

Package ‘mlxR’

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Type Package

Version 4.2.0

Title Simulation of Longitudinal Data

Description Simulation and visualization of complex models for longitudinal data. The models are encoded using the model coding language 'Mlxtran' and automatically converted into C++ codes. That allows one to implement very easily complex ODE-based models and complex statistical models, including mixed effects models, for continuous, count, categorical, and time-to-event data.

URL <http://simulx.webpopix.org>

BugReports <https://github.com/MarcLavielle/mlxR/issues>

Depends R (>= 3.0.1), ggplot2

Suggests XML, Rcpp (>= 0.11.3), reshape2, gridExtra, shiny

Imports tools, methods, graphics, grDevices, utils, stats

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NeedsCompilation no

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Collate 'apiTools.R' 'apiManager.R' 'mlxComputeRCleaner.R' 'mlxComputeRLibraryBuilder.R' 'addFieldsFromHeader.R' 'catplotmlx.R' 'commentModel.R' 'convertmlx.R' 'exposure.R' 'generateModelFromPkModel.R' 'ggplotmlx.R' 'hformat.R' 'hgdata.R' 'inlineDataFrame.R' 'inlineModel.R' 'isfield.R' 'kmpplotmlx.R' 'mlxplore.R' 'monolix2simulx.R' 'pkmodel.R' 'prctilemlx.R' 'processing_monolix.R' 'processing_setting.R' 'processing_target.R' 'readVector.R' 'readdatamlx.R' 'shinymlx.R' 'simpopmlx.R' 'simulR.R' 'simulx.R' 'statmlx.R' 'toolsmlx.R' 'toolsmlx.R' 'translateIOV.R' 'translateIOVind.R' 'uuid.R' 'writeDatamlx.R'

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catplotmlx

Plot Categorical Longitudinal Data

Description

Plot the empirical distribution of categorical longitudinal data.

Usage

```
catplotmlx(
  r,
  col = NULL,
  breaks = NULL,
  plot = TRUE,
  color = "#194280",
  group = NULL,
```

```

    facet = TRUE,
    labels = NULL
  )

```

Arguments

<code>r</code>	a data frame with a column 'id', a column 'time', a column with values and possibly Hk[ja column 'group'.
<code>col</code>	a vector of 3 column numbers: ('id', 'time/x', 'y'. Default = c(1, 2,3).
<code>breaks</code>	one of: • a vector giving the breakpoints, • a single number giving the number of segments.
<code>plot</code>	if TRUE the empirical distribution is displayed, if FALSE the values are returned
<code>color</code>	a color to be used for the plots (default="#194280")
<code>group</code>	variable to be used for defining groups (by default, 'group' is used when it exists)
<code>facet</code>	makes subplots for different groups if TRUE
<code>labels</code>	vector of strings

Details

See <http://simulx.webpopix.org/mlxr/catplotmlx/> for more details.

Value

a ggplot object if plot=TRUE ; otherwise, a list with fields:

- color a vector of colors used for the plot
- y a data frame with the values of the empirical distribution computed at each time point

Examples

```

## Not run:
catModel <- inlineModel("
[LONGITUDINAL]
input = {a,b}
EQUATION:
lp1=a-b*t
lp2=a-b*t/2
DEFINITION:
y = {type=categorical, categories={1,2,3},
logit(P(y<=1))=lp1, logit(P(y<=2))=lp2}
")

y.out <- list(name='y', time=seq(0, 100, by=4))

Ng <- 1000
g1 <- list(size=Ng, parameter=c(a=6,b=0.2))

```

```

res <- simulx(model=catModel, output=y.out, group=g1)
catplotmlx(res$y)
catplotmlx(res$y, breaks=seq(-2,102,by=8), color="purple")
catplotmlx(res$y, breaks=5, color="#490917")

g2 <- list(size=Ng, parameter=c(a=10,b=0.2))
res <- simulx(model=catModel, output=y.out, group=list(g1,g2))
catplotmlx(res$y)
catplotmlx(res$y, group="none")

g3 <- list(size=Ng, parameter=c(a=6,b=0.4))
g4 <- list(size=Ng, parameter=c(a=10,b=0.4))
res <- simulx(model=catModel, output=y.out, group=list(g1,g2,g3,g4))
catplotmlx(res$y)

cov <- data.frame(id=levels(res$y$id), a=rep(c(6,10,6,10),each=Ng),
                b=rep(c(0.2,0.2,0.4,0.4),each=Ng))
catplotmlx(res$y, group=cov)

## End(Not run)

```

exposure

Computation of AUC, Cmax and Cmin

Description

Compute the area under the curve, the maximum and minimum values of a function of time over a given interval or at steady state

Usage

```

exposure(
  model = NULL,
  output = NULL,
  group = NULL,
  treatment = NULL,
  parameter = NULL,
  data = NULL,
  project = NULL,
  settings = NULL,
  regressor = NULL,
  varlevel = NULL
)

