, using the model and estimated parameters from a fit.

**Usage**

```r
## S3 method for class 'SaemixObject'
simulate(
  object,
  nsim,
  seed,
  predictions = TRUE,
```
simulate.SaemixObject

```
res.var = TRUE,
uncertainty = FALSE,
"
```

Arguments

- **object**: an saemixObject object returned by the `saemix` function
- **nsim**: Number of simulations to perform. Defaults to the `nb.simpred` element in options
- **seed**: if non-null, seed used to initiate the random number generator (defaults to NULL)
- **predictions**: Whether the simulated parameters should be used to compute predictions. Defaults to TRUE. If FALSE, only individual parameters are simulated.
- **res.var**: Whether residual variability should be added to the predictions. Defaults to TRUE.
- **uncertainty**: Uses uncertainty (currently not implemented). Defaults to FALSE
- **...**: additional arguments, unused (included for compatibility with the generic)

Details

The simulated data can then be used to produce Visual Predictive Check graphs, as well as to compute the normalised prediction distribution errors (npde).

This function replaces the previous function (simul.saemix), which will be deprecated in future versions but can still be called as previously for compatibility purposes.

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr, Audrey Lavenu, Marc Lavielle.

References


See Also

`SaemixObject, saemix, saemix.plot.data, saemix.plot.convergence, saemix.plot.llis, saemix.plot.randeff, saemix.plot.obsvspred, saemix.plot.fits, saemix.plot.parcov, saemix.plot.distpsi, saemix.plot.scatterres, saemix.plot.vpc`
simulateDiscreteSaemix

Perform simulations under the model for an saemixObject object defined by its log-likelihood

Description

This function is used to simulate from discrete response models.

Usage

simulateDiscreteSaemix(
  object,
  nsim,
  seed,
  uncertainty = FALSE,
  verbose = FALSE
)

Arguments

object an saemixObject object returned by the saemix function. The model must contain a slot simulate.function, containing a function with the same structure as the model functions (arguments (psi, id, xidep)) which returns simulated responses when given a set of individual parameters (psi) and the corresponding predictors (xidep)

nsim Number of simulations to perform. Defaults to the nb.simpred element in options

seed if non-null, seed used to initiate the random number generator (defaults to NULL)

uncertainty Uses uncertainty (currently not implemented). Defaults to FALSE

verbose if TRUE, prints messages (defaults to FALSE)

Details

To call this function, the user needs to define a simulate.function matching the model function in the object. The function will then be used to simulate data under the empirical design, using the model and estimated parameters from a fit.

This function calls simulate.SaemixObject with the prediction=FALSE option to simulate individual parameters, then the simulate.function to obtain corresponding predictions.

Author(s)

Emmanuelle Comets emmanuelle.comets@inserm.fr

See Also

SaemixObject, saemix, simulate.SaemixObject
Stepwise procedure for joint selection of covariates and random effects

Description

Joint selection of covariates and random effects in a nonlinear mixed effects model by a stepwise-type algorithm based on two different versions of BIC for covariate selection and random effects selection respectively. Selection is made among the covariates as such specified in the SaemixData object. Only uncorrelated random effects structures are considered.

Usage

```r
step.saemix(
  saemixObject,
  trace = TRUE,
  direction = "forward",
  covariate.init = NULL
)
```

Arguments

- `saemixObject`: An object returned by the `saemix` function
- `trace`: If TRUE, a table summarizing the steps of the algorithm is printed. Default "TRUE"
- `direction`: The mode of stepwise search, can be one of "both", "backward", or "forward", with a default of "forward".
- `covariate.init`: A matrix specifying the initial covariate structure to be considered in the algorithm. `covariate.init` is only used if the `direction` argument is "both".

Value

An object of the SaemixObject class storing the covariate model and the covariance structure of random effects of the final model.

