Package ‘Rsmlx’

March 4, 2022

Type Package

Title R Speaks 'Monolix'

Version 4.0.2

Description Provides methods for model building and model evaluation of mixed effects models using 'Monolix' <https://monolix.lixoft.com>. 'Monolix' is a software tool for nonlinear mixed effects modeling that must have been installed in order to use 'Rsmlx'. Among other tasks, 'Rsmlx' provides a powerful tool for automatic PK model building, performs statistical tests for model assessment, bootstrap simulation and likelihood profiling for computing confidence intervals. 'Rsmlx' also proposes several automatic covariate search methods for mixed effects models.

URL http://rsmlx.webpopix.org

SystemRequirements 'Monolix' (<https://monolix.lixoft.com>)

Depends R (>= 3.0.0)

Imports graphics, grDevices, utils, stats, MASS, ggplot2, gridExtra, dplyr

Suggests mlxR,

confintmlx.R correlationModelSelection.R
_covariateModelSelection.R covariateSearch.R
whichPKmodel.R writeDataSmlx.R

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NeedsCompilation no

Encoding UTF-8

LazyData true

RoxygenNote 7.1.2
Description

Generate replicates of the original data using random sampling with replacement. Population parameters are then estimated from each replicate.
Usage

```r
bootmlx(
  project,
  nboot = 100,
  dataFolder = NULL,
  parametric = FALSE,
  tasks = c(populationParameterEstimation = TRUE),
  settings = NULL
)
```

Arguments

- **project**: Monolix project
- **nboot**: [optional] number of bootstrap replicates (default=100)
- **dataFolder**: [optional] folder where already generated datasets are stored, e.g. `dataFolder="./dummy_project/boot/"` (default: data set are generated by `bootmlx`)
- **parametric**: [optional] boolean to define if parametric bootstrap is performed (new data is drawn from the model), (default: false)
- **settings**: [optional] a list of settings for the resampling and the results:
  - **N**: the number of individuals in each bootstrap data set (default value is the number of individuals in the original data set).
  - **newResampling**: boolean to generate the data sets again if they already exist (default=FALSE).
  - **covStrat**: a categorical covariate of the project. The original distribution of this covariate is maintained in each resampled data set if covStrat is defined (default=NULL). Notice that if the categorical covariate is varying within the subject (in case of IOV), it will not be taken into account.
  - **plot**: boolean to choose if the distribution of the bootstraped estimates is displayed (default = FALSE)
  - **level**: level of the bootstrap confidence intervals of the population parameters (default = 0.90)
  - **seed**: seed for the generation of the data sets (default = NA)

Value

a data frame with the bootstrap estimates

Examples

```r
## Not run:
# RsmlxDemo1 mlxtran is a Monolix project for modelling the PK of warfarin using a PK model
# with parameters ka, V, Cl.
```
buildAll                              Automatic complete statistical model building

Description

buildAll builds the complete statistical model by iteratively calling functions buildmlx and buildVar. Penalties can be either a custom penalty of the form gamma*(number of parameters), AIC (gamma=2) or BIC (gamma=log(N)).

Usage

buildAll(
  project,
  final.project = NULL,
  model = "all",
  paramToUse = "all",
  covToTest = "all",
  covToTransform = "none",
  center.covariate = FALSE,
  criterion = "BICc",
  linearization = FALSE,
  ll = T,
  pen.cov = 1,
  direction = NULL,
  steps = 1000,
  max.iter = 20,
  explor.iter = 2,
  seq.cov = FALSE,
  seq.corr = TRUE,
buildAll

seq.cov.iter = 0,
p.max = 0.1,
p.min = c(0.075, 0.05, 0.1),
print = TRUE,
nb.model = 1,
fix.param1 = NULL,
fix.param0 = NULL,
remove = T,
add = T,
delta = c(30, 5),
omega.set = NULL,
pop.set1 = NULL,
pop.set2 = NULL
)