```

Arguments

model	a Mlxtran model used for the simulation
output	a list with fields:

	• name: a vector of output names • time: = 'steady.state' • ntp: number of time points used for computing the exposure (default=100) • tol: tolerance number, between 0 and 1, for approximating steady-state (default=0.01) • ngc: number of doses used for estimating the convergence rate to steady-state (default=5)
group	a list, or a list of lists, with fields: • size : size of the group (default=1), • level : level(s) of randomization, • parameter : if different parameters per group are defined, • output : if different outputs per group are defined, • treatment : if different treatments per group are defined, • regressor : if different regression variables per group are defined.
treatment	a list with fields • tfd : time of first dose (default=0), • ii : inter dose interval (mandatory), • amount : the amount for each dose, • rate : the infusion rate (default=Inf), • tinf : the time of infusion (default=0), • type : the type of input (default=1), • target : the target compartment (default=NULL).
parameter	a vector of parameters with their names and values
data	a list
project	the name of a Monolix project
settings	a list of optional settings • result.file : name of the datafile where the simulated data is written (string), • seed : initialization of the random number generator (integer), • load.design : TRUE/FALSE (if load.design is not defined, a test is automatically performed to check if a new design has been defined), • data.in : TRUE/FALSE (default=FALSE) • id.out : add columns id (when N=1) and group (when #group=1), TRUE/FALSE (default=FALSE) • Nmax : maximum group size used in a single call of mlxCompute (default=100)
regressor	a list, or a list of lists, with fields • name : a vector of regressor names, • time : a vector of times, • value : a vector of values.
varlevel	a list, or a list of lists, with fields • name : a vector of names of variability levels, • time : a vector of times that define the occasions.

Details

Input arguments are the input arguments of Simulx (<http://simulx.webpopix.org>)

Specific input arguments can be also used for computing the exposure at steady state, i.e. after the administration of an "infinite" number of doses. See <http://simulx.webpopix.org/exposure/> for more details.

Value

A list of data frames. One data frame per output is created with columns `id` (if number of subject >1), `group` (if number of groups >1), `t1` (beginning of time interval), `t2` (end of time interval), `step` (time step), `auc` (area under the curve), `tmax` (time of maximum value), `cmax` (maximum value), `tmin` (time of minimum value), `cmin` (minimum value).

<code>ggplotmlx</code>	<i>mlxR wrapper for ggplot</i>
------------------------	--------------------------------

Description

mlxR wrapper around [ggplot](#) with a custom theme

Usage

```
ggplotmlx(...)
```

Arguments

`...` parameters passed to [ggplot](#)

Value

see [ggplot](#)

<code>initMlxR</code>	<i>Initialize mlxR library</i>
-----------------------	--------------------------------

Description

Initialize mlxR library

Usage

```
initMlxR(path = NULL, ...)
```

Arguments

- path *(character)* [optional] Path to installation directory of the Lixoft suite. If mlxR library is not already loaded and no path is given, the directory written in the lixoft.ini file is used for initialization.
- ... [optional] Extra arguments passed to lixoftConnectors package when mlxR is used with a version of Lixoft(/@) software suite higher or equal to 2019R1.
- force *(bool)* [optional] Should mlxR initialization overpass lixoftConnectors software switch security or not. Equals FALSE by default.

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:
initMlxR(path = "/path/to/lixoftRuntime/")

## End(Not run)
```

inlineDataFrame	<i>inline data frame</i>
-----------------	--------------------------

Description

utility function to inline creation of a data frame

Usage

```
inlineDataFrame(str, header = TRUE, colClasses = NA, ...)
```

Arguments

- str text representation of the data frame
- header see [read.table](#)
- colClasses see [read.table](#)
- ... see [read.table](#)

inlineModel	<i>inline model</i>
-------------	---------------------

Description

Define a model "inline"

Usage

```
inlineModel(str, filename = NULL)
```

Arguments

str	model
filename	name of the temporary model file

Details

A temporary model file filename is created. Default name is "tempModel.txt". filename="random" generates a random name.

Value

A Shiny app with files ui.R, server.R and model.txt

Examples

```
## Not run:
myModel1 <- inlineModel("
[LONGITUDINAL]
EQUATION:
f = 10*exp(-0.2*t)
")

print(myModel1)

r <- simlux(model=myModel1, output=list(name="f", time=0:100))

myModel2 <- inlineModel("
[LONGITUDINAL]
EQUATION:
f = 10*exp(-0.2*t)
", filename="random")

print(myModel2)

## End(Not run)
```

kmplotmlx*Kaplan Meier plot*

Description

Plot empirical survival functions using the Kaplan Meier estimate.

Usage