Author(s)

Maud Delattre

References

Examples

data(theo.saemix)

saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA, name.group=c("Id"),name.predictors=c("Dose","Time"), name.response=c("Concentration"),name.covariates=c("Weight","Sex"), units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")

# Definition of models to be compared
model1cpt<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}

saemix.model1<-saemixModel(model=model1cpt,modeltype="structural", description="One-compartment model with first-order absorption", psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))), transform.par=c(1,1,1), covariate.model=matrix(c(0,0,1,0,0,0),ncol=3,byrow=TRUE))

saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE)

saemix.fit1<-saemix(saemix.model1,saemix.data,saemix.options)

## Not run:
res.forward <- step.saemix(saemix.fit1, direction = "forward")
res.backward <- step.saemix(saemix.fit1, direction = "backward")
covariate.init <- matrix(c(1,0,0,0,1,0),ncol=3,nrow=2)
res.stepwise <- step.saemix(saemix.fit1, direction="both")
## End(Not run)

stepwise.procedure

Stepwise procedure for joint selection of covariates and random effects

Description

Joint selection of covariates and random effects in a nonlinear mixed effects model by a stepwise-type algorithm based on two different versions of BIC for covariate selection and random effects selection respectively. Selection is made among the covariates as such specified in the SaemixData object. Only uncorrelated random effects structures are considered.

Usage

stepwise.procedure(saemixObject, covariate.init = NULL, trace = TRUE)
subset

Arguments

- saemixObject: An object returned by the saemix function
- covariate.init: A matrix specifying the initial covariate structure to be considered in the algorithm.
- trace: If TRUE, a table summarizing the steps of the algorithm is printed. Default "TRUE"

Value

An object of the SaemixObject class storing the covariate model and the covariance structure of random effects of the final model.

Author(s)

Maud Delattre

References


Description

Return an SaemixData object containing the subset of data which meets conditions.

Usage

```r
## S3 method for class 'SaemixData'
subset(x, subset, ...)
```

Arguments

- x: saemixData object
- subset: logical expression indicating elements or rows to keep: missing values are taken as false
- ...: additional parameters (ignored)

Value

an object of class "SaemixData"

Examples

```r
# TODO
```
summary-methods

Methods for Function summary

Description

Methods for function summary

summary method for class SaemixData

Usage

## S4 method for signature 'SaemixData'
summary(object, print = TRUE, ...)

## S4 method for signature 'SaemixModel'
summary(object, print = TRUE, ...)

## S4 method for signature 'SaemixRes'
summary(object, print = TRUE, ...)

## S4 method for signature 'SaemixObject'
summary(object, print = TRUE, ...)

Arguments

object an object of class SaemixData
print a boolean controlling whether to print the output or return it silently
... additional arguments (ignored)

Value

a list with a number of elements extracted from the dataset

N number of subjects
nobs the total number of observations
nind.obs a vector giving the number of observations for each subject
id subject ID; x: predictors; y: response, and, if present in the data, covariates: the covariates (as many lines as observations) and ind.covariates: the individual covariates (one

Methods

list("signature(x = \"ANY\")") default summary function ?
list("signature(x = \"SaemixData\")") summary of the data
list("signature(x = \"SaemixModel\")") summary of the model
list("signature(x = \"SaemixObject\")") summary of an SaemixObject
Description
Performs tests for the normalised prediction distribution errors returned by npde

Usage
testnpde(npde)

Arguments
npde the vector of prediction distribution errors

Details
Given a vector of normalised prediction distribution errors (npde), this function compares the npde to the standardised normal distribution N(0,1) using a Wilcoxon test of the mean, a Fisher test of the variance, and a Shapiro-Wilks test for normality. A global test is also reported. The helper functions kurtosis and skewness are called to compute the kurtosis and skewness of the distribution of the npde.

Value
a list containing 4 components:
| Wilcoxon test of mean=0 | compares the mean of the npde to 0 using a Wilcoxon test |
| variance test | compares the variance of the npde to 1 using a Fisher test |
| SW test of normality | compares the npde to the normal distribution using a Shapiro-Wilks test |
| global test | an adjusted p-value corresponding to the minimum of the 3 previous p-values multiplied by the number of tests (3), or 1 if this p-value is larger than 1. |

Author(s)
Emmanuelle Comets emmanuelle.comets@inserm.fr

References

See Also
saemix, saemix.plot.npde
Description

The theo.saemix data frame has 132 rows and 6 columns of data from an experiment on the pharmacokinetics of theophylline. A column with gender was added to the original data for demo purposes, and contains simulated data.