Arguments

- **project**: a string: the initial Monolix project
- **final.project**: a string: the final Monolix project (default adds "_built" to the original project)
- **model**: components of the model to optimize c("residualError", "covariate", "correlation"), (default="all")
- **paramToUse**: list of parameters possibly function of covariates (default="all")
- **covToTest**: components of the covariate model that can be modified (default="all")
- **covToTransform**: list of (continuous) covariates to be log-transformed (default="none")
- **center.covariate**: TRUE/FALSE center the covariates of the final model (default=FALSE)
- **criterion**: penalization criterion to optimize c("AIC", "BIC", "BICc", gamma)
- **linearization**: TRUE/FALSE whether the computation of the likelihood is based on a linearization of the model (default=FALSE, deprecated)
- **ll**: TRUE/FALSE compute the observe likelihood and the criterion to optimize at each iteration
- **pen.cov**: multiplicative penalty term for the covariate model (default=1)
- **direction**: method for covariate search c("full", "both", "backward", "forward"), (default="full" or "both")
- **steps**: maximum number of iteration for stepAIC (default=1000)
- **max.iter**: maximum number of iterations (default=20)
- **explor.iter**: number of iterations during the exploratory phase (default=2)
- **seq.cov**: TRUE/FALSE whether the covariate model is built before the correlation model
- **seq.corr**: TRUE/FALSE whether the correlation model is built iteratively (default=TRUE)
- **seq.cov.iter**: number of iterations before building the correlation model (only when seq.cov=F, default=0)
- **p.max**: maximum p-value used for removing non significant relationships between covariates and individual parameters (default=0.1)
buildAll

p.min minimum p-values used for testing the components of a new model (default=c(0.075, 0.05, 0.1))

print TRUE/FALSE display the results (default=TRUE)

nb.model number of models to display at each iteration (default=1)

fix.param1 parameters with variability that cannot be removed (default=NULL)

fix.param0 parameters without variability that cannot be added (default=NULL)

remove try to remove random effects (default=T)

add try to add random effects (default=T)

delta maximum difference in criteria for testing a new model (default=c(30,5))

omega.set settings to define how a variance varies during iterations of SAEM

pop.set1 Monolix settings 1

pop.set2 Monolix settings 2

Details

See http://rsmlx.webpopix.org for more details.

Value

a new Monolix project with a new statistical model.

Examples

```r
## Not run:
# Build the complete statistical model using the default settings
r1 <- buildAll(project="warfarinPK_project.mlxtran")

# Force parameter Tlag to be fixed (no variability) and parameter Cl to vary
r2 <- buildAll(project="warfarinPK_project.mlxtran", fix.param0="Tlag", fix.param1="Cl")

# Estimate the log-likelihood by linearization of the model (faster)
r3 <- buildAll(project="warfarinPK_project.mlxtran", linearization=T)
```

## End(Not run)

# See http://rsmlx.webpopix.org/userguide/buildmlx/ for detailed examples of use of buildmlx
# Download the demo examples here: http://rsmlx.webpopix.org/installation
buildmlx

Automatic statistical model building

Description

buildmlx uses SAMBA (Stochastic Approximation for Model Building Algorithm), an iterative procedure to accelerate and optimize the process of model building by identifying at each step how best to improve some of the model components. This method allows to find the optimal statistical model which minimizes some information criterion in very few steps.

Usage

buildmlx(
  project = NULL,
  final.project = NULL,
  model = "all",
  paramToUse = "all",
  covToTest = "all",
  covToTransform = "none",
  center.covariate = FALSE,
  criterion = "BICc",
  linearization = FALSE,
  ll = T,
  pen.cov = 1,
  direction = NULL,
  steps = 1000,
  max.iter = 20,
  explor.iter = 2,
  seq.cov = FALSE,
  seq.cov.iter = 0,
  seq.corr = TRUE,
  p.max = 0.1,
  p.min = c(0.075, 0.05, 0.1),
  print = TRUE,
  nb.model = 1
)

Arguments

project a string: the initial Monolix project
final.project a string: the final Monolix project (default adds "_.built" to the original project)
model components of the model to optimize c("residualError", "covariate", "correlation"), (default="all")
paramToUse list of parameters possibly function of covariates (default="all")
covToTest components of the covariate model that can be modified (default="all")
**buildmlx**

- **covToTransform**: list of (continuous) covariates to be log-transformed (default="none")
- **center.covariate**: TRUE/FALSE center the covariates of the final model (default=FALSE)
- **criterion**: penalization criterion to optimize c("AIC", "BIC", "BICc", gamma)
- **linearization**: TRUE/FALSE whether the computation of the likelihood is based on a linearization of the model (default=FALSE)
- **ll**: TRUE/FALSE compute the observe likelihood and the criterion to optimize at each iteration
- **pen.cov**: multiplicative penalty term for the covariate model (default=1)
- **direction**: method for covariate search c("full", "both", "backward", "forward"), (default="full" or "both")
- **steps**: maximum number of iteration for stepAIC (default=1000)
- **max.iter**: maximum number of iterations (default=20)
- **explor.iter**: number of iterations during the exploratory phase (default=2)
- **seq.cov**: TRUE/FALSE whether the covariate model is built before the correlation model
- **seq.cov.iter**: number of iterations before building the correlation model (only when seq.cov=F, default=0)
- **seq.corr**: TRUE/FALSE whether the correlation model is built iteratively (default=TRUE)
- **p.max**: maximum p-value used for removing non significant relationships between covariates and individual parameters (default=0.1)
- **p.min**: vector of 3 minimum p-values used for testing the components of a new model (default=c(0.075, 0.05, 0.1))
- **print**: TRUE/FALSE display the results (default=TRUE)
- **nb.model**: number of models to display at each iteration (default=1)