```
kmplotmlx(  
  r,  
  index = 1,  
  level = NULL,  
  time = NULL,  
  cens = TRUE,  
  plot = TRUE,  
  color = "#e05969",  
  group = NULL,  
  facet = TRUE,  
  labels = NULL  
)
```

Arguments

<code>r</code>	a data frame with a column 'id', a column 'time', a column with values and possibly a column 'group'.
<code>index</code>	an integer: <code>index=k</code> means that the survival function for the k-th event is displayed. Default is <code>index=1</code> .
<code>level</code>	a number between 0 and 1: confidence interval level.
<code>time</code>	a vector of time points where the survival function is evaluated.
<code>cens</code>	if TRUE right censoring times are displayed.
<code>plot</code>	if TRUE the estimated survival function is displayed, if FALSE the values are returned
<code>color</code>	color to be used for the plots (default="#e05969")
<code>group</code>	variable to be used for defining groups (by default, 'group' is used when it exists)
<code>facet</code>	makes subplots for different groups if TRUE
<code>labels</code>	vector of strings

Details

See <http://simulx.webpopix.org/mlxr/kmplotmlx/> for more details.

Value

a ggplot object if plot=TRUE ; otherwise, a list with fields:

- surv a data frame with columns T (time), S (survival), possibly (S1, S2) (confidence interval) and possibly group
- cens a data frame with columns T0 (time), S0 (survival) and possibly group

Examples

```
## Not run:
tteModel1 <- inlineModel("
  [LONGITUDINAL]
  input = {beta,lambda}
  EQUATION:
  h=(beta/lambda)*(t/lambda)^(beta-1)
  DEFINITION:
  e = {type=event, maxEventNumber=1, rightCensoringTime=70, hazard=h}
")

p1 <- c(beta=2.5,lambda=50)
e <- list(name='e', time=0)
res1 <- simulx(model=tteModel1, parameter=p1, output=e, group=list(size=100))
p11 <- kmplotmlx(res1$e, level=0.95)
print(p11)

p2 <- c(beta=2,lambda=45)
g1 <- list(size=50, parameter=p1)
g2 <- list(size=100, parameter=p2)
res2 <- simulx(model=tteModel1, output=e, group=list(g1,g2))
p12 <- kmplotmlx(res2$e)
print(p12)

## End(Not run)
```

lixoft.read.table	<i>Read Lixoft@ files</i>
-------------------	---------------------------

Description

Utility function to read Lixoft@ formatted input/output files

Usage

```
lixoft.read.table(file, sep = "", ...)
```

Arguments

file	file path of the file to read
sep	separator
...	see read.table

`mlxplore`*Explore and visualize models*

Description

Explore and visualize ‘Mlxtran’ models with the ‘Mlxlore’ software.

Usage

```
mlxplore(  
  model,  
  parameter = NULL,  
  output = NULL,  
  group = NULL,  
  treatment = NULL  
)
```

Arguments

<code>model</code>	a Mlxtran model
<code>parameter</code>	a vector of parameters with their names and values
<code>output</code>	a list with fields: • <code>name</code>: a vector of output names • <code>time</code>: a vector of times
<code>group</code>	a list with unique field: • <code>treatment</code> : a list,
<code>treatment</code>	a list with fields • <code>time</code> : a vector of input times, • <code>amount</code> : a scalar or a vector of amounts, • <code>rate</code> : a scalar or a vector of infusion rates (default=Inf), • <code>type</code> : the type of input (default=1), • <code>target</code> : the target compartment (default=NULL).

Details

See <http://simulx.webpopix.org/mlxr/mlxplore/> for more details.

monolix2simulx

*Convert a Monolix Project into an executable for the simulator Simulx***Description**

Convert a Monolix Project into an executable for the simulator Simulx

Usage

```
monolix2simulx(
  project,
  parameter = NULL,
  group = NULL,
  open = FALSE,
  r.data = TRUE,
  fim = NULL
)
```

Arguments

project	: the name of a Monolix project
parameter	: string \$(NameOfTypeOfParameter), the type of specific parameters to use example: "mode", "mean"...
group	: a list with the number of subjects
open	: load the R script created if open=TRUE
r.data	: read the data if r.data=TRUE
fim	: Fisher information matrix

Value

creates a folder projectNameR containing files :

- projectName.R : executable R code for the simulator,
- treatment.txt : contains the treatment informations,
- populationParameter.txt : contains the population parameters estimated from Monolix,
- individualParameter.txt : contains the individual parameters (mode/mean) estimated from Monolix (if used for the simulation),
- individualCovariate.txt : contains the individual covariates,
- originalId.txt : contains the original id's when group is used with a different size than the original one,
- outputi.txt : contains the output number i informations (time, id),
- \$(NameOfTypeOfParameter)s.txt : contains the specific parameter used.

Examples