Usage

theo.saemix

Format

This data frame contains the following columns:

- **Id**: an ordered factor with levels 1, ..., 12 identifying the subject on whom the observation was made. The ordering is by time at which the observation was made.
- **Dose**: dose of theophylline administered orally to the subject (mg/kg).
- **Time**: time since drug administration when the sample was drawn (hr).
- **Concentration**: theophylline concentration in the sample (mg/L).
- **Weight**: weight of the subject (kg).
- **Sex**: gender (simulated, 0=male, 1=female)

Details

Boeckmann, Sheiner and Beal (1994) report data from a study by Dr. Robert Upton of the kinetics of the anti-asthmatic drug theophylline. Twelve subjects were given oral doses of theophylline then serum concentrations were measured at 11 time points over the next 25 hours.

These data are analyzed in Davidian and Giltinan (1995) and Pinheiro and Bates (2000) using a two-compartment open pharmacokinetic model.

These data are also available in the library datasets under the name Theoph in a slightly modified format and including the data at time 0. Here, we use the file in the format provided in the NONMEM installation path (see the User Guide for that software for details).

Source


References


Examples

data(theo.saemix)

# Plotting the theophylline data
plot(Concentration~Time, data=theo.saemix, xlab="Time after dose (hr)", ylab="Theophylline concentration (mg/L)"

saemix.data<-saemixData(name.data=theo.saemix, header=TRUE, sep=" ", na=NA, name.group=c("Id"), name.predictors=c("Dose","Time"), name.response=c("Concentration"), name.covariates=c("Weight","Sex"), units=list(x="hr", y="mg/L"), covariates=list(c("kg","-")), name.X="Time")

model1cpt<-function(psi,id,xidep) {
dose<-xidep[,1]
tim<-xidep[,2]
ka<-psi[id,1]
V<-psi[id,2]
CL<-psi[id,3]
k<-CL/V
ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
return(ypred)
}

# Default model, no covariate
saemix.model<-saemixModel(model=model1cpt, description="One-compartment model with first-order absorption", psi0=matrix(c(1.,20,0.5,0.1,0.01),ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))), transform.par=c(1,1,1))

# Note: remove the options save=FALSE and save.graphs=FALSE
# to save the results and graphs
saemix.options<-list(seed=632545, save=FALSE, save.graphs=FALSE, displayProgress=FALSE)

# Not run (strict time constraints for CRAN)
saemix.fit<-saemix(saemix.model, saemix.data, saemix.options)

# Model with covariates
saemix.model<-saemixModel(model=model1cpt, description="One-compartment model with first-order absorption", psi0=matrix(c(1.,20,0.5,0.1,0.01),ncol=3,byrow=TRUE, dimnames=list(NULL, c("ka","V","CL"))), transform.par=c(1,1,1), covariate.model=matrix(c(0,0,1,0,0,0),ncol=3,byrow=TRUE), fixed.estim=c(1,1,1), covariance.model=matrix(c(1,0,0,1,1,0,0,1,1),ncol=3,byrow=TRUE), omega.init=matrix(c(1,0,0,0,1,1,0,1,1),ncol=3,byrow=TRUE), error.model="combined")

saemix.options<-list(seed=39546, save=FALSE, save.graphs=FALSE, displayProgress=FALSE)

# Not run (strict time constraints for CRAN)
saemix.fit<-saemix(saemix.model, saemix.data, saemix.options)
Description

The toenail.saemix data are from a multicenter study comparing two oral treatments for toe-nail infection, including information for 294 patients measured at 7 weeks, comprising a total of 1908 measurements. The outcome binary variable "onycholysis" indicates the degree of separation of the nail plate from the nail-bed (none or mild versus moderate or severe). Patients were evaluated at baseline (week 0) and at weeks 4, 8, 12, 24, 36, and 48 thereafter.

Usage
toenail.saemix

Format

This data frame contains the following columns:

id  subject index in file
time time of measurement (in months)
y  degree of onycholysis (0 if none or mild, 1 if moderate or severe)
treatment treatment indicator (1=Treatment A, 0= Treatment B)
visit visit number (visit numbers 1-7 correspond to scheduled visits at 0, 4, 8, 12, 24, 36, and 48 weeks)

#'

Details

The data in the toenail.saemix was copied from the Toenail dataset provided by the prLogistic package. Different models and analyses have been performed to describe this dataset in Molenberg and Verbeke (2000). Please refer to the PDF documentation (chapter 4, section 4.6) for more details on the analysis, including how to obtain diagnostic plots.

