Package 'nonmem2rx'

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Type Package

Title Converts 'NONMEM' Models to 'rxode2'

Version 0.1.6

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Description 'NONMEM' has been a tool for running nonlinear mixed effects models since the 80s and is still used today (Bauer 2019 <doi:10.1002/psp4.12404>). This tool allows you to convert 'NONMEM' models to 'rxode2' (Wang, Hallow and James (2016) <doi:10.1002/psp4.12052>) and with simple models 'nlmixr2' syntax (Fidler et al (2019) <doi:10.1002/psp4.12445>). The 'nlmixr2' syntax requires the residual specification to be included and it is not always translated. If available, the 'rxode2' model will read in the 'NONMEM' data and compare the simulation for the population model ('PRED') individual model ('IPRED') and residual model ('IWRES') 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 'NONMEM' and can convert the objects converted from 'nonmem2rx' to a full 'nlmixr2' fit.

License GPL (>= 3)

URL https://nlmixr2.github.io/nonmem2rx/,
 https://github.com/nlmixr2/nonmem2rx/

Encoding UTF-8

LinkingTo dparser, Rcpp, rxode2

Imports checkmate, digest, dparser, lotri, Rcpp, rxode2 (> 2.0.13), magrittr, cli, tools, utils, data.table, qs, xml2, ggplot2, ggforce, crayon

Suggests devtools, testthat (>= 3.0.0), nonmemica, NMdata, nonmem2R, withr, nlme, dplyr, xgxr, vdiffr, knitr, rmarkdown, spelling

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Config/testthat/edition 3 RoxygenNote 7.3.2 VignetteBuilder knitr Language en-US NeedsCompilation yes Author Matthew Fidler [aut, cre] (https://orcid.org/0000-0001-8538-6691), Philip Delff [ctb], Gabriel Staples [ctb] (string insensitive compare) Repository CRAN Date/Publication 2024-11-28 19:30:02 UTC Contents as.nonmem2rx nmcov nmext nminfo

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Description

Convert a model to a nonmem2rx model

Usage

```
as.nonmem2rx(model1, model2, compress = TRUE)
```

Arguments

model1 Input model 1 model 2 Input model 2

compress boolean to compress the ui at the end

Value

nonmem2rx model

nmcov 3

Author(s)

Matthew L. Fidler

```
mod <- nonmem2rx(system.file("mods/cpt/runODE032.ctl", package="nonmem2rx"),</pre>
                   determineError=FALSE, 1st=".res", save=FALSE)
 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 - c1 * CENTRAL/v1
     d/dt(PERI) <- -k21 * PERI + k12 * CENTRAL
     f <- CENTRAL/scale1</pre>
     f ~ prop(RSV)
   })
 }
new <- try(as.nonmem2rx(mod2, mod))</pre>
if (!inherits(new, "try-error")) print(new, page=1)
```

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Description

Read in data file

Usage

```
nmcov(file, ...)
```

Arguments

file file name to read the results from

... other parameters passed to data.table::fread

Value

A matrix with covariance step from NONMEM

Author(s)

Philip Delff and Matthew L. Fidler

Examples

```
nmcov(system.file("mods/cpt/runODE032.cov", package="nonmem2rx"))
```

nmext

Reads the NONMEM .ext file for final parameter information

Description

Reads the NONMEM .ext file for final parameter information

Usage

```
nmext(file)
```

Arguments

file

File where the list is located

Value

return a list with \$theta, \$eta and \$eps

Author(s)

Matthew L. Fidler

```
nmext(system.file("run001.ext", package="nonmem2rx"))
```

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nminfo

Get the most accurate information you can get from NONMEM

Description

Get the most accurate information you can get from NONMEM

Usage

```
nminfo(
  file,
 mod = ".mod",
 xml = ".xml",
 ext = ".ext",
  cov = ".cov",
 phi = ".phi",
  lst = ".lst",
 useXml = TRUE,
 useExt = TRUE,
 useCov = TRUE,
 usePhi = TRUE,
 useLst = TRUE,
 strictLst = FALSE,
  verbose = FALSE
)
```

Arguments

file	nonmem file, like control stream, phi. This function will remove the extension to try to get the right information. It preferentially selects the most accurate estimates from the file.
mod	the NONMEM output extension, defaults to .mod
xml	the NONMEM xml file extension, defaults to .xml
ext	the NONMEM ext file extension, defaults to .ext
cov	the NONMEM covariance file extension, defaults to .cov
phi	the NONMEM eta/phi file extension, defaults to .phi
lst	the NONMEM output extension, defaults to .1st
useXml	if present, use the NONMEM xml file to import much of the NONMEM information
useExt	if present, use the NONMEM ext file to extract parameter estimates (default TRUE), otherwise defaults to parameter estimates extracted in the NONMEM output
useCov	if present, use the NONMEM cov file to import the covariance, otherwise import the covariance with list file

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usePhi if present, use the NONMEM phi file to extract etas (default TRUE), otherwise

defaults to etas in the tables (if present)

useLst if present, use the NONMEM lst file to extract NONMEM information strictLst The list parsing needs to be correct for a successful load (default FALSE).

verbose this is a flag to be more verbose when reading information in, by default this is

FALSE

Value

list of NONMEM information

Author(s)

Matthew L. Fidler

Examples

```
nminfo(system.file("mods/cpt/runODE032.res", package="nonmem2rx"))
```

nmlst

Reads the NONMEM .1st file for final parameter information

Description

Reads the NONMEM .1st file for final parameter information

Usage

```
nmlst(file, strictLst = FALSE)
```

Arguments

file File where the list is located

strictLst The list parsing needs to be correct for a successful load (default FALSE).

Value

return a list with \$theta, \$eta and \$eps and other information about the control stream

Author(s)

Matthew L. Fidler

```
nmlst(system.file("mods/DDMODEL00000322/HCQ1CMT.lst", package="nonmem2rx"))
nmlst(system.file("mods/DDMODEL00000302/run1.lst", package="nonmem2rx"))
nmlst(system.file("mods/DDMODEL00000301/run3.lst", package="nonmem2rx"))
nmlst(system.file("mods/cpt/runODE032.res", package="nonmem2rx"))
```

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nmtab

Read nonmem table file

Description

Read nonmem table file

Usage

```
nmtab(file, ...)
```

Arguments

```
file file name to read the results from
... other parameters passed to data.table::fread
```

Value

data frame of the read table

Author(s)

Philip Delff, Matthew L. Fidler

Examples

```
nmtab(system.file("mods/cpt/runODE032.csv", package="nonmem2rx"))
```

nmxml

Read a nonmem xml and create output similar to the nmlst()

Description

Read a nonmem xml and create output similar to the nmlst()

Usage

```
nmxml(xml)
```

Arguments

xml

xml file

Value

list of nonmem information

Author(s)

Matthew L. Fidler

Examples

```
nmxml(system.file("mods/cpt/runODE032.xml", package="nonmem2rx"))
```

nonmem2rx

Convert a NONMEM source file to a rxode model (nlmixr2-syle)

Description

Convert a NONMEM source file to a rxode model (nlmixr2-syle)

Usage

```
nonmem2rx(
  file,
  inputData = NULL,
  nonmemOutputDir = NULL,
  rename = NULL,
  tolowerLhs = TRUE,
  thetaNames = TRUE,
  etaNames = TRUE,
  cmtNames = TRUE,
  updateFinal = TRUE,
  determineError = TRUE,
  validate = getOption("nonmem2rx.validate", TRUE),