```
## Not run:
project.file <- 'monolixRuns/theophylline1_project.mlxtran' #relative path
monolix2simulx(project=project.file,open=TRUE)
monolix2simulx(project=project.file,parameter=list("mean",c(a=0, b=0)),open=TRUE)

## End(Not run)
```

pkmodel	<i>Easy simulation of PK models Easy simulation of basic PK models See http://simulx.webpopix.org/mlxr/pkmodel/ for more details.</i>
---------	--

Description

Easy simulation of PK models

Easy simulation of basic PK models

See <http://simulx.webpopix.org/mlxr/pkmodel/> for more details.

Usage

```
pkmodel(time, treatment, parameter)
```

Arguments

time	a vector
treatment	a list with fields - time : a vector of input times, - amount : a scalar or a vector of amounts, - rate : a scalar or a vector of infusion rates (default=Inf), - tinf : a scalar or a vector of infusion times (default=0),
parameter	vector of parameters with their names and values

Examples

```
## Not run:
adm <- list(time=c(2,14,20), amount=40)
p <- c(V=8, Cl=0.5,k12=0.3, k21=0.2)
t <- seq(0, 30, by=0.1)

res <- pkmodel(time = t, treatment = adm, parameter = p)

print(ggplot(data=res, aes(x=time, y=cc)) + geom_line(size=1) +
  xlab("time (h)") + ylab("concentration (mg/L)"))

adm <- list(time = c(1,23,37,45), amount = c(1,0.5,2,0.3))
p <- c(Mtt=5, Ktr=1, ka=0.5, V=10, Vm=1, Km=0.6, p=0.5)
```

```

t <- seq(0, 80, by=0.1)

res <- pkmodel(t,adm,p)

print(ggplot(data=res, aes(x=time, y=cc)) + geom_line(size=1) +
      xlab("time (h)") + ylab("concentration (mg/L)"))

adm <- list( time = 2, amount = 40)

p <- inlineDataFrame("
id  ka  V  Cl
1   0.5  4   1
2    1  6   1
3   1.5  6  1.5
")

t <- seq(0, 30, by=0.1)

res <- pkmodel(t,adm,p)

print(ggplot(data=res, aes(x=time, y=cc, colour=id)) + geom_line(size=1) +
      xlab("time (h)") + ylab("concentration (mg/L)"))
adm <- list(time=seq(2, 100, by=24), amount=40, rate=5)
p <- c(V=8, Cl=0.5, k12=0.3, k21=0.2, ke0=0.2)
t <- seq(0, 50, by=0.1)

res <- pkmodel(t,adm,p)

if( require("reshape2") ){
  r <- melt(res, id='time', variable.name='c')
  print(ggplot(r, aes(time,value)) + geom_line(aes(colour = c),size=1) +
        ylab('concentration') + guides(colour=guide_legend(title=NULL)) +
        theme(legend.position=c(.9, .8)))
}

## End(Not run)

```

prctilemlx

Percentiles of the empiricial distribution of longitudinal data

Description

Compute and display percentiles of the empiricial distribution of longitudinal data.

Usage

```

prctilemlx(
  r,
  col = NULL,
  number = 8,

```

```

    level = 80,
    plot = TRUE,
    color = "#9a35ff",
    group = NULL,
    facet = TRUE,
    labels = NULL,
    band = NULL
  )

```

Arguments

<code>r</code>	a data frame with a column 'id', a column 'time' and a column with values. The times should be the same for each individual.
<code>col</code>	a vector with the three column indexes for 'id', 'time/x' and 'y'. Default = c(1, 2, 3).
<code>number</code>	the number of intervals (i.e. the number of percentiles minus 1).
<code>level</code>	the largest interval (i.e. the difference between the lowest and the highest percentile).
<code>plot</code>	if TRUE the empirical distribution is displayed, if FALSE the values are returned
<code>color</code>	a color to be used for the plots (default="#9a35ff")
<code>group</code>	variable to be used for defining groups (by default, 'group' is used when it exists)
<code>facet</code>	makes subplots for different groups if TRUE
<code>labels</code>	vector of strings
<code>band</code>	is deprecated (use number and level instead) ; a list with two fields • <code>number</code> the number of intervals (i.e. the number of percentiles minus 1). • <code>level</code> the largest interval (i.e. the difference between the lowest and the highest percentile).

Details

See <http://simulx.webpopix.org/mlxr/prtilemlx/> for more details.

Value

a ggplot object if `plot=TRUE` ; otherwise, a list with fields:

- `proba` a vector of probabilities of length `band$number+1`
- `color` a vector of colors used for the plot of length `band$number`
- `y` a data frame with the values of the empirical percentiles computed at each time point

Examples

```
## Not run:
myModel <- inlineModel("
[LONGITUDINAL]
input = {ka, V, Cl}
EQUATION:
C = pkmodel(ka,V,Cl)