**Details**

Penalization criterion can be either a custom penalization of the form \( \gamma \times \text{(number of parameters)} \), AIC (\( \gamma = 2 \)) or BIC (\( \gamma = \text{log}(N) \)).

Several strategies can be used for building the covariate model at each iteration of the algorithm: direction="full" means that all the possible models are compared (default when the number of covariates is less than 10). Otherwise, direction is the mode of stepwise search of stepAIC (MASS), can be one of "both", "backward", or "forward", with a default of "both" when there are at least 10 covariates. See http://rsmlix.webpopix.org for more details.

**Value**

a new Monolix project with a new statistical model.
Examples

# RsmlxDemo1.mlxtran is a Monolix project for modelling the pharmacokinetics (PK) of warfarin
# using a PK model with parameters ka, V, Cl.

# By default, buildmlx will compute the best statistical model in term of BIC, i.e,
# the best covariate model, the best correlation model for the three random effects and the best
# residual error model in terms of BIC.
# In this example, three covariates (wt, age, sex) are available with the data and will be used
# for building the covariate model for the three PK parameters:
# r1 <- buildmlx(project="RsmlxDemo1.mlxtran")

# Here, the covariate model will be built for V and Cl only and log-transformation of all
# continuous covariates will also be considered:
# r2 <- buildmlx(project="RsmlxDemo1.mlxtran", paramToUse=c("V", "Cl"), covToTransform="all")

# Only the covariate model will be built, using AIC instead of BIC:
# r3 <- buildmlx(project="RsmlxDemo1.mlxtran", model="covariate", criterion="AIC")

# See http://rsmlx.webpopix.org/userguide/buildmlx/ for detailed examples of use of buildmlx
# Download the demo examples here: http://rsmlx.webpopix.org/installation

buildVar

Automatic model variance building

Description

buildVar is designed to build the best variance model for the random effects by selecting which
individual parameters vary and which ones are fixed.

Usage

buildVar(
  project,
  final.project = NULL,
  fix.param1 = NULL,
  fix.param0 = NULL,
  criterion = "BICc",
  linearization = F,
  remove = T,
  add = T,
  delta = c(30, 5),
  omega.set = NULL,
  pop.set1 = NULL,
  pop.set2 = NULL,
  print = TRUE
)
**Arguments**

- **project**: a string: the initial Monolix project
- **final.project**: a string: the final Monolix project (default adds "_var" to the original project)
- **fix.param1**: parameters with variability that cannot be removed (default=NULL)
- **fix.param0**: parameters without variability that cannot be added (default=NULL)
- **criterion**: penalization criterion to optimize c("AIC", "BIC", "BICc", gamma)
- **linearization**: TRUE/FALSE whether the computation of the likelihood is based on a linearization of the model (default=FALSE)
- **remove**: try to remove random effects (default=T)
- **add**: try to add random effects (default=T)
- **delta**: maximum difference in criteria for testing a new model (default=c(30,5))
- **omega.set**: settings to define how a variance varies during iterations of SAEM
- **pop.set1**: Monolix settings 1
- **pop.set2**: Monolix settings 2
- **print**: TRUE/FALSE display the results (default=TRUE)

**Details**

Penalization criterion can be either a custom penalization of the form gamma*(number of parameters), AIC (gamma=2) or BIC (gamma=log(N)).

See [http://rsmlx.webpopix.org](http://rsmlx.webpopix.org) for more details.

**Value**

a new Monolix project with a new inter individual variability model.