Source

prLogistic package in R

References


Examples

data(toenail.saemix)
saemix.data<-saemixData(name.data=toenail.saemix, name.group=c("id"), name.predictors=c("time","y"), name.response="y", name.covariates=c("treatment"),name.X=c("time"))
binary.model<-function(psi,id,xidep) {
  tim<-xidep[,1]
  y<-xidep[,2]
  inter<-psi[id,1]
  slope<-psi[id,2]
  logit<-inter+slope*tim
  pevent<-exp(logit)/(1+exp(logit))
  pobs = (y==0)*(1-pevent)+(y==1)*pevent
  logpdf <- log(pobs)
  return(logpdf)
}
saemix.model<-saemixModel(model=binary.model,description="Binary model", modeltype="likelihood",
psi0=matrix(c(-5,-.1,0,0),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1","theta2"))),
transform.par=c(0,0), covariate.model=c(0,1),
covariance.model=matrix(c(1,0,0,1),ncol=2))
saemix.options<-list(seed=1234567,save=FALSE,save.graphs=FALSE, displayProgress=FALSE,
  nb.chains=10, fim=FALSE)
binary.fit<-saemix(saemix.model,saemix.data,saemix.options)
plot(binary.fit, plot.type="convergence")

transform

Transform covariates

Description

Transform and/or center a vector

Usage

## S3 method for class 'numeric'
transform(
  _data,
  transformation = function(x) x,
  centering = "median",
  verbose = FALSE,
  ...
)
Transform covariates

transformCatCov

Arguments

_data a vector with values of type numeric

transformation transformation function. Defaults to no transformation

centering string, giving the value used to center the covariate; can be "mean" or "median", in which case this value will be computed from the data, 'none' or 0 for no centering, or a value given by the user. Defaults to the median value over the dataset.

verbose a boolean, prints messages during the execution of the function if TRUE. Defaults to FALSE.

... unused, for consistency with the generic method

Value

a vector

Examples

# TODO

transformCatCov

Transform covariates

Description

Regroup categorical covariates

Usage

transformCatCov(object, covariate, group, reference, verbose = FALSE)

Arguments

object saemixData object

covariate name of the covariate

group a vector giving the categories to which the initial values of the covariates should be mapped. If the resulting covariate is binary, it will be stored as 0/1. If it has more than 2 categories, dummy covariates will be created for the analysis.

reference the reference group

verbose a boolean, prints messages during the execution of the function if TRUE. Defaults to FALSE.

Value

an object of class "SaemixData"
transformContCov

Transform covariates

Description
Transform and/or center continuous covariates

Usage
transformContCov(
  object,  
  covariate, 
  transformation = function(x) x, 
  centering = "median", 
  verbose = FALSE
)

Arguments
- object: saemixData object
- covariate: name of the covariate
- transformation: transformation function. Defaults to no transformation
- centering: string, giving the value used to center the covariate; can be "mean" or "median", in which case this value will be computed from the data, 'none' or 0 for no centering, or a value given by the user. Defaults to the median value over the dataset.
- verbose: a boolean, prints messages during the execution of the function if TRUE. Defaults to FALSE.

Value
an object of class "SaemixData"

Examples
# TODO
validate.covariance.model

Validate the structure of the covariance model

Description

Check that a matrix corresponds to a structure defining a covariance model for a non-linear mixed effect model. Such a matrix should be composed of only 0s and 1s, with at least one element set to 1, and should be square and symmetrical. 1s on the diagonal indicate that the corresponding parameter has interindividual variability and that its variance will be estimated. 1s as off-diagonal elements indicate that a covariance between the two corresponding parameters will be estimated.

Usage

validate.covariance.model(x, verbose = TRUE)

Arguments

x a matrix
verbose a boolean indicating whether warnings should be output if x is not a valid covariance model

Value

a boolean, TRUE if x is an acceptable structure and FALSE if not. Messages will be output to describe why x isn’t a valid covariance model if the argument verbose is TRUE.