[INDIVIDUAL]
input = {ka_pop, V_pop, Cl_pop, omega_ka, omega_V, omega_Cl}
DEFINITION:
ka = {distribution=lognormal, reference=ka_pop, sd=omega_ka}
V = {distribution=lognormal, reference=V_pop, sd=omega_V }
Cl = {distribution=lognormal, reference=Cl_pop, sd=omega_Cl}
")

N=2000

pop.param <- c(
  ka_pop = 1,    omega_ka = 0.5,
  V_pop  = 10,   omega_V  = 0.4,
  Cl_pop = 1,    omega_Cl = 0.3)

res <- simulx(model      = myModel,
               parameter = pop.param,
               treatment = list(time=0, amount=100),
               group     = list(size=N, level='individual'),
               output     = list(name='C', time=seq(0,24,by=0.1)))
# res$C is a data.frame with 2000x241=482000 rows and 3 columns
head(res$C)
# we can display the empirical percentiles of C using the default
# settings (i.e. percentiles of order 10%, 20%, ... 90%)
prctilemlx(res$C)
# The 3 quartiles (i.e. percentiles of order 25%, 50% and 75%) are displayed by
# selecting a 50% interval splitted into 2 subintervals
prctilemlx(res$C, number=2, level=50)
# A one 90% interval can be displayed using only one interval
prctilemlx(res$C, number=1, level=90)
# or 75 subintervals in order to better represent the continuous distribution
# of the data within this interval
prctilemlx(res$C, number=75, level=90)
# prctilemlx produces a ggplot object that can be modified
pl <- prctilemlx(res$C, number=4, level=80)
pl + ylab("concentration") + ggtitle("predictive distribution")
# The percentiles are not plotted by setting plot=FALSE
pr.out <- prctilemlx(res$C, number=4, level=80, plot=FALSE)
print(pr.out$proba)
print(pr.out$color)
print(pr.out$y[1:5,])

## End(Not run)
```

read.vector	<i>Reads a table into a vector</i>
-------------	------------------------------------

Description

Reads a table into a vector

Usage

```
read.vector(f, header = FALSE, sep = "", quote = "\"'")
```

Arguments

f	: the table
header	: bool, use the header or not
sep	: the separator
quote	: the quote character

Value

the vector

readDatamlx	<i>Read formatted data file</i>
-------------	---------------------------------

Description

Read data in a Monolix/NONMEM format

Usage

```
readDatamlx(  
  project = NULL,  
  data = NULL,  
  out.data = FALSE,  
  nbSSDoses = 10,  
  obs.rows = FALSE,  
  error.iov = FALSE,  
  filter = NULL,  
  datafile = NULL,  
  header = NULL,  
  infoProject = NULL,  
  addl.ss = NULL  
)
```

Arguments

project	a Monolix project
data	a list with fields • dataFile: path of a formatted data file • headerTypes: a vector of strings
out.data	TRUE/FALSE (default=FALSE) returns the original data as a table and some information about the Monolix project
nbSSDoses	number of additional doses to use for steady-state (default=10)
obs.rows	a list of observation indexes
error.iov	TRUE/FALSE (default=TRUE) returns an error message if occasions are overlapping
filter	filter to apply to the data (string)
datafile	(deprecated) a formatted data file
header	(deprecated) a vector of strings
infoProject	(deprecated) an xmlfile
addl.ss	(deprecated) number of additional doses to use for steady-state (default=10)

Details

See <http://simulx.webpopix.org/mlxr/readdatamlx/> for more details.

Value

A list of data frames

Examples

```
## Not run:
# using a Monolix project:
d <- readDatamlx(project='projects/warfarinPK.mlxtran')

# using a data file:
warfarinPK <- list(dataFile = "data/warfarinPK.csv",
                  headerTypes = c("id", "time", "observation", "amount",
                                   "contcov", "contcov", "catcov"),
                  administration = "oral")
d <- readDatamlx(data=warfarinPK)

## End(Not run)
```

shinymlx

*Automatic code generation for Shiny applications***Description**

Creates a Shiny application for longitudinal data model

Usage