**Examples**

```r
## Not run:
# Build the variability model using the default settings
r1 <- buildVar(project="warfarinPK_project.mlxtran")

# Force parameter Tlag to be fixed (no variability) and parameter Cl to vary
r2 <- buildVar(project="warfarinPK_project.mlxtran", fix.param0="Tlag", fix.param1="Cl")

# Estimate the log-likelihood by linearization of the model (faster)
r3 <- buildVar(project="warfarinPK_project.mlxtran", linearization=T)

## End(Not run)
```

# See [http://rsmlx.webpopix.org/userguide/buildvar/](http://rsmlx.webpopix.org/userguide/buildvar/) for detailed examples of use of buildvar
# Download the demo examples here: [http://rsmlx.webpopix.org/installation](http://rsmlx.webpopix.org/installation)
Description

Compute confidence intervals for the population parameters estimated by Monolix.

Usage

confintmlx(
  project,
  parameters = "all",
  method = "fim",
  level = 0.9,
  linearization = TRUE,
  nboot = 100,
  parametric = FALSE,
  settings = NULL
)

Arguments

- **project**: a Monolix project
- **parameters**: list of parameters for which confidence intervals are computed (default="all")
- **method**: method c("fim", "proflike", "bootstrap")
- **level**: confidence level, a real number between 0 and 1 (default=0.90)
- **linearization**: TRUE/FALSE whether the calculation of the standard errors or the profile likelihood is based on a linearization of the model (default=TRUE)
- **nboot**: number of bootstrap replicates (default=100, used when method="bootstrap")
- **parametric**: boolean to define if parametric bootstrap is performed (new data is drawn from the model), (default: false)
- **settings**: a list of settings for the profile likelihood method:
  - **max.iter**: maximum number of iterations to find the solution (default=10)
  - **tol.LL**: absolute tolerance for -2LL (default=0.001)
  - **tol.param**: relative tolerance for the parameter (default=0.01)
  - **print**: TRUE/FALSE display the results (default=TRUE)

Details

The method used for computing the confidence intervals can be either based on the standard errors derived from an estimation of the Fisher Information Matrix ("fim"), on the profile likelihood ("proflike") or on nonparametric bootstrap estimate ("bootstrap"). method="fim" is used by default.

When method="fim", the FIM can be either estimated using a linearization of the model or a stochastic approximation. When method="proflike", the observed likelihood can be either estimated using a linearization of the model or an importance sampling Monte Carlo procedure. When method="bootstrap", the bootstrap estimates are obtained using the bootmlx function.
Value

A list with the computed confidence intervals, the method used and the level.

Examples

```r
# RsmlxDemo2.mltrans is a Monolix project for modelling the PK of warfarin using a PK model
# with parameters ka, V, Cl.

# confintmlx will compute a 90% confidence interval for all the population parameters
# using the population estimates obtained by Monolix and the Fisher Information Matrix
# estimated by linearization
r1 <- confintmlx(project="RsmlxDemo2.mltrans")

# 95% confidence intervals are now computed, using the FIM estimated by Monolix using a
# stochastic approximation algorithm:
r2 <- confintmlx(project="RsmlxDemo2.mltrans", linearization=FALSE, level=0.95)

# Confidence intervals are computed for ka_pop and omega_ka only,
# using the profile likelihood method:
r <- confintmlx(project = "RsmlxDemo2.mltrans",
method = "proflike",
parameters = c("ka_pop","omega_ka"))

# Confidence intervals are computed using 200 bootstrap samples:
r3 <- confintmlx(project="RsmlxDemo2.mltrans", method="bootstrap", nboot=200)

# See http://rsmlx.webpopix.org/userguide/confintmlx/ for detailed examples of use of confintmlx
# Download the demo examples here: http://rsmlx.webpopix.org/installation
```

covariateSearch  Covariate model building

Description

Automatic search of the best covariate model. Two methods for covariate model building are proposed:

- SCM: stepwise covariate modeling method. In the forward selection, at each step, each of the remaining (i.e. not yet included) parameter-covariate relationships are added to the model in an univariate model (one model per relationship), and run. Among all models, the model that improves some criteria (LRT, BIC or AIC) most is selected and taken forward to the next step. During backward elimination, parameter-covariate relationships are removed in an univariate manner.

- COSSAC: Conditional Sampling for Stepwise Approach based on Correlation tests method. COSSAC makes use of the information contained in the base model run to choose which covariate to try first (instead of trying all covariates "blindly" as in SCM). Indeed, the correlation between the individual parameters (or random effects) and the covariates hints at possibly
relevant parameter-covariate relationships. If the EBEs (empirical Bayes estimates) are used, shrinkage may bias the result. COSSAC instead uses samples from the a posteriori conditional distribution (available as “conditional distribution” task in MonolixSuite2018) to calculate the correlation between the random effects and covariates. A p-value can be derived using the Pearson’s correlation test for continuous covariate and ANOVA for categorical covariate. The p-values are used to sort all the random effect-covariate relationships. Relationships with the lowest p-value are added first, run and confirmed using a likelihood ratio test, AIC or BIC criteria.

Usage

covariateSearch(
  project,
  final.project = NULL,
  method = NULL,
  covToTest = NULL,
  covToTransform = NULL,
  paramToUse = NULL,
  testRelations = NULL,
  settings = NULL
)

Arguments

project  a Monolix project
final.project  [optional] string corresponding to the final Monolix project (default: 'runFinal.mlxtran' in covariate search output folder)
method  [optional] string correspondig to the method. It can be 'COSSAC' or 'SCM'. By default, COSSAC' is used.
covToTest  [optional] vector of covariates to test. Cannot be used if testRelations is defined. By default, all covariates are tested.
covToTransform  [optional] vector of covariates to transform. The transformation consists in a log transform of the covariate with centering by the mean value (ex: WT is transformed into log(WT/mean) with mean the mean WT value over the individuals of the data set). Both the transformed and untransformed covariate are tested by the algorithm. By default, no covariate is transformed. Note: adding a non-transformed covariate on a lognormally distributed parameter results in an exponential relationship: log(V) = log(Vpop) + beta*WT + eta <=> V = Vpop * exp(beta*WT) * exp(eta) adding a log-transformed covariate on a lognormally distributed parameter results in a power law relationship: log(V) = log(Vpop) + beta*log(WT/70) + eta <=> V = Vpop * (WT/70)^beta * exp(eta)
paramToUse  [optional] vector of parameters which may be function of covariates. Cannot be used if testRelations is defined. By default, all parameters are tested.
testRelations  [optional] list of parameter-covariate relationships to test, ex: list(V=c("WT","SEX"),Cl=c("CRCL")). Cannot be used if covToTest or paramToUse is defined. By default, all parameter-covariate relationships are tested.
settings  [optional] list of settings for the covariate search:
getEstimatedCovarianceMatrix

Get estimated covariance and correlation matrices

Description

Get estimated covariance and correlation matrices

Examples

# RsmlxDemo1.mlxtran is a Monolix project for modelling the pharmacokinetics (PK) of warfarin
# using a PK model with parameters ka, V, Cl.

# In this example, three covariates (wt, age, sex) are available with the data
# covariatesearch will compute the best covariate model, in term of BIC,
# for the three PK parameters using the three covariates.
r1 <- covariateSearch(project="RsmlxDemo1.mlxtran")

# Instead of using the COSSAC method, we can use the SCM method:
r2 <- covariateSearch(project="RsmlxDemo1.mlxtran", method = 'SCM')

# Here, the covariate model is built using age and wt only, for V and Cl only:
r3 <- covariateSearch(project = "RsmlxDemo1.mlxtran",
                      paramToUse = c("V","Cl"),
                      covToTest = c("age","wt"))

# See http://rsmlx.webpopix.org/userguide/covariatesearch/ for detailed examples of covariatesearch
# Download the demo examples here: http://rsmlx.webpopix.org/installation
Usage

getEstimatedCovarianceMatrix()

Value

a list of two matrices.

Examples

```r
## Not run:
# Assume that the Monolix project "warfarinPKPD.mlxtran" has been loaded
r = getEstimatedCovarianceMatrix() # r is a list with elements "cor.matrix" and "cov.matrix"

# See http://rsmlx.webpopix.org/userguide/newconnectors/ for more detailed examples
# Download the demo examples here: http://rsmlx.webpopix.org/installation

## End(Not run)
```

---

getEstimatedIndividualParameters2

*Get estimated individual and population parameters*

Description

Get the individual individual parameters, the population parameters with the population covariates and the population parameters with the individual covariates.

Usage

getEstimatedIndividualParameters2()

Value

a list of data frames.

Examples

```r
## Not run:
# Assume that the Monolix project "warfarinPKPD.mlxtran" has been loaded
r = getEstimatedIndividualParameters2()

# r is a list with elements "saem", "conditionalMean", "conditionalSD", "conditionalMode",
# "popPopCov" and "popIndCov"

# See http://rsmlx.webpopix.org/userguide/newconnectors/ for more detailed examples
# Download the demo examples here: http://rsmlx.webpopix.org/installation

## End(Not run)
```
getEstimatedPredictions

Get estimated predictions

Description

Get the individual predictions obtained with the estimated individual parameters:

Usage

getEstimatedPredictions()

Value

a list of data frames (one data frame per output).

