# Package 'monolix2rx'

October 24, 2024

Version 0.0.3 **Description** 'Monolix' is a tool for running mixed effects model using 'saem'. This tool allows you to convert 'Monolix' models to 'rxode2' (Wang, Hallow and James (2016) <doi:10.1002/psp4.12052>) using the form compatible with 'nlmixr2' (Fidler et al (2019) <doi:10.1002/psp4.12445>). If available, the 'rxode2' model will read in the 'Monolix' data and compare the simulation for the population model individual model and residual model to immediately show how well the translation is performing. This saves the model development time for people who are creating an 'rxode2' model manually. Additionally, this package reads in all the information to allow simulation with uncertainty (that is the number of observations, the number of subjects, and the covariance matrix) with a 'rxode2' model. This is complementary to the 'babelmixr2' package that translates 'nlmixr2' models to 'Monolix' and can convert the objects converted from 'monolix2rx' to a full 'nlmixr2' fit. While not required, you can get/install the 'lixoftConnectors' package in the 'Monolix' installation, as described at the following url <a href="https://monolixsuite.slp-software.com/">https://monolixsuite.slp-software.com/</a> r-functions/2024R1/installation-and-initialization>. When 'lixoftConnectors' is available, 'Monolix' can be used to load its model library instead manually setting up text files (which only works with old versions of

License MIT + file LICENSE

'Monolix').

URL https://nlmixr2.github.io/monolix2rx/,
 https://github.com/nlmixr2/monolix2rx/

**Title** Converts 'Monolix' Models to 'rxode2'

**Encoding** UTF-8 **RoxygenNote** 7.3.2

LinkingTo dparser, rxode2, Rcpp

**Imports** Rcpp, checkmate, cli, dparser, withr, ggplot2, ggforce, stringi, crayon, lotri, magrittr, rxode2 (>= 3.0.0),

**Suggests** devtools, testthat (>= 3.0.0), xgxr, vdiffr, lixoftConnectors

Config/testthat/edition 3

2 mlxtran

#### Config/Needs/website rmarkdown

**NeedsCompilation** yes

**Author** Matthew Fidler [aut, cre] (<a href="https://orcid.org/0000-0001-8538-6691">https://orcid.org/0000-0001-8538-6691</a>),
Justin Wilkins [ctb] (<a href="https://orcid.org/0000-0002-7099-9396">https://orcid.org/0000-0002-7099-9396</a>)

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

**Repository** CRAN

**Date/Publication** 2024-10-24 10:40:03 UTC

# **Contents**

mlxtran														 				2
mlxTxt														 				3
monolix	2rx													 				4

Index

mlxtran

Read and parse mlxtran lines

#### **Description**

Read and parse mlxtran lines

# Usage

```
mlxtran(file, equation = FALSE, update = FALSE)
```

# Arguments

file mlxtran file to process

 $equation \qquad \qquad parse \ the \ equation \ block \ to \ rxode2 \ (some \ models \ cannot \ be \ translated)$ 

update when true, try to update the parameter block to the final parameter estimates

### Value

mlxtran object

#### Author(s)

Matthew L. Fidler

mlxTxt 3

#### **Examples**

```
# First load in the model; in this case the theo model
# This is modified from the Monolix demos by saving the model
# File as a text file (hence you can access without model library)
# setup.
#
# This example is also included in the monolix2rx package, so
# you refer to the location with `system.file()`:

pkgTheo <- system.file("theo", package="monolix2rx")

mlx <- mlxtran(file.path(pkgTheo, "theophylline_project.mlxtran"))
mlx</pre>
```

mlxTxt

Get equation block from a Monolix model txt file

#### **Description**

Get equation block from a Monolix model txt file

#### Usage

```
mlxTxt(file, retFile = FALSE)
```

#### **Arguments**

file string representing the model text file. Can be lib:fileName.txt if library setup/available retFile boolean that tells mlxTxt() to return the file name instead of error if the file does not exist

#### Value

parsed equation or file name

#### Author(s)

Matthew L. Fidler

#### **Examples**

```
# First load in the model; in this case the theo model
# This is modified from the Monolix demos by saving the model
# File as a text file (hence you can access without model library)
# setup.
#
# This example is also included in the monolix2rx package, so
# you refer to the location with `system.file()`:
```

4 monolix2rx

```
pkgTheo <- system.file("theo", package="monolix2rx")
mod <- mlxTxt(file.path(pkgTheo, "oral1_1cpt_kaVCl.txt"))
mod</pre>
```

monolix2rx

Translate a monolix file to rxode2

# Description

Translate a monolix file to rxode2

# Usage

```
monolix2rx(
  mlxtran,
  update = TRUE,
  thetaMatType = c("sa", "lin"),
  sd = 1,
  cor = 1e-05,
  theta = 0.5,
  ci = 0.95,
  sigdig = 3,
  envir = parent.frame()
)
```

# Arguments

mlxtran	file name for mlxtran to translate to rxode2
update	is a boolean that represents if the final parameter estimates should be used for the translation (when present) $\frac{1}{2}$
thetaMatType	This lists the preferred source for thetaMat covariance matrix. By default it is sa for simulated annealing, though you could use lin for linearized covariance calculation. If only one is present, then use whatever is present
sd	Default standard deviation for between subject variability/inter-occasion variability that are missing.
cor	Default correlation for missing correlations estimate
theta	default population estimate
ci	confidence interval for validation, by default 0.95
sigdig	number of significant digits for validation, by default 3
envir	represents the environment used for evaluating the corresponding rxode2 function

monolix2rx 5

#### Value

rxode2 model

#### Author(s)

Matthew L. Fidler

# **Examples**

```
# First load in the model; in this case the theo model
# This is modified from the Monolix demos by saving the model
# File as a text file (hence you can access without model library)
# setup.
#
# This example is also included in the monolix2rx package, so
# you refer to the location with `system.file()`:

pkgTheo <- system.file("theo", package="monolix2rx")

rx <- monolix2rx(file.path(pkgTheo, "theophylline_project.mlxtran"))

pkgCov <- system.file("cov", package="monolix2rx")

rx <- monolix2rx(file.path(pkgCov, "warfarin_covariate3_project.mlxtran"))
rx</pre>
```

# **Index**

mlxtran, 2
mlxTxt, 3
monolix2rx, 4