Author(s)

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See Also

SaemixModel

Examples

covarmodel<-diag(c(1,0))
validate.covariance.model(covarmodel) # should return TRUE
validate.names  

Description

Helper function, checks if the names given by the user match to the names in the dataset. If not, automatic recognition is attempted when automatic=TRUE.

Arguments

- **usernames**  
  vector of strings

- **datanames**  
  vector of strings

- **recognisednames**  
  vector of strings, values for automatic recognition

- **verbose**  
  logical, whether to print warning messages

- **automatic**  
  a boolean indicating whether to attempt automatic name recognition when some column names are missing or wrong (defaults to TRUE)

Value

a vector with valid names

Examples

# TODO

vcov

Description

Extracts the Variance-Covariance Matrix for a Fitted Model Object

Usage

```r
## S3 method for class 'SaemixRes'
vcov(object, ...)
```

```r
## S3 method for class 'SaemixObject'
vcov(object, ...)
```

```r
## S3 method for class 'SaemixObject'
vcov(object, ...)
```
**Arguments**

- **object** a fitted object from a call to saemix
- ... further arguments to be passed to or from other methods

**Value**

A matrix of the estimated covariances between the parameter estimates in model. In saemix, this matrix is obtained as the inverse of the Fisher Information Matrix computed by linearisation.

---

### xbinning

**Description**

Functions used by plot functions to define the boundaries of the bins on the X-axis.

**Usage**

```r
xbinning(xvec, plot.opt, verbose = FALSE)
```

**Arguments**

- **xvec** a vector of values for the X-axis of the plot
- **plot.opt** graphical options
- **verbose** boolean (defaults to FALSE). If TRUE, a table showing how the binning was performed

**Details**

These functions are normally not called by the end-user.

**Value**

a list with 3 elements, xgrp (the number of the bin associated with each element of xvec), xcent (a named vector containing the mean of the elements of xvec contained in each bin; the name of the bin is the interval), and xgroup (a vector with the group associated to each element of xvec after binning) If verbose is TRUE, a table showing the bins is shown, giving the interval of xvec associated with each bin, the mean value of xvec in each bin, and the number of observations

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yield.saemix

References

See Also
npde, autonpde

<table>
<thead>
<tr>
<th>yield.saemix</th>
<th>Wheat yield in crops treated with fertiliser, in SAEM format</th>
</tr>
</thead>
</table>

Description
The yield.saemix contains data from winter wheat experiments.

Usage
yield.saemix

Format
This data frame contains the following columns:

- **site**: the site number
- **dose**: dose of nitrogen fertiliser (kg/ha)
- **yield**: grain yield (kg/ha)
- **soil.nitrogen**: end-of-winter mineral soil nitrogen (NO3- plus NH4+) in the 0 to 90 cm layer was measured on each site/year (kg/ha)

Details
The data in the yield.saemix comes from 37 winter wheat experiments carried out between 1990 and 1996 on commercial farms near Paris, France. Each experiment was from a different site. Two soil types were represented, a loam soil and a chalky soil. Common winter wheat varieties were used. Each experiment consisted of five to eight different nitrogen fertiliser rates, for a total of 224 nitrogen treatments. Nitrogen fertilizer was applied in two applications during the growing season. For each nitrogen treatment, grain yield (adjusted to 150 g.kg-1 grain moisture content) was measured. In addition, end-of-winter mineral soil nitrogen (NO3- plus NH4+) in the 0 to 90 cm layer was measured on each site-year during February when the crops were tillering. Yield and end-of-winter mineral soil nitrogen measurements were in the ranges 3.44-11.54 t.ha-1, and 40-180 kg.ha-1 respectively.

Source
Examples

data(yield.saemix)

saemix.data<-saemixData(name.data=yield.saemix, header=TRUE, name.group=c("site"),
    name.predictors=c("dose"), name.response=c("yield"),
    name.covariates=c("soil.nitrogen"), units=list(x="kg/ha", y="t/ha", covariates=c("kg/ha")))