```
shinymlx(
  model,
  parameter = NULL,
  output = NULL,
  treatment = NULL,
  regressor = NULL,
  group = NULL,
  data = NULL,
  appname = "shinymlxApp",
  style = "basic",
  settings = NULL,
  title = " "
)
```

Arguments

model	a Mlxtran model used for the simulation
parameter	a vector, or a list of shiny widgets
output	a list - or a list of lists - with fields: • name: a vector of output names • time: a vector of times, or a vector (min, max, step)
treatment	a list with fields • tfd : first time of dose, • amount : amount, • nd : number of doses, • ii : interdose interval, • type : the type of input, Input argument of Simulx can also be used, i.e. a list with fields time, amount, rate, tinf, type, target.
regressor	a list, or a list of lists, with fields • name : a vector of regressor names, • time : a vector of times, • value : a vector of values.
group	a list, or a list of lists, with fields:

	• <code>size</code> : size of the group (default=1), • <code>level</code> : level(s) of randomization, • <code>parameter</code> : if different parameters per group are defined, • <code>output</code> : if different outputs per group are defined, • <code>treatment</code> : if different treatments per group are defined, • <code>regressor</code> : if different regression variables per group are defined.
<code>data</code>	data to display with the plot (either a data frame or the name of a file)
<code>appname</code>	the name of the application (and possibly its path)
<code>style</code>	the style of the Shiny app • <code>"basic"</code>: basic Shiny app with a single side bar (default) • <code>"navbar1"</code>: navigation bar and tabPanels (including outputs) • <code>"navbar2"</code>: navigation bar and tabPanels (outputs separated) • <code>"dashboard1"</code> : Shiny dashboard,
<code>settings</code>	a list of settings • <code>"tabstyle"</code> : look of the tabs c("tabs","pills"), • <code>"select.x"</code> : display the list of variables available for the x-axis c(TRUE,FALSE), • <code>"select.y"</code> : display the list of variables available for the y-axis c(TRUE,FALSE), • <code>"select.log"</code> : log scale option c(TRUE,FALSE), • <code>"select.ref"</code> : reference curves option c(TRUE,FALSE)
<code>title</code>	the title of the application

Details

shinymlx automatically generates files `ui.R` and `server.R` required for a Shiny application.

Elements of parameters and treatment can be either scalars or lists. A scalar automatically generates a slider with default minimum and maximum values and default step. A list may contain the type of widget (`"slider"`, `"select"`, `"numeric"`) and the settings defining the widget: (value, min, max, step) for slider, (selected, choices) for select and value for numeric.

See <http://simulx.webpopix.org/mlxr/shinymlx/> for more details.

Value

A Shiny app with files `ui.R`, `server.R` and `model.txt`

Examples

```
## Not run:
library(mlxR)
PKPDmodel <- inlineModel("
[LONGITUDINAL]
input={ka,V,C1,Imax,IC50,S0,kout}
EQUATION:
C      = pkmodel(ka, V, C1)
E_0    = S0
ddt_E  = kout*((1-Imax*C/(C+IC50))*S0- E)
```

```

")

p1 <- c(ka=0.5, V=10, Cl=1)
p2 <- c(Imax=0.5, IC50=0.03, S0=100, kout=0.1)
adm <- list(tfd=5, nd=15, ii=12, amount=1)
f1 <- list(name = 'C', time = seq(0, 250, by=1))
f2 <- list(name = 'E', time = seq(0, 250, by=1))
f <- list(f1, f2)

shinymlx(model=PKPDmodel, treatment=adm, parameter=list(p1,p2), output=f,
          style="dashboard1")

#-----
p1 <- list(
  ka = list(widget="slider", value=0.5, min=0.1, max=2, step=0.1),
  V = list(widget="slider", value=10, min=2, max=20, step=2),
  Cl = list(widget="slider", value=1, min=0.1, max=2, step=0.1)
)
adm <- list(
  tfd = list(widget="slider", value=5, min=0, max=100, step=5),
  nd = list(widget="numeric", value=15),
  ii = list(widget="select", selected=12, choices=c(3,6,12,18,24)),
  amount = list(widget="slider", value=40, min=0, max=50, step=5)
)
s <- list(select.x=FALSE, select.y=FALSE)
shinymlx(model=PKPDmodel, treatment=adm, parameter=list(p1,p2), output=f,
          style="navbar1", settings=s)

## End(Not run)

```

simpopmlx

Population parameters simulation

Description

Draw population parameters using the covariance matrix of the estimates.

Usage

```

simpopmlx(
  n = 1,
  project = NULL,
  fim = "needed",
  parameter = NULL,
  corr = NULL,
  kw.max = 100
)

```

Arguments

n	the number of vectors of population parameters (default = 1),
project	a Monolix project, assuming that the Fisher information Matrix was estimated by Monolix.
fim	the Fisher Information Matrix estimated by Monolix. fim="sa", "lin" (default="sa")
parameter	a data frame with a column 'pop.param' (no default), a column 'sd' (no default), possibly a column 'trans' (default='N') and possibly columns 'lim.a' (default=0) and 'lim.b' (default=1). Only when project is not used.
corr	correlation matrix of the population parameters (default = identity). Only when project is not used.
kw.max	maximum number of trials for generating a positive definite covariance matrix (default = 100)

Details

See <http://simulx.webpopix.org/mlxr/simpopmlx/> for more details.

Examples

```
## Not run:
project.file <- 'monolixRuns/theophylline1_project.mlxtran' #relative path
pop1 <- simpopmlx(n=3, project=project.file)