Examples

## Not run:
# Assume that the Monolix project "warfarinPKPD.mlxtran" has been loaded
r = getEstimatedPredictions() # r is a list with elements "y1" and "y2"

# See http://rsmlx.webpopix.org/userguide/newconnectors/ for more detailed examples
# Download the demo examples here: http://rsmlx.webpopix.org/installation

## End(Not run)

getEstimatedResiduals

Get estimated residuals

Description

Get the residuals computed from the individual predictions obtained with the estimated individual parameters:

Usage

getEstimatedResiduals()

Value

a list of data frames (one data frame per output).
getSimulatedPredictions

## Description

Get the individual predictions obtained with the simulated individual parameters:

## Usage

```r
getSimulatedPredictions()
```

## Value

A list of data frames (one data frame per output).

## Examples

```r
## Not run:
# Assume that the Monolix project "warfarinPKPD.mlxtran" has been loaded
r = getSimulatedPredictions() # r is a list with elements "y1" and "y2"

# See http://rsmlx.webpopix.org/userguide/newconnectors/ for more detailed examples
# Download the demo examples here: http://rsmlx.webpopix.org/installation

## End(Not run)
```

---

getSimulatedResiduals

## Description

Get the residuals computed from the individual predictions obtained with the simulated individual parameters:

## Usage

```r
getSimulatedResiduals()
```

## Value

A list of residuals (one residual per output).

## Examples

```r
## Not run:
# Assume that the Monolix project "warfarinPKPD.mlxtran" has been loaded
r = getSimulatedResiduals() # r is a list with elements "Cc" and "E"

# See http://rsmlx.webpopix.org/userguide/newconnectors/ for more detailed examples
# Download the demo examples here: http://rsmlx.webpopix.org/installation

## End(Not run)
```
Usage

getSimulatedResiduals()

Value

a list of data frames (one data frame per output).

Examples

## Not run:
# Assume that the Monolix project "warfarinPKPD.mlxtran" has been loaded
r = getSimulatedResiduals()  # r is a list with elements "y1" and "y2"

# See http://rsmlx.webpopix.org/userguide/newconnectors/ for more detailed examples
# Download the demo examples here: http://rsmlx.webpopix.org/installation

## End(Not run)

initRsmlx

**Initialize Rsmlx library**

Description

Initialize Rsmlx library

Usage

initRsmlx(path = NULL)

Arguments

path Monolix path

Value

A list:

- software: the software that is used (should be monolix with Rsmlx)
- path: the path to MonolixSuite
- version: the version of MonolixSuite that is used
- status: boolean equaling TRUE if the initialization has been successful.

Examples

## Not run:
initRsmlx()  # print the info about Monolix and lixoftConnectors
initRsmlx(path="C:/ProgramData/Lixoft/MonolixSuite2019R1")  # use MonolixSuite 2019R1

## End(Not run)
pkbuild

Automatic PK model building

Description

Fit several structural PK models and select the best one based on a Bayesian Information Criterion. Models to compare can be defined by rate constants and/or clearances and can include or not nonlinear elimination models. See http://rsmlx.webpopix.org/pkbuild/ for more details.

Usage

pkbuild(
  data = NULL,
  project = NULL,
  stat = FALSE,
  param = "clearance",
  new.dir = ".",
  MM = FALSE,
  level = NULL,
  settings.stat = NULL
)

Arguments

data a list with fields
  • dataFile: path of a formatted data file
  • headerTypes: a vector of strings
  • administration ("iv", "bolus", "infusion", "oral", "ev"): route of administration

project a Monolix project

stat (FALSE, TRUE): the statistical model is also built (using buildmlx)

param ("clearance", "rate", "both"): parameterization

new.dir name of the directory where the created files are stored (default is the current working directory)

MM (FALSE, TRUE): tested models include or not Michaelis Menten elimination models

level an integer between 1 and 9 (used by setSettings)

settings.stat list of settings used by buildmlx (only if stat=TRUE)

Value

A list of results
Examples

```r
## Not run:
# Build a PK model for the warfarin PK data.
# By default, only models using clearance (and inter compartmental clearances) are used.
warf.pk1 <- pkbuild(data=warfarin)

# Models using elimination and transfer rate constants are used,
# as well as nonlinear elimination models
warf.pk2 <- pkbuild(data=warfarin, new.dir="warfarin", param="rate", MM=TRUE)

# Both models using clearances and rates are used.
# Level is set to 7 in order to get accurate results.
warf.pk3 <- pkbuild(data=warfarin, new.dir="warfarin", param="both", level=7)

## End(Not run)
```

---

**pkpopini**  
*Compute initial population PK parameters*

Description

Use the pooled PK data to derive population PK parameters for a "standard" PK model (i.e. a model of the Monolix PK library). The structural model is automatically defined using the names of the PK parameters. Allowed names are: 'Tlag', 'Mtt', 'Ktr', 'ka', 'Tk0', 'V', 'V1', 'V2', 'V3', 'Q', 'Q2', 'Q3', 'Cl', 'k', 'k12', 'k21', 'k13', 'k31', 'Vm', 'Km'.

