Package 'babelmixr2'

September 23, 2024

```
Type Package
Title Use 'nlmixr2' to Interact with Open Source and Commercial
      Software
Version 0.1.4
Description Run other estimation and simulation software via the 'nlmixr2' (Fidler et al (2019)
      <doi:10.1002/psp4.12445>) interface including 'PKNCA', 'NONMEM' and 'Mono-
      lix'. While not required, you can
      get/install the 'lixoftConnectors' package in the 'Monolix' installation, as
      described at the following url
      <https://monolixsuite.slp-software.com/r-functions/2024R1/</pre>
      installation-and-initialization>. When
      'lixoftConnectors' is available, 'Monolix' can be run directly instead of setting up
      command line usage.
License GPL (>= 3)
URL https://nlmixr2.github.io/babelmixr2/,
      https://github.com/nlmixr2/babelmixr2/
NeedsCompilation yes
Encoding UTF-8
Suggests testthat, nlmixr2data, withr, lixoftConnectors, PKNCA (>=
      0.10.0), knitr, rmarkdown, spelling, PopED, units, vdiffr,
Depends R (>= 3.5), nlmixr2 (>= 2.0.8)
Imports checkmate, cli, digest, lotri, nlmixr2est (>= 2.1.6),
      nonmem2rx (>= 0.1.3), monolix2rx, methods, qs, rex, rxode2 (>=
      3.0.0)
RoxygenNote 7.3.2
Config/testthat/edition 3
LinkingTo Rcpp, rxode2, RcppArmadillo, RcppEigen
Language en-US
VignetteBuilder knitr
```

2 .setupPopEDdatabase

Author Matthew Fidler [aut, cre] (https://orcid.org/0000-0001-8538-6691), Bill Denney [aut] (https://orcid.org/0000-0002-5759-428X), Nook Fulloption [ctb] (goldfish art)

Maintainer Matthew Fidler <matthew.fidler@gmail.com>

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Contents

pPopEDomixr2 . l.poped.o atToMor tandardC elUnitCo	 latabase iolix . olName	· · · · · · · · · · · · · · · · · · ·	 	· ·																						3
atToMor tandardC elUnitCo	olix . olName																									
tandardC elUnitCo	olName										•	•		٠		•			•					•		5
elUnitCo		2.5																								5
																										8
	nversio																									9
olixContı	ol																									10
xr2Est.pl	enca .																									13
nemCont	rol																									14
aControl																										17
dGetMul	tipleEn	dpoir	ιtΜ	ode	ling	gTir	nes																			28
					_																					
Monolix																										30
Nonmem																										31
lifyUnit																										31
																										33
i	ixr2Est.pk memCont caControl edControl edGetMul edMultipl bMonolix bNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpo bMonolix bNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix	ixr2Est.pknca	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	ixr2Est.pknca memControl caControl edControl edGetMultipleEndpointModelingTimes edMultipleEndpointResetTimeIndex oMonolix oNonmem	isolixControl ixr2Est.pknca memControl caControl caControl cdControl cdGetMultipleEndpointModelingTimes cdMultipleEndpointResetTimeIndex coMonolix coNonmem						

Setup the poped database

Description

Setup the poped database

.setupPopEDdatabase

Usage

.setupPopEDdatabase(ui, data, control)

Arguments

ui rxode2 ui function
data babelmixr2 design data

control PopED control

as.nlmixr2

Value

PopED database

Author(s)

Matthew L. Fidler

as.nlmixr2

Convert an object to a nlmixr2 fit object

Description

Convert an object to a nlmixr2 fit object

Usage

```
as.nlmixr2(
    x,
    ...,
    table = nlmixr2est::tableControl(),
    rxControl = rxode2::rxControl(),
    ci = 0.95
)

as.nlmixr(
    x,
    ...,
    table = nlmixr2est::tableControl(),
    rxControl = rxode2::rxControl(),
    ci = 0.95
)
```

Arguments

Х	Object to convert
	Other arguments
table	is the nlmixr2est::tableControl() options
rxControl	is the rxode2::rxControl() options, which is generally needed for how addl doses are handled in the translation
ci	is the confidence interval of the residual differences calculated (by default 0.95)

Value

nlmixr2 fit object

4 as.nlmixr2

Author(s)

Matthew L. Fidler

Examples

```
# First read in the model (but without residuals)
mod <- nonmem2rx(system.file("mods/cpt/run0DE032.ctl", package="nonmem2rx"),</pre>
                  determineError=FALSE, 1st=".res", save=FALSE)
# define the model with residuals (and change the name of the
# parameters) In this step you need to be careful to not change the
# estimates and make sure the residual estimates are correct (could
# have to change var to sd).
 mod2 <-function() {</pre>
   ini({
     lcl <- 1.37034036528946
     lvc <- 4.19814911033061
     lq <- 1.38003493562413
     lvp <- 3.87657341967489
     RSV <- c(0, 0.196446108190896, 1)
     eta.cl ~ 0.101251418415006
     eta.v ~ 0.0993872449483344
     eta.q ~ 0.101302674763154
     eta.v2 ~ 0.0730497519364148
   })
   model({
     cmt(CENTRAL)
     cmt(PERI)
     cl <- exp(lcl + eta.cl)</pre>
     v <- exp(lvc + eta.v)</pre>
     q \leftarrow exp(lq + eta.q)
     v2 \leftarrow exp(lvp + eta.v2)
     v1 <- v
     scale1 <- v
     k21 < - q/v2
     k12 \leftarrow q/v
     d/dt(CENTRAL) <- k21 * PERI - k12 * CENTRAL - cl * CENTRAL/v1
     d/dt(PERI) <- -k21 * PERI + k12 * CENTRAL
     f <- CENTRAL/scale1
     f ~ prop(RSV)
   })
 }
# now we create another nonmem2rx object that validates the model above:
new <- as.nonmem2rx(mod2, mod)</pre>
# once that is done, you can translate to a full nlmixr2 fit (if you wish)
```

babel.poped.database 5

```
fit <- as.nlmixr2(new)
print(fit)</pre>
```

babel.poped.database Expand a babelmixr2 PopED database

Description

Expand a babelmixr2 PopED database

Usage

```
babel.poped.database(popedInput, ..., optTime = NA)
```

Arguments

popedInput The babelmixr2 generated PopED database

... other parameters sent to PopED::create.poped.database()

optTime boolean to indicate if the global time indexer inside of babelmixr2 is reset if

the times are different. By default this is TRUE. If FALSE you can get slightly better run times and possibly slightly different results. When optTime is FALSE the global indexer is reset every time the PopED rxode2 is setup for a problem or

when a poped dataset is created. You can manually reset with popedMultipleEndpointResetTimeIndex