## End(Not run)
```

simulx

Simulation of mixed effects models and longitudinal data

Description

Compute predictions and sample data from Mlxtran and R models

Usage

```
simulx(
  model = NULL,
  parameter = NULL,
  output = NULL,
  treatment = NULL,
  regressor = NULL,
  varlevel = NULL,
  group = NULL,
  data = NULL,
  project = NULL,
  nrep = 1,
  npop = NULL,
```

```

    fim = NULL,
    result.folder = NULL,
    result.file = NULL,
    stat.f = "statmlx",
    addlines = NULL,
    settings = NULL
)

```

Arguments

model	a Mlxtran, or R model used for the simulation
parameter	a vector of parameters with their names and values
output	a list (or list of lists) with fields: • name: a vector of output names • time: a vector of times (only for the longitudinal outputs) • lloq: lower limit of quantification (only for the longitudinal outputs) • uloq: upper limit of quantification (only for the longitudinal outputs) • limit: lower bound of the censoring interval (only for the longitudinal outputs)
treatment	a list with fields • time : a vector of input times, • amount : a scalar or a vector of amounts, • rate : a scalar or a vector of infusion rates (default=Inf), • tinf : a scalar or a vector of infusion times (default=0), • type : the type of input (default=1), • target : the target compartment (default=NULL).
regressor	a list, or a list of lists, with fields • name : a vector of regressor names, • time : a vector of times, • value : a vector of values.
varlevel	(IOV supported by mlxR >= 3.2.2) a list (or a dataframe) with fields: • name : name of the variable which defines the occasions, • time : a vector of times (beginnings of occasions) ,
group	a list, or a list of lists, with fields: • size : size of the group (default=1), • level : level(s) of randomization, • parameter : if different parameters per group are defined, • output : if different outputs per group are defined, • treatment : if different treatments per group are defined, • regressor : if different regression variables per group are defined.
data	a list (output of simulx when settings\$data.in==TRUE)
project	the name of a Monolix project

nrep	number of replicates
npop	number of population parameters to draw randomly
fim	a string with the Fisher Information Matrix to be used
result.folder	the name of the folder where the outputs of simulx should be stored
result.file	the name of the single file where the outputs of simulx should be saved
stat.f	a R function for computing some summary (mean, quantiles, survival,...) of the simulated data. Default = "statmlx".
addlines	a list with fields: • section: a string (default = "[LONGITUDINAL]"), • block: a string (default = "EQUATION:"), • formula: string, or vector of strings, to be inserted .
settings	a list of optional settings • seed : initialization of the random number generator (integer), • load.design : TRUE/FALSE (if load.design is not defined, a test is automatically performed to check if a new design has been defined), • data.in : TRUE/FALSE (default=FALSE) • id.out : add columns id (when N=1) and group (when #group=1), TRUE/FALSE (default=FALSE) • kw.max : maximum number of trials for generating a positive definite covariance matrix (default = 100) • sep : the field separator character (default = ",") • digits : number of decimal digits in output files (default = 5) • disp.iter : TRUE/FALSE (default = FALSE) display replicate and population numbers • replacement : TRUE/FALSE (default = FALSE) sample id's with/without replacement • out.trt : TRUE/FALSE (default = TRUE) output of simulx includes treatment • format.original : TRUE/FALSE (default = FALSE) with a Monolix project, write data in result.file using the original format of the data file

Details

simulx takes advantage of the modularity of hierarchical models for simulating different components of a model: models for population parameters, individual covariates, individual parameters and longitudinal data.

Furthermore, simulx allows to draw different types of longitudinal data, including continuous, count, categorical, and time-to-event data.

The models are encoded using either the model coding language 'Mlxtran', or 'R'. 'Mlxtran' models are automatically converted into C++ codes, compiled on the fly and linked to R using the 'Rcpp' package. That allows one to implement very easily complex models and to take advantage of the numerical solvers used by the C++ 'mlxLibrary'.

See <http://simulx.webpopix.org> for more details.

Value

A list of data frames. Each data frame is an output of simulx

Examples

```
## Not run:
myModel <- inlineModel("
[LONGITUDINAL]
input = {A, k, c, a}
EQUATION:
t0      = 0
f_0     = A
ddt_f   = -k*f/(c+f)
DEFINITION:
y = {distribution=normal, prediction=f, sd=a}
[INDIVIDUAL]
input = {k_pop, omega}
DEFINITION:
k = {distribution=lognormal, prediction=k_pop, sd=omega}
")
f <- list(name='f', time=seq(0, 30, by=0.1))
y <- list(name='y', time=seq(0, 30, by=2))
res <- simulx(model      = 'model/home.txt',
              parameter = c(A=100, k_pop=6, omega=0.3, c=10, a=2),
              output    = list(f,y,"k"),
              group      = list(size=4, level='individual'))

plot(ggplotmlx() + geom_line(data=res$f, aes(x=time, y=f, colour=id)) +
     geom_point(data=res$y, aes(x=time, y=y, colour=id)))
print(res$parameter)