Usage

```r
pkpopini(
  data = NULL,
  project = NULL,
  parameter = NULL,
  new.project = NULL,
  new.dir = NULL,
  par.ini = NULL
)
```

Arguments

- `data`: a list with fields
  - `dataFile`: path to a formatted data file
  - `headerTypes`: a vector of strings
- `project`: a Monolix project
- `parameter`: a vector of strings (names of the PK parameters)
- `new.project`: name of the new Monolix project (a default name is created if not provided)
- `new.dir`: name of the directory where the created files are stored (default is the current working directory)
- `par.ini`: a vector of PK parameter values
Details

A Monolix project is then automatically created using these values as initial population parameters. See http://rsmlx.webpopix.org/pkpopini/ for more details.

Value

A list of results

Examples

```r
## Not run:
# Create in the working directory a Monolix project for a 1 cpt model with
# lag time, 0 order absorption and linear elimination
warf.ini1 <- pkpopini(data=warfarin, param=c("Tlag", "Tk0", "V", "Cl"))

# Create in directory 'warfarin' a Monolix project called 'warfPK2.mlxtran'
# for a 2 cpt model with 1st order absorption and nonlinear elimination
warf.ini3 <- pkpopini(data=warfarin, param=c("ka", "V", "k12", "k21", "Vm", "Km"),
                        new.dir="warfarin", new.project="warfPK2.mlxtran")
## End(Not run)
```

---

<table>
<thead>
<tr>
<th>resMonolix</th>
<th>Monolix results</th>
</tr>
</thead>
</table>

Description

Monolix results used by the Rsmlx examples

Usage

```r
resMonolix
```

Format

A R list

Source

Monolix demos

References

Rsmlx website: http://rsmlx.webpopix.org
RsmlxDemo2.project

RsmlxDemo1.project  Monolix project for warfarin PK - 1

Description
RsmlxDemo2.mlxtran is a Monolix project for modelling the pharmacokinetics (PK) of warfarin using a PK model with parameters ka, V, Cl. There is no covariate in the model.

Usage
RsmlxDemo1.project

Format
A text file

Source
Monolix project

References
Rsmlx documentation

RsmlxDemo2.project  Monolix project for warfarin PK - 2

Description
RsmlxDemo2.mlxtran is a Monolix project for modelling the pharmacokinetics (PK) of warfarin using a PK model with parameters ka, V, Cl. Here, V and Cl are function of weight.

Usage
RsmlxDemo2.project

Format
A text file

Source
Monolix project

References
Rsmlx documentation
**setSettings**

*Easy tuning of the settings of a Monolix project*

**Description**

Use a single accuracy level, between 1 and 9, to automatically tune all the settings of a Monolix project. When the accuracy level is equal to 1, the algorithms are very fast but the results may be not precise. When the accuracy level is equal to 9, the algorithms are slow but the results are accurate. Default Monolix settings are obtained with level=5.

**Usage**

```r
setSettings(project = NULL, new.project = NULL, level = 5)
```

**Arguments**

- **project**: a string: a Monolix project (the loaded project if NULL)
- **new.project**: a string: the new created Monolix project (default is the original project)
- **level**: an integer between 1 and 9 (default=5)

**Examples**

```r
# Not run:
# RsmlxDemo1.mlxtran is a Monolix project for modelling the PK of warfarin.
# All settings of the project are set so that algorithms used by Monolix converge as
# quickly as possible possible:
setSettings(project="RsmlxDemo1.mlxtran", level=1)

# A new project will be created with settings set in order to obtain the most
# precise results possible:
new.project= file.path(tempdir(),"RsmlxDemoNew.mlxtran")
setSettings(project="RsmlxDemo1.mlxtran", new.project=new.project, level=9)

# See http://rsmlx.webpopix.org/userguide/setSettings/ for detailed examples of use of setSettings
# Download the demo examples here: http://rsmlx.webpopix.org/installation
```