# Model: linear + plateau
yield.LP<-function(psi,id,xidep) {
  x<-xidep[,1]
  ymax<-psi[id,1]
  xmax<-psi[id,2]
  slope<-psi[id,3]
  f<-ymax+slope*(x-xmax)
  # cat(length(f)," ",length(ymax),"\n")
  f[x>xmax]<-ymax[x>xmax]
  return(f)
}

saemix.model<-saemixModel(model=yield.LP, description="Linear plus plateau model",
    psi0=matrix(c(8,100,0.2,0,0,0), ncol=3, byrow=TRUE, dimnames=list(NULL,
        c("Ymax","Xmax","slope")), covariance.model=matrix(c(1,0,0,0,1,0,0,0,1), ncol=3, byrow=TRUE),
        error.model="constant")
)

saemix.data$site<-

saemix.options<-list(algorithms=c(1,1,1), nb.chains=1, seed=666,
    save=FALSE, save.graphs=FALSE, displayProgress=FALSE)

# Plotting the data
plot(saemix.data,xlab="Fertiliser dose (kg/ha)", ylab="Wheat yield (t/ha)")

saemix.fit<-saemix(saemix.model, saemix.data, saemix.options)

# Comparing the likelihoods obtained by linearisation and importance sampling
# to the likelihood obtained by Gaussian Quadrature
saemix.fit<-llgq.saemix(saemix.fit)
{
  cat("LL by Importance sampling, LL_IS=", saemix.fit["results"]["ll.is"], "\n")
  cat("LL by linearisation, LL_lin=", saemix.fit["results"]["ll.lin"], "\n")
  cat("LL by Gaussian Quadrature, LL_GQ=", saemix.fit["results"]["ll.gq"], "\n")
}

# Testing for an effect of covariate soil.nitrogen on Xmax
saemix.model2<-saemixModel(model=yield.LP, description="Linear plus plateau model",
    psi0=matrix(c(8,100,0.2,0,0,0), ncol=3, byrow=TRUE, dimnames=list(NULL,
        c("Ymax","Xmax","slope")), covariance.model=matrix(c(1,0,0,0,1,0,0,0,1), ncol=3, byrow=TRUE),
        error.model="constant")
)

saemix.fit2<-saemix(saemix.model2, saemix.data, saemix.options)

# BIC for the two models
{
  cat("Model without covariate, BIC=", saemix.fit["results"]["bic.is"], "\n")
$$\text{Model with covariate, BIC} = \text{saemix.fit2["results"]['bic.is'], }"$
$$
pval<-1-pchisq(-2*saemix.fit["results"]['ll.is']+2*saemix.fit2["results"]['ll.is'],1)
$$
cat("\text{LRT: p=}, pval,"\n")
$$
}

# @keywords datasets

---

### Get/set methods for SaemixData object

**Description**

Access slots of a SaemixData object using the object["slot"] format

**Usage**

```r
## S4 method for signature 'SaemixData'
x[i, j, drop]

## S4 method for signature 'SaemixRepData'
x[i, j, drop]

## S4 replacement method for signature 'SaemixRepData'
x[i, j] <- value

## S4 method for signature 'SaemixSimData'
x[i, j, drop]

## S4 replacement method for signature 'SaemixSimData'
x[i, j] <- value

## S4 replacement method for signature 'SaemixRes'
x[i, j] <- value
```

**Arguments**

- `x`  
  object

- `i`  
  element to be replaced

- `j`  
  element to replace with

- `drop`  
  whether to drop unused dimensions

- `value`  
  value to replace with
[,SaemixModel-method  Get/set methods for SaemixModel object

Description
Access slots of a SaemixModel object using the object["slot"] format

Usage
## S4 method for signature 'SaemixModel'
x[i, j, drop]

Arguments
x  object
i  element to be replaced
j  element to replace with
drop  whether to drop unused dimensions

[,SaemixObject-method  Get/set methods for SaemixObject object

Description
Access slots of a SaemixObject object using the object["slot"] format

Usage
## S4 method for signature 'SaemixObject'
x[i, j, drop]

Arguments
x  object
i  element to be replaced
j  element to replace with
drop  whether to drop unused dimensions
### [,SaemixRes-method](#,SaemixRes-method)

**Get/set methods for SaemixRes object**

**Description**

Access slots of a SaemixRes object using the object["slot"] format.

**Usage**

```r
## S4 method for signature 'SaemixRes'
x[i, j, drop]
```

**Arguments**

- `x`  
  - object
- `i`  
  - element to be replaced
- `j`  
  - element to replace with
- `drop`  
  - whether to drop unused dimensions
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