## End(Not run)
```

statmlx

*Summary of data***Description**

Compute statistical summaries (mean, quantile, variance, survival rate,...)

Usage

```
statmlx(r, FUN = "mean", probs = c(0.05, 0.5, 0.95), surv.time = NULL)
```

Arguments

r a data frame

FUN	a string, or a vector of strings, with the name of the functions to apply to the result of the simulation
probs	a vector of quantiles between 0 and 1. Only used when "quantile" has been defined in FUN
surv.time	a scalar or a vector of times. Only used when "event" has been defined in type

Details

See <http://simulx.webpopix.org/statmlx> for more details.

Value

A data frame.

Examples

```
## Not run:
modelPK <- inlineModel("
[LONGITUDINAL]
input={V,C1,alpha, beta,b}

EQUATION:
C = pkmodel(V, C1)
h = alpha*exp(beta*C)
g = b*C

DEFINITION:
y = {distribution=normal, prediction=C, sd=g}
e = {type=event, maxEventNumber=1, rightCensoringTime=30, hazard=h}

[INDIVIDUAL]
input={V_pop,C1_pop,omega_V,omega_C1}

DEFINITION:
V      = {distribution=lognormal,    prediction=V_pop,    sd=omega_V}
C1     = {distribution=lognormal,    prediction=C1_pop,   sd=omega_C1}
")

adm <- list(amount=100, time=0)
p <- c(V_pop=10, C1_pop=1, omega_V=0.2, omega_C1=0.2, alpha=0.02, beta=0.1, b=0.1)
out.y <- list(name=c('y'), time=seq(0,to=25,by=5))
out.e <- list(name=c('e'), time=0)
out.p <- c("V", "C1")
out <- list(out.y, out.e, out.p)
g <- list(size=100, level='individual')
res1 <- simulx(model=modelPK, treatment=adm, parameter=p, output=out, group=g)

statmlx(res1$parameter, FUN = "mean", probs = c(0.05, 0.5, 0.95))
statmlx(res1$parameter, FUN = "quantile", probs = c(0.05, 0.5, 0.95))
statmlx(res1$parameter, FUN = c("sd", "quantile"), probs = c(0.05, 0.95))
statmlx(res1$y, FUN = c("mean", "sd", "quantile"), probs = c(0.05, 0.95))
statmlx(res1$e, surv.time=c(10,20))
```

```

res2 <- simulx(model=modelPK, treatment=adm, parameter=p, output=out, group=g, nrep=3)
statmlx(res2$parameter, FUN = c("sd", "quantile"), probs = c(0.05, 0.95))
statmlx(res2$y, FUN = c("mean", "sd", "quantile"), probs = c(0.05, 0.95))
statmlx(res2$e, surv.time=c(10,20,30))

## End(Not run)

```

writeDatamlx	<i>Write formatted data file</i>
--------------	----------------------------------

Description

Write data contained in a list of dataframes in a single file (NONMEM/Monolix format) or in several files as tables

Usage

```

writeDatamlx(
  r,
  result.file = NULL,
  result.folder = NULL,
  sep = ",",
  ext = NULL,
  digits = 5,
  app.file = F,
  app.dir = F,
  project = NULL
)

```

Arguments

<code>r</code>	a list of dataframes
<code>result.file</code>	a string with the name of the file
<code>result.folder</code>	a string with the name of the folder
<code>sep</code>	(default = ",")
<code>ext</code>	a string with the extension of the file names
<code>digits</code>	(default = 5)
<code>app.file</code>	TRUE/FALSE (default=FALSE) append to file
<code>app.dir</code>	TRUE/FALSE (default=FALSE) append to dir
<code>project</code>	A Monolix project

Details

See <http://simulx.webpopix.org/mlxr/writedatamlx/> for more details.

Examples

```
## Not run:
modelPK <- inlineModel("
[LONGITUDINAL]
input = {V, Cl, a1}
EQUATION:
Cc = pkmodel(V, Cl)
DEFINITION:
y1 ={distribution=lognormal, prediction=Cc, sd=a1}
")
adm <- list(amount=100, time=seq(0,50,by=12))
p <- c(V=10, Cl=1, a1=0.1)
y1 <- list(name=c('y1'), time=seq(5,to=50,by=5))
res <- simulx(model=modelPK, treatment=adm, parameter=p, output=y1)
writeDatamlx(res, result.file="res.csv")
writeDatamlx(res, result.file="res.txt", sep="\t")
writeDatamlx(res, result.folder="res")

## End(Not run)
```

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