---

**testmlx**

*Statistical tests for model assessment*

**Description**

Perform several statistical tests using the results of a Monolix run to assess the statistical components of the model in use.
Usage

testmlx(
    project = NULL,
    tests = c("covariate", "randomEffect", "correlation", "residual"),
    plot = FALSE,
    adjust = "edf",
    n.sample = NULL
)

Arguments

project a Monolix project

tests a vector of strings: the list of tests to perform among c("covariate","randomEffect","correlation","residual")

plot FALSE/TRUE display some diagnostic plots associated to the tests (default=FALSE)

adjust method to take into account the dependency of MCMC sample c("edf","BH")

n.sample number of samples from the conditional distribution to be used (default = number of available samples in the project)

Details

The tests used are: 1) F-tests (or, equivalently, correlation tests) to evaluate the effect of each covariate on each parameter ("covariate"), 2) correlation tests to assess the correlation structure of the random effects ("correlation"), 3) Shapiro-Wilk and Miao-Gel-Gastwirth tests to assess, respectively the normality and the symmetry of the distribution of the random effects ("randomEffect"), 4) Shapiro-Wilk and Miao-Gel-Gastwirth tests to assess, respectively the normality and the symmetry of the distribution of residual errors ("residual"). By default, the four tests are performed.

When several samples of the conditional distributions are used, two methods are proposed in order to take into the dependance of the samples for the Shapiro-Wilk and Miao-Gel-Gastwirth tests: "edf" computes an effective degrees of freedom, "BH" performs one test per replicates and adjust the smallest p-value using the Benjamini-Hochberg correction.

Value

a list of data frames and ggplot objects if plot=TRUE

Examples

# RsmlxDemo2.mlxtran is a Monolix project for modelling the PK of warfarin using a PK model with parameters ka, V, Cl.

#testmlx will perform statistical tests for the different component of the statistical model:
r1 <- testmlx(project="RsmlxDemo2.mlxtran")

#testmlx will perform statistical tests for the covariate model and the correlation model only.
r2 <- testmlx(project="RsmlxDemo2.mlxtran", tests=c("covariate","correlation"))

# See http://rsmlx.webpopix.org/userguide/testmlx/ for detailed examples of use of testmlx
warfarin.data

# Download the demo examples here: http://rsmlx.webpopix.org/installation

---

### warfarin.data  
**warfarin PKPD data**

**Description**

The warfarin PK and PD data for 32 patients

**Usage**

```r
warfarin.data
```

**Format**

A csv file

**Source**

Monolix demos

**References**


---

### whichPKmodel  
**Find a Monolix PK model**

**Description**

Return the path of the Monolix PK model defined by a list of parameter names See [http://rsmlx.webpopix.org/whichPKmodel/](http://rsmlx.webpopix.org/whichPKmodel/) for more details.

**Usage**

```r
whichPKmodel(parameter, mlxPath = NULL, pkPath = NULL, lib = FALSE)
```

**Arguments**

- `parameter`  
a vector of PK parameter names
- `mlxPath`  
path to Monolix install
- `pkPath`  
path to the Monolix PK library
- `lib`  
boolean to define if the absolute path is returned
### Examples

```r
## Not run:
whichPKmodel(parameter=c("Tlag", "Tk0", "V", "Cl"))
## End(Not run)
```

---

#### writeDataSmlx

Write Simulx Dataset

#### Description

Format outputs of simulx simulations and write datasets in monolix and pkanalix project format.

#### Usage

```r
writeDataSmlx(
  filename = "simulated_dataset.csv",
  sep = ",",
  ext = "csv",
  nbdigits = 5,
  mapObservation = NULL
)
```

#### Arguments

- `filename` *(string) (optional)*: file path to dataset. (default "simulated_dataset.csv") In case of multiple replicates, the function creates one dataset per replicate with name `filename_repi` If filename contains an extension, it must be "csv" or "txt". If it does not, extension is defined by `ext` argument.
- `sep` *(string) (optional)*: Separator used to write dataset file. (default ",," ) It must be one of ",", ",", ",", ",".
- `ext` *(bool) (optional)*: Extension used to write dataset file. (default "csv") It must be one of "csv", "txt" To defined only if filename with no extension
- `nbdigits` *(integer) (optional)*: number of decimal digits in output file. (default = 5)
- `mapObservation` *(name vector) (optional)*: mapping of observation name

#### Details

WARNING: ‘writeData’ function is not implemented for simulx project with regressors in MonolixSuite version 2020R1

#### Value

- a dataframe if one single simulation, a list of dataframe if multiple replicates.
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