Value

babelmixr2 PopED database (with \$babelmixr2 in database)

Author(s)

Matthew L. Fidler

bblDatToMonolix Convert nlmixr2-compatible data to other formats (if possible)

Description

Convert nlmixr2-compatible data to other formats (if possible)

6 bblDatToMonolix

Usage

```
bblDatToMonolix(
 model,
  data,
  table = nlmixr2est::tableControl(),
  rxControl = rxode2::rxControl(),
  env = NULL
)
bblDatToNonmem(
 model,
  data,
  table = nlmixr2est::tableControl(),
  rxControl = rxode2::rxControl(),
  env = NULL
)
bblDatToRxode(
 model,
  data,
  table = nlmixr2est::tableControl(),
  rxControl = rxode2::rxControl(),
  env = NULL
)
bblDatToMrgsolve(
 model,
  data,
  table = nlmixr2est::tableControl(),
  rxControl = rxode2::rxControl(),
  env = NULL
)
bblDatToPknca(
 model,
  data,
  table = nlmixr2est::tableControl(),
  rxControl = rxode2::rxControl(),
  env = NULL
)
```

Arguments

model rxode2 model for conversion

data Input dataset.

table is the table control; this is mostly to figure out if there are additional columns to

keep.

bblDatToMonolix 7

rxControl is the rxode2 control options; This is to figure out how to handle the addl dosing

information.

env When NULL (default) nothing is done. When an environment, the function nlmixr2est::.foceiPreProce

env, model, rxControl) is called on the provided environment.

Value

With the function bblDatToMonolix() return a list with:

- Monolix compatible dataset (\$monolix)
- Monolix ADM information (\$adm)

With the function nlmixrDataToNonmem() return a dataset that is compatible with NONMEM.

With the function nlmixrDataToMrgsolve() return a dataset that is compatible with mrgsolve. Unlike NONMEM, it supports replacement events with evid=8 (note with rxode2 replacement evid is 5).

With the function nlmixrDataToRxode() this will normalize the dataset to use newer evid definitions that are closer to NONMEM instead of any classic definitions that are used at a lower level

Author(s)

Matthew L. Fidler

Examples

```
pk.turnover.emax3 <- function() {</pre>
  ini({
    tktr < -log(1)
    tka <- log(1)
    tcl <- log(0.1)
    tv <- log(10)
    ##
    eta.ktr ~ 1
    eta.ka ~ 1
    eta.cl ~ 2
    eta.v ~ 1
    prop.err <- 0.1
    pkadd.err <- 0.1
    ##
    temax <- logit(0.8)</pre>
    tec50 < -log(0.5)
    tkout <- log(0.05)
    te0 < -log(100)
    eta.emax \sim .5
    eta.ec50 ~ .5
    eta.kout ~ .5
    eta.e0 \sim .5
    pdadd.err <- 10
  })
```

```
model({
   ktr <- exp(tktr + eta.ktr)</pre>
   ka <- exp(tka + eta.ka)
   cl <- exp(tcl + eta.cl)</pre>
    v <- exp(tv + eta.v)</pre>
    emax = expit(temax+eta.emax)
    ec50 = exp(tec50 + eta.ec50)
    kout = exp(tkout + eta.kout)
    e0 = exp(te0 + eta.e0)
    ##
   DCP = center/v
   PD=1-emax*DCP/(ec50+DCP)
    effect(0) = e0
    kin = e0*kout
    ##
    d/dt(depot) = -ktr * depot
    d/dt(gut) = ktr * depot -ka * gut
    d/dt(center) = ka * gut - cl / v * center
    d/dt(effect) = kin*PD -kout*effect
    ##
    cp = center / v
   cp ~ prop(prop.err) + add(pkadd.err)
    effect ~ add(pdadd.err) | pca
  })
}
bblDatToMonolix(pk.turnover.emax3, nlmixr2data::warfarin)
bblDatToNonmem(pk.turnover.emax3, nlmixr2data::warfarin)
bblDatToMrgsolve(pk.turnover.emax3, nlmixr2data::warfarin)
bblDatToRxode(pk.turnover.emax3, nlmixr2data::warfarin)
```

getStandardColNames

Determine standardized rxode2 column names from data

Description

Determine standardized rxode2 column names from data

Usage

```
getStandardColNames(data)
```

Arguments

data

A data.frame as the source for column names

modelUnitConversion 9

Value

A named character vector where the names are the standardized names and the values are either the name of the column from the data or NA if the column is not present in the data.

Examples

```
getStandardColNames(data.frame(ID=1, DV=2, Time=3, CmT=4))
```

modelUnitConversion

Unit conversion for pharmacokinetic models

Description

Unit conversion for pharmacokinetic models

Usage

```
modelUnitConversion(
  dvu = NA_character_,
  amtu = NA_character_,
  timeu = NA_character_,
  volumeu = NA_character_
)
```

Arguments

```
dvu, amtu, timeu The units for the DV, AMT, and TIME columns in the data volumeu The units for the volume parameters in the model
```

Value

A list with names for the units associated with each parameter ("amtu", "clearanceu", "volumeu", "timeu", "dvu") and the numeric value to multiply the modeled estimate (for example, cp) so that the model is consistent with the data units.

See Also

```
Other Unit conversion: simplifyUnit()
```

Examples

```
modelUnitConversion(dvu = "ng/mL", amtu = "mg", timeu = "hr", volumeu = "L")
```

10 monolixControl

monolixControl

Monolix Controller for nlmixr2

Description

Monolix Controller for nlmixr2

Usage

```
monolixControl(
  nbSSDoses = 7,
  useLinearization = FALSE,
  stiff = FALSE,
  addProp = c("combined2", "combined1"),
  exploratoryAutoStop = FALSE,
  smoothingAutoStop = FALSE,
  burnInIterations = 5,
  smoothingIterations = 200,
  exploratoryIterations = 250,
  simulatedAnnealingIterations = 250,
  exploratoryInterval = 200,
  exploratoryAlpha = 0,
  omegaTau = 0.95,
  errorModelTau = 0.95,
  variability = c("none", "firstStage", "decreasing"),
  runCommand = getOption("babelmixr2.monolix", ""),
  rxControl = NULL,
  sumProd = FALSE,
  optExpression = TRUE,
  calcTables = TRUE,
  compress = TRUE,
  ci = 0.95,
  sigdigTable = NULL,
  absolutePath = FALSE,
 modelName = NULL,
 muRefCovAlg = TRUE,
  run = TRUE,
)
```

Arguments

```
nbSSDoses Number of steady state doses (default 7)
useLinearization
Use linearization for log likelihood and fim.
stiff boolean for using the stiff ODE solver
```

monolixControl 11

addProp

specifies the type of additive plus proportional errors, the one where standard deviations add (combined1) or the type where the variances add (combined2). The combined1 error type can be described by the following equation:

$$y = f + (a + b \times f^c) \times \varepsilon$$

The combined2 error model can be described by the following equation:

$$y = f + \sqrt{a^2 + b^2 \times f^{2 \times c}} \times \varepsilon$$

Where:

- y represents the observed value

- f represents the predicted value

- a is the additive standard deviation

- b is the proportional/power standard deviation

- c is the power exponent (in the proportional case c=1)

exploratoryAutoStop

logical to turn on or off exploratory phase auto-stop of SAEM (default 250)

smoothingAutoStop

Boolean indicating if the smoothing should automatically stop (default FALSE)

burnInIterations

Number of burn in iterations

smoothingIterations

Number of smoothing iterations

exploratory Iterations

Number of iterations for exploratory phase (default 250)

simulated Annealing Iterations

Number of simulating annealing iterations

exploratoryInterval

Minimum number of iterations in the exploratory phase (default 200)

exploratoryAlpha

Convergence memory in the exploratory phase (only used when exploratoryAutoStop

is TRUE)

omegaTau Proportional rate on variance for simulated annealing

errorModelTau Proportional rate on error model for simulated annealing

variability This describes the methodology for parameters without variability. It could be:

- Fixed throughout (none) - Variability in the first stage (firstStage) - Decreasing

until it reaches the fixed value (decreasing)

runCommand is a shell command or function to run monolix; You can specify the default by

options("babelmixr2.monolix"="runMonolix"). If it is empty and 'lixoft-Connectors' is available, use lixoftConnectors to run monolix. See details for

function usage.

rxControl 'rxode2' ODE solving options during fitting, created with 'rxControl()'

sumProd Is a boolean indicating if the model should change multiplication to high pre-

cision multiplication and sums to high precision sums using the PreciseSums

package. By default this is FALSE.

12 monolixControl

optExpression	Optimize the rxode2 expression to speed up calculation. By default this is turned on.
calcTables	This boolean is to determine if the foceiFit will calculate tables. By default this is TRUE
compress	Should the object have compressed items
ci	Confidence level for some tables. By default this is 0.95 or 95% confidence.
sigdigTable	Significant digits in the final output table. If not specified, then it matches the significant digits in the 'sigdig' optimization algorithm. If 'sigdig' is NULL, use 3.
absolutePath	Boolean indicating if the absolute path should be used for the monolix runs
modelName	Model name used to generate the NONMEM output. If NULL try to infer from the model name (could be x if not clear). Otherwise use this character for outputs.
muRefCovAlg	This controls if algebraic expressions that can be mu-referenced are treated as mu-referenced covariates by:
	1. Creating a internal data-variable 'nlmixrMuDerCov#' for each algebraic mureferenced expression
	2. Change the algebraic expression to 'nlmixrMuDerCov# * mu_cov_theta'
	3. Use the internal mu-referenced covariate for saem
	4. After optimization is completed, replace 'model()' with old 'model()' expression
	5. Remove 'nlmixrMuDerCov#' from nlmix2 output
	In general, these covariates should be more accurate since it changes the system to a linear compartment model. Therefore, by default this is 'TRUE'.
run	Should monolix be run and the results be imported to nlmixr2? (Default is TRUE)
	Ignored parameters

Details

If runCommand is given as a string, it will be called with the system() command like: runCommand mlxtran.

For example, if runCommand="'/path/to/monolix/mlxbsub2021'-p" then the command line used would look like the following:

'/path/to/monolix/mlxbsub2021' monolix.mlxtran

If runCommand is given as a function, it will be called as FUN(mlxtran, directory, ui) to run Monolix. This allows you to run Monolix in any way that you may need, as long as you can write it in R. babelmixr2 will wait for the function to return before proceeding.

If runCommand is NA, nlmixr() will stop after writing the model files and without starting Monolix.

Note that you can get the translated monolix components from a parsed/compiled rxode2 ui object with ui\$monolixModel and ui\$mlxtran

Value

A monolix control object

nlmixr2Est.pknca 13

Author(s)

Matthew Fidler

nlmixr2Est.pknca

Estimate starting parameters using PKNCA

Description

Estimate starting parameters using PKNCA

Usage

```
## S3 method for class 'pknca'
nlmixr2Est(env, ...)
```

Arguments

env Environment for the nlmixr2 estimation routines.

This needs to have:

- rxode2 ui object in '\$ui'

- data to fit in the estimation routine in '\$data'

- control for the estimation routine's control options in '\$ui'

Other arguments provided to 'nlmixr2Est()' provided for flexibility but not cur-

rently used inside nlmixr

Details

Parameters are estimated as follows:

- ka 4 half-lives to Tmax but not higher than 3: log(2)/(tmax/4)
- · vc Inverse of dose-normalized Cmax
- cl Estimated as the median clearance
- vp, vp22- and 4-fold the vc, respectively by default, controlled by the vpMult and vp2Mult arguments to pkncaControl
- q,q2 0.5- and 0.25-fold the cl, respectively by default, controlled by the qMult and q2Mult arguments to pkncaControl

The bounds for the parameter estimates are set to 10% of the first percentile and 10 times the 99th percentile. (For ka, the lower bound is set to the lower of 10% of the first percentile or 0.03 and the upper bound is not modified from 10 times the 99th percentile.)

Parameter estimation methods may be changed in a future version.

Value

A model with updated starting parameters. In the model a new element named "nca" will be available which includes the PKNCA results used for the calculation.

14 nonmemControl

nonmemControl

NONMEM estimation control

Description

NONMEM estimation control

Usage

```
nonmemControl(
  est = c("focei", "imp", "its", "posthoc"),
 advan0de = c("advan13", "advan8", "advan6"),
  cov = c("r,s", "r", "s", ""),
 maxeval = 1e+05,
  tol = 6,
  atol = 12,
  sstol = 6,
  ssatol = 12,
  sigl = 12,
  sigdig = 3,
  print = 1,
  extension = getOption("babelmixr2.nmModelExtension", ".nmctl"),
  outputExtension = getOption("babelmixr2.nmOutputExtension", ".lst"),
  runCommand = getOption("babelmixr2.nonmem", ""),
  iniSigDig = 5,
  protectZeros = FALSE,
 muRef = TRUE,
  addProp = c("combined2", "combined1"),
  rxControl = NULL,
  sumProd = FALSE,
  optExpression = TRUE,
  calcTables = TRUE,
  compress = TRUE,
  ci = 0.95,
  sigdigTable = NULL,
  readRounding = FALSE,
  readBadOpt = FALSE,
  niter = 100L,
  isample = 1000L,
  iaccept = 0.4,
  iscaleMin = 0.1,
  iscaleMax = 10,
  df = 4,
  seed = 14456,
 mapiter = 1,
 mapinter = 0,
  noabort = TRUE,
```

nonmemControl 15

```
modelName = NULL,
muRefCovAlg = TRUE,
run = TRUE,
...
)
```

Arguments

est NONMEM estimation method

advan0de The ODE solving method for NONMEM

cov The NONMEM covariance method

maxeval NONMEM's maxeval (for non posthoc methods)

tol NONMEM tolerance for ODE solving advan
atol NONMEM absolute tolerance for ODE solving
sstol NONMEM tolerance for steady state ODE solving

ssatol NONMEM absolute tolerance for steady state ODE solving

sigl NONMEM sigl estimation option sigdig the significant digits for NONMEM print The print number for NONMEM

extension NONMEM file extensions

outputExtension

Extension to use for the NONMEM output listing

runCommand Command to run NONMEM (typically the path to "nmfe75") or a function. See

the details for more information.

iniSigDig How many significant digits are printed in \$THETA and \$OMEGA when the

estimate is zero. Also controls the zero protection numbers

protectZeros Add methods to protect divide by zero

muRef Automatically mu-reference the control stream

addProp, sumProd, optExpression, calcTables, compress, ci, sigdigTable

Passed to nlmixr2est::foceiControl

rxControl Options to pass to rxode2::rxControl for simulations

readRounding Try to read NONMEM output when NONMEM terminated due to rounding

errors

readBadOpt Try to read NONMEM output when NONMEM terminated due to an apparent

failed optimization

niter number of iterations in NONMEM estimation methods isample Isample argument for NONMEM ITS estimation method

iaccept Iaccept for NONMEM ITS estimation methods

iscaleMin parameter for IMP NONMEM method (ISCALE_MIN)
iscaleMax parameter for IMP NONMEM method (ISCALE_MAX)

df degrees of freedom for IMP method

16 nonmemControl

seed is the seed for NONMEM methods

mapiter the number of map iterations for IMP method mapinter is the MAPINTER parameter for the IMP method

noabort Add the NOABORT option for \$EST

modelName Model name used to generate the NONMEM output. If NULL try to infer from the

model name (could be x if not clear). Otherwise use this character for outputs.

muRefCovAlg This controls if algebraic expressions that can be mu-referenced are treated as

mu-referenced covariates by:

1. Creating a internal data-variable 'nlmixrMuDerCov#' for each algebraic mu-

referenced expression

2. Change the algebraic expression to 'nlmixrMuDerCov# * mu_cov_theta'

3. Use the internal mu-referenced covariate for saem

4. After optimization is completed, replace 'model()' with old 'model()' expres-

sion

5. Remove 'nlmixrMuDerCov#' from nlmix2 output

In general, these covariates should be more accurate since it changes the system

to a linear compartment model. Therefore, by default this is 'TRUE'.

run Should NONMEM be run (and the files imported to nlmixr2); default is TRUE,

but FALSE will simply create the NONMEM control stream and data file.

... optional genRxControl argument controlling automatic rxControl generation.

Details

If runCommand is given as a string, it will be called with the system() command like: runCommand controlFile outputFile.

For example, if runCommand="'/path/to/nmfe75'" then the command line used would look like the following:

'/path/to/nmfe75' one.cmt.nmctl one.cmt.lst

If runCommand is given as a function, it will be called as FUN(ctl, directory, ui) to run NON-MEM. This allows you to run NONMEM in any way that you may need, as long as you can write it in R. babelmixr2 will wait for the function to return before proceeding.

If runCommand is NA, nlmixr() will stop after writing the model files and without starting NON-MEM.

Value

babelmixr2 control option for generating NONMEM control stream and reading it back into babelmixr2/nlmixr2

Author(s)

Matthew L. Fidler

Examples

nonmemControl()

pkncaControl 17

pkncaControl

PKNCA estimation control

Description

PKNCA estimation control

Usage

```
pkncaControl(
  concu = NA_character_,
  doseu = NA_character_,
  timeu = NA_character_,
  volumeu = NA_character_,
  vpMult = 2,
  qMult = 1/2,
  vp2Mult = 4,
  q2Mult = 1/4
  dvParam = "cp",
  groups = character(),
  sparse = FALSE,
  ncaData = NULL,
 ncaResults = NULL,
  rxControl = rxode2::rxControl()
)
```

Arguments

concu, doseu, timeu

concentration, dose, and time units from the source data (passed to PKNCA::pknca_units_table()).

volumeu compartment volume for the model (if NULL, simplified units from source data

will be used)

vpMult, qMult, vp2Mult, q2Mult

Multipliers for vc and cl to provide initial estimates for vp, q, vp2, and q2

dvParam The parameter name in the model that should be modified for concentration unit

conversions. It must be assigned on a line by itself, separate from the residual

error model line.

groups Grouping columns for NCA summaries by group (required if sparse = TRUE)

sparse Are the concentration-time data sparse PK (commonly used in small nonclinical

species or with terminal or difficult sampling) or dense PK (commonly used in

clinical studies or larger nonclinical species)?

ncaData Data to use for calculating NCA parameters. Typical use is when a subset of the

original data are informative for NCA.

ncaResults Already computed NCA results (a PKNCAresults object) to bypass automatic

calculations. At least the following parameters must be calculated in the NCA:

tmax, cmax.dn, cl.last

rxControl Control options sent to rxode2::rxControl()

Value

A list of parameters

popedControl

Control for a PopED design task

Description

Control for a PopED design task

Usage

```
popedControl(
  stickyRecalcN = 4,
 maxOdeRecalc = 5,
  odeRecalcFactor = 10^{\circ}(0.5),
 maxn = NULL,
  rxControl = NULL,
  sigdig = 4,
  important = NULL,
  unimportant = NULL,
 iFIMCalculationType = c("reduced", "full", "weighted", "loc", "reducedPFIM", "fullABC",
    "largeMat", "reducedFIMABC"),
  iApproximationMethod = c("fo", "foce", "focei", "foi"),
  iFOCENumInd = 1000,
  prior_fim = matrix(0, 0, 1),
  d_switch = c("d", "ed"),
  ofv_calc_type = c("lnD", "d", "a", "Ds", "inverse"),
  strEDPenaltyFile = "",
  ofv_fun = NULL,
  iEDCalculationType = c("mc", "laplace", "bfgs-laplace"),
  ED_samp_size = 45,
  bLHS = c("hypercube", "random"),
  bUseRandomSearch = TRUE,
  bUseStochasticGradient = TRUE,
  bUseLineSearch = TRUE,
  bUseExchangeAlgorithm = FALSE,
  bUseBFGSMinimizer = FALSE,
  bUseGrouped_xt = FALSE,
  EACriteria = c("modified", "fedorov"),
  strRunFile = "",
  poped_version = NULL,
  modtit = "PopED babelmixr2 model",
  output_file = "PopED_output_summary",
```

```
output_function_file = "PopED_output_";
strIterationFileName = "PopED_current.R",
user_data = NULL,
ourzero = 1e-05,
dSeed = NULL,
line_opta = NULL,
line_optx = NULL,
bShowGraphs = FALSE,
use_logfile = FALSE,
m1_switch = c("central", "complex", "analytic", "ad"),
m2_switch = c("central", "complex", "analytic", "ad"),
hle_switch = c("central", "complex", "ad"),
gradff_switch = c("central", "complex", "analytic", "ad"),
gradfg_switch = c("central", "complex", "analytic", "ad"),
grad_all_switch = c("central", "complex"),
rsit_output = 5,
sgit_output = 1,
hm1 = 1e-05,
hlf = 1e-05,
hlg = 1e-05,
hm2 = 1e-05,
hgd = 1e-05,
hle = 1e-05,
AbsTol = 1e-06,
RelTol = 1e-06,
iDiffSolverMethod = NULL,
bUseMemorySolver = FALSE,
rsit = 300,
sgit = 150,
intrsit = 250,
intsgit = 50,
maxrsnullit = 50,
convergence_eps = 1e-08,
rslxt = 10,
rsla = 10,
cfaxt = 0.001,
cfaa = 0.001,
bGreedyGroupOpt = FALSE,
EAStepSize = 0.01,
EANumPoints = FALSE,
EAConvergenceCriteria = 1e-20,
bEANoReplicates = FALSE,
BFGSProjectedGradientTol = 1e-04,
BFGSTolerancef = 0.001,
BFGSToleranceg = 0.9,
BFGSTolerancex = 0.1,
ED_diff_it = 30,
ED_diff_percent = 10,
```

```
line_search_it = 50,
Doptim_iter = 1,
iCompileOption = c("none", "full", "mcc", "mpi"),
compileOnly = FALSE,
iUseParallelMethod = c("mpi", "matlab"),
MCC_Dep = NULL,
strExecuteName = "calc_fim.exe",
iNumProcesses = 2,
iNumChunkDesignEvals = -2,
Mat_Out_Pre = "parallel_output",
strExtraRunOptions = "",
dPollResultTime = 0.1,
strFunctionInputName = "function_input",
bParallelRS = FALSE,
bParallelSG = FALSE,
bParallelMFEA = FALSE,
bParallelLS = FALSE,
groupsize = NULL,
time = "time",
timeLow = "low"
timeHi = "high",
id = "id",
m = NULL,
x = NULL,
ni = NULL,
maxni = NULL,
minni = NULL,
maxtotni = NULL,
mintotni = NULL,
maxgroupsize = NULL,
mingroupsize = NULL,
maxtotgroupsize = NULL,
mintotgroupsize = NULL,
xt\_space = NULL,
a = NULL,
maxa = NULL,
mina = NULL,
a_{space} = NULL,
x_space = NULL,
use_grouped_xt = FALSE,
grouped_xt = NULL,
use_grouped_a = FALSE,
grouped_a = NULL,
use\_grouped\_x = FALSE,
grouped_x = NULL,
our_zero = NULL,
auto_pointer = "",
user_distribution_pointer = "",
```

```
minxt = NULL,
maxxt = NULL,
discrete_xt = NULL,
discrete_a = NULL,
fixRes = FALSE,
script = NULL,
overwrite = TRUE,
literalFix = TRUE,
opt_xt = FALSE,
opt_a = FALSE,
opt_x = FALSE,
opt_samps = FALSE,
optTime = TRUE,
...
)
```

Arguments

stickyRecalcN

The number of bad ODE solves before reducing the atol/rtol for the rest of the

problem.

maxOdeRecalc

Maximum number of times to reduce the ODE tolerances and try to resolve the

system if there was a bad ODE solve.

odeRecalcFactor

The ODE recalculation factor when ODE solving goes bad, this is the factor the

rtol/atol is reduced

maxn

Maximum number of design points for optimization; By default this is declared

by the maximum number of design points in the babelmixr2 dataset (when NULL)

rxControl

'rxode2' ODE solving options during fitting, created with 'rxControl()'

sigdig

Optimization significant digits. This controls:

- The tolerance of the inner and outer optimization is 10^-sigdig
- The tolerance of the ODE solvers is 0.5*10^(-sigdig-2); For the sensitivity equations and steady-state solutions the default is 0.5*10^(-sigdig-1.5) (sensitivity changes only applicable for liblsoda)
- The tolerance of the boundary check is $5 \times 10^{\circ}$ (-sigdig + 1)

important

character vector of important parameters or NULL for default. This is used with Ds-optimality

unimportant

character vector of unimportant parameters or NULL for default. This is used with Ds-optimality

iFIMCalculationType

can be either an integer or a named value of the Fisher Information Matrix type:

- 0/"full" = Full FIM
- 1/"reduced" = Reduced FIM
- 2/"weighted" = weighted models
- 3/"loc" = Loc models
- 4/"reducedPFIM" = reduced FIM with derivative of SD of sigma as in PFIM

- 5/"fullABC" = FULL FIM parameterized with A,B,C matrices & derivative of variance
- 6/"largeMat" = Calculate one model switch at a time, good for large matrices
- 7/"reducedFIMABC" = =Reduced FIM parameterized with A,B,C matrices & derivative of variance

iApproximationMethod

Approximation method for model, 0=FO, 1=FOCE, 2=FOCEI, 3=FOI

iFOCENumInd integer; number of individuals in focei solve

prior_fim matrix; prior FIM

d_switch integer or character option:

- 0/"ed" = ED design
- 1/"d" = D design

ofv_calc_type objective calculation type:

- 1/"d" = D-optimality". Determinant of the FIM: det(FIM)
- 2/"a" = "A-optimality". Inverse of the sum of the expected parameter variances: 1/trace_matrix(inv(FIM))
- 4/"lnD" = "lnD-optimality". Natural logarithm of the determinant of the FIM: log(det(FIM))
- 6/"Ds" = "Ds-optimality". Ratio of the Determinant of the FIM and the Determinant of the uninteresting rows and columns of the FIM: det(FIM)/det(FIM_u)
- 7/"inverse" = Inverse of the sum of the expected parameter RSE: 1/sum(get_rse(FIM,poped.db,use_parameter RSE)

strEDPenaltyFile

Penalty function name or path and filename, empty string means no penalty. User defined criterion can be defined this way.

ofv_fun

User defined function used to compute the objective function. The function must have a poped database object as its first argument and have "..." in its argument list. Can be referenced as a function or as a file name where the function defined in the file has the same name as the file. e.g. "cost.txt" has a function named "cost" in it.

iEDCalculationType

ED Integral Calculation type:

- 0/"mc" = Monte-Carlo-Integration
- 1/"laplace" = Laplace Approximation
- 2/"bfgs-laplace" = BFGS Laplace Approximation

ED_samp_size Sample size for E-family sampling

bLHS How to sample from distributions in E-family calculations. 0=Random Sampling, 1=LatinHyperCube –

bUseRandomSearch

******START OF Optimization algorithm SPECIFICATION OPTIONS*********

Use random search (1=TRUE, 0=FALSE)

bUseStochasticGradient

Use Stochastic Gradient search (1=TRUE, 0=FALSE)

bUseLineSearch Use Line search (1=TRUE, 0=FALSE)

bUseExchangeAlgorithm

Use Exchange algorithm (1=TRUE, 0=FALSE)

bUseBFGSMinimizer

Use BFGS Minimizer (1=TRUE, 0=FALSE)

bUseGrouped_xt Use grouped time points (1=TRUE, 0=FALSE).

EACriteria Exchange Algorithm Criteria:

• 1/"modified" = Modified

• 2/"fedorov" = Fedorov

strRunFile Filename and path, or function name, for a run file that is used instead of the

regular PopED call.

poped_version • ******START OF Labeling and file names SPECIFICATION OPTIONS*********

The current PopED version

modtit The model title

output_file Filename and path of the output file during search

output_function_file

Filename suffix of the result function file

strIterationFileName

Filename and path for storage of current optimal design

user_data ******START OF Miscellaneous SPECIFICATION OPTIONS********

User defined data structure that, for example could be used to send in data to the

model

ourzero Value to interpret as zero in design

dSeed The seed number used for optimization and sampling – integer or -1 which cre-

ates a random seed as.integer(Sys.time()) or NULL.

line_opta Vector for line search on continuous design variables (1=TRUE,0=FALSE)

line_optx Vector for line search on discrete design variables (1=TRUE,0=FALSE)

bShowGraphs Use graph output during search

use_logfile If a log file should be used (0=FALSE, 1=TRUE)

m1_switch Method used to calculate M1:

• 1/"central" = Central difference

• 0/"complex" = Complex difference

• 20/"analytic" = Analytic derivative

• 30/"ad" = Automatic differentiation

m2_switch Method used to calculate M2:

• 1/"central" = Central difference

• 0/"complex" = Complex difference

• 20/"analytic" = Analytic derivative

• 30/"ad" = Automatic differentiation

hle_switch Method used to calculate linearization of residual error:

	• 1/"central" = Central difference
	• 0/"complex" = Complex difference
	• 30/"ad" = Automatic differentiation
gradff_switch	Method used to calculate the gradient of the model:
	• 1/"central" = Central difference
	• 0/"complex" = Complex difference
	• 20/"analytic" = Analytic derivative
	• 30/"ad" = Automatic differentiation
gradfg_switch	Method used to calculate the gradient of the parameter vector g:
	• 1/"central" = Central difference
	• 0/"complex" = Complex difference
	• 20/"analytic" = Analytic derivative
anad all awita	• 30/"ad" = Automatic differentiation
grad_all_switc	Method used to calculate all the gradients:
	• 1/"central" = Central difference
	• 0/"complex" = Complex difference
rsit_output	Number of iterations in random search between screen output
sgit_output	Number of iterations in stochastic gradient search between screen output
hm1	Step length of derivative of linearized model w.r.t. typical values
hlf	Step length of derivative of model w.r.t. g
hlg	Step length of derivative of g w.r.t. b
hm2	Step length of derivative of variance w.r.t. typical values
hgd 	Step length of derivative of OFV w.r.t. time
hle	Step length of derivative of model w.r.t. sigma
AbsTol	The absolute tolerance for the diff equation solver
RelTol	The relative tolerance for the diff equation solver
iDiffSolverMet	
hllooMomony Cc 1	The diff equation solver method, NULL as default.
bUseMemorySolv	If the differential equation results should be stored in memory (1) or not (0)
rsit	Number of Random search iterations
1511	number of Nandolli Scaleli Reladolls

rsit Number of Random search iterations
sgit Number of stochastic gradient iterations
intrsit Number of Random search iterations with discrete optimization.
Intsgit Number of Stochastic Gradient search iterations with discrete optimization
maxrsnullit Iterations until adaptive narrowing in random search

convergence_eps

Stochastic Gradient convergence value, (difference in OFV for D-optimal, difference in gradient for ED-optimal)

rslxt Random search locality factor for sample times

rsla Random search locality factor for covariates

cfaxt Stochastic Gradient search first step factor for sample times
cfaa Stochastic Gradient search first step factor for covariates

bGreedyGroupOpt

Use greedy algorithm for group assignment optimization

EAStepSize Exchange Algorithm StepSize
EANumPoints Exchange Algorithm NumPoints

 ${\tt EAConvergenceCriteria}$

Exchange Algorithm Convergence Limit/Criteria

bEANoReplicates

Avoid replicate samples when using Exchange Algorithm

BFGSProjectedGradientTol

BFGS Minimizer Convergence Criteria Normalized Projected Gradient Toler-

ance

BFGSTolerancef BFGS Minimizer Line Search Tolerance f
BFGSToleranceg BFGS Minimizer Line Search Tolerance g
BFGSTolerancex BFGS Minimizer Line Search Tolerance x

ED_diff_it Number of iterations in ED-optimal design to calculate convergence criteria

ED_diff_percent

ED-optimal design convergence criteria in percent

line_search_it Number of grid points in the line search

Doptim_iter Number of iterations of full Random search and full Stochastic Gradient if line

search is not used

iCompileOption Compile options for PopED

• "none"/-1 = No compilation

• "full/0 or 3 = Full compilation

• "mcc"/1 or 4 = Only using MCC (shared lib)

• "mpi"/2 or 5 = Only MPI,

When using numbers, option 0,1,2 runs PopED and option 3,4,5 stops after compilation.

When using characters, the option compileOnly determines if the model is only compiled (and PopED is not run).

compileOnly logical; only compile the model, do not run PopED (in conjunction with iCompileOption) iUseParallelMethod

Parallel method to use

• 0/"matlab"= Matlab PCT

• 1/"mpi" = MPI

MCC_Dep Additional dependencies used in MCC compilation (mat-files), if several space

separated

strExecuteName Compilation output executable name

iNumProcesses Number of processes to use when running in parallel (e.g. 3 = 2 workers, 1 job

manager)

iNumChunkDesignEvals

Number of design evaluations that should be evaluated in each process before

getting new work from job manager

Mat_Out_Pre The prefix of the output mat file to communicate with the executable

strExtraRunOptions

Extra options send to e\$g. the MPI executable or a batch script, see execute_parallel\$m

for more information and options

dPollResultTime

Polling time to check if the parallel execution is finished

strFunctionInputName

The file containing the popedInput structure that should be used to evaluate the

designs

bParallelRS If the random search is going to be executed in parallel

bParallelSG If the stochastic gradient search is going to be executed in parallel

bParallelMFEA If the modified exchange algorithm is going to be executed in parallel

bParallelLS If the line search is going to be executed in parallel

groupsize Vector defining the size of the different groups (num individuals in each group).

If only one number then the number will be the same in every group.

time string that represents the time in the dataset (ie xt)
timeLow string that represents the lower design time (ie minxt)
timeHi string that represents the upper design time (ie maxmt)

id The id variable

m Number of groups in the study. Each individual in a group will have the same

design.

x A matrix defining the initial discrete values for the model Each row is a group/individual.

ni Vector defining the number of samples for each group.

maxni • ******START OF DESIGN SPACE OPTIONS********

Max number of samples per group/individual

minni Min number of samples per group/individual

Mumber defining the maximum number of samples allowed in the experiment.

Number defining the minimum number of samples allowed in the experiment.

maxgroupsize Vector defining the max size of the different groups (max number of individuals

in each group)

mingroupsize Vector defining the min size of the different groups (min num individuals in each

group) –

maxtotgroupsize

The total maximal groupsize over all groups

mintotgroupsize

The total minimal groupsize over all groups

xt_space	Cell array cell defining the discrete variables allowed for each xt value. Can also be a vector of values c(1:10) (same values allowed for all xt), or a list of lists list(1:10, 2:23, 4:6) (one for each value in xt in row major order or just for one row in xt, and all other rows will be duplicated).
a	Matrix defining the initial continuous covariate values. n_rows=number of groups, n_cols=number of covariates. If the number of rows is one and the number of groups > 1 then all groups are assigned the same values.
maxa	Vector defining the max value for each covariate. If a single value is supplied then all a values are given the same max value
mina	Vector defining the min value for each covariate. If a single value is supplied then all a values are given the same max value
a_space	Cell array cell defining the discrete variables allowed for each a value. Can also be a list of values list(1:10) (same values allowed for all a), or a list of lists list(1:10, 2:23, 4:6) (one for each value in a).
x_space	Cell array cell defining the discrete variables for each x value.
use_grouped_xt	Group sampling times between groups so that each group has the same values (TRUE or FALSE).
grouped_xt	Matrix defining the grouping of sample points. Matching integers mean that the points are matched. Allows for finer control than use_grouped_xt
use_grouped_a	Group continuous design variables between groups so that each group has the same values (TRUE or FALSE).
grouped_a	Matrix defining the grouping of continuous design variables. Matching integers mean that the values are matched. Allows for finer control than use_grouped_a.
use_grouped_x	Group discrete design variables between groups so that each group has the same values (TRUE or FALSE).
grouped_x	Matrix defining the grouping of discrete design variables. Matching integers mean that the values are matched. Allows for finer control than use_grouped_x.
our_zero	Value to interpret as zero in design.
auto_pointer	Filename and path, or function name, for the Autocorrelation function, empty string means no autocorrelation
user_distribut	
	Filename and path, or function name, for user defined distributions for E-family designs
minxt	Matrix or single value defining the minimum value for each xt sample. If a single value is supplied then all xt values are given the same minimum value
maxxt	Matrix or single value defining the maximum value for each xt sample. If a single value is supplied then all xt values are given the same maximum value.
discrete_xt	Cell array cell defining the discrete variables allowed for each xt value. Can also be a list of values list(1:10) (same values allowed for all xt), or a list of lists list(1:10, 2:23, 4:6) (one for each value in xt). See examples in create_design_space.
discrete_a	Cell array cell defining the discrete variables allowed for each a value. Can also be a list of values list(1:10) (same values allowed for all a), or a list of lists list(1:10, 2:23, 4:6) (one for each value in a). See examples in create_design_space.

when a poped dataset is created. You can manually reset with popedMultipleEndpointResetTimeIndex

fixRes	boolean; Fix the residuals to what is specified by the model
script	write a PopED/rxode2 script that can be modified for more fine control. The default is NULL.
	When script is TRUE, the script is returned as a lines that would be written to a file and with the class babelmixr2popedScript. This allows it to be printed as the script on screen.
	When script is a file name (with an R extension), the script is written to that file.
overwrite	[logical(1)] If TRUE, an existing file in place is allowed if it it is both readable and writable. Default is FALSE.
literalFix	boolean, substitute fixed population values as literals and re-adjust ui and parameter estimates after optimization; Default is 'TRUE'.
opt_xt	boolean to indicate if this is meant for optimizing times
opt_a	boolean to indicate if this is meant for optimizing covariates
opt_x	boolean to indicate if the discrete design variables be optimized
opt_samps	boolean to indicate if the sample optimizer is used (not implemented yet in PopED)
optTime	boolean to indicate if the global time indexer inside of babelmixr2 is reset if the times are different. By default this is TRUE. If FALSE you can get slightly better run times and possibly slightly different results. When optTime is FALSE the global indexer is reset every time the PopED rxode2 is setup for a problem or

Value

. . .

popedControl object

Author(s)

Matthew L. Fidler

 ${\tt popedGetMultipleEndpointModelingTimes}$

Get Multiple Endpoint Modeling Times

other parameters for PopED control

Description

This function takes a vector of times and a corresponding vector of IDs, groups the times by their IDs, initializes an internal C++ global TimeIndexer, that is used to efficiently lookup the final output from the rxode2 solve and then returns the sorted unique times.

The popedMultipleEndpointIndexDataFrame() function can be used to visualize the internal data structure inside R, but it does not show all the indexes in the case of time ties for a given ID. Rather it shows one of the indexs and the total number of indexes in the data.frame

Usage

```
popedGetMultipleEndpointModelingTimes(times, modelSwitch, sorted = FALSE)
popedMultipleEndpointIndexDataFrame(print = FALSE)
```

Arguments

times A numeric vector of times.

modelSwitch An integer vector of model switch indicator corresponding to the times

sorted A boolean indicating if the returned times should be sorted

print boolean for popedMultipleEndpointIndexDataFrame() when TRUE show each

id/index per time even though it may not reflect in the returned data.frame

Value

A numeric vector of unique times.

Examples

```
times <- c(1.1, 1.2, 1.3, 2.1, 2.2, 3.1)
modelSwitch <- c(1, 1, 1, 2, 2, 3)
sortedTimes <- popedGetMultipleEndpointModelingTimes(times, modelSwitch, TRUE)
print(sortedTimes)

# now show the output of the data frame representing the model
# switch to endpoint index

popedMultipleEndpointIndexDataFrame()

# now show a more complex example with overlaps etc.

times <- c(1.1, 1.2, 1.3, 0.5, 2.2, 1.1, 0.75,0.75)
modelSwitch <- c(1, 1, 1, 2, 2, 2, 3, 3)
sortedTimes <- popedGetMultipleEndpointModelingTimes(times, modelSwitch, TRUE)
print(sortedTimes)

popedMultipleEndpointIndexDataFrame(TRUE) # Print to show individual matching</pre>
```

30 rxToMonolix

popedMultipleEndpointResetTimeIndex

Reset the Global Time Indexer for Multiple Endpoint Modeling

Description

This clears the memory and resets the global time indexer used for multiple endpoint modeling.

Usage

```
popedMultipleEndpointResetTimeIndex()
```

Value

NULL, called for side effects

Examples

popedMultipleEndpointResetTimeIndex()

rxToMonolix

Convert RxODE syntax to monolix syntax

Description

Convert RxODE syntax to monolix syntax

Usage

```
rxToMonolix(x, ui)
```

Arguments

x Expression ui rxode2 ui

Value

Monolix syntax

Author(s)

Matthew Fidler

rxToNonmem 31

rxToNonmem

Convert RxODE syntax to NONMEM syntax

Description

Convert RxODE syntax to NONMEM syntax

Usage

```
rxToNonmem(x, ui)
```

Arguments

x Expression ui rxode2 ui

Value

NONMEM syntax

Author(s)

Matthew Fidler

simplifyUnit

Simplify units by removing repeated units from the numerator and denominator

Description

Simplify units by removing repeated units from the numerator and denominator

Usage

```
simplifyUnit(numerator = "", denominator = "")
```

Arguments

numerator The numerator of the units (or the whole unit specification)

denominator The denominator of the units (or NULL if numerator is the whole unit specifi-

cation)

Details

NA or "" for numerator and denominator are considered unitless.

32 simplifyUnit

Value

The units specified with units that are in both the numerator and denominator cancelled.

See Also

Other Unit conversion: modelUnitConversion()

Examples

```
simplifyUnit("kg", "kg/mL")
# units that don't match exactly are not cancelled
simplifyUnit("kg", "g/mL")
```

Index

```
* Unit conversion
    modelUnitConversion, 9
    simplifyUnit, 31
.setupPopEDdatabase, 2
as.nlmixr(as.nlmixr2), 3
as.nlmixr2, 3
babel.poped.database, 5
bblDatToMonolix, 5
bblDatToMrgsolve (bblDatToMonolix), 5
bblDatToNonmem (bblDatToMonolix), 5
bblDatToPknca (bblDatToMonolix), 5
bblDatToRxode (bblDatToMonolix), 5
cell, 27
create_design_space, 27
getStandardColNames, 8
modelUnitConversion, 9, 32
monolixControl, 10
nlmixr2Est.pknca, 13
nonmemControl, 14
pkncaControl, 17
popedControl, 18
poped {\tt GetMultipleEndpointModelingTimes},
poped \verb|MultipleEndpointIndexDataFrame|
        (popedGetMultipleEndpointModelingTimes),
poped \verb|MultipleEndpointResetTimeIndex|,
rxToMonolix, 30
rxToNonmem, 31
simplifyUnit, 9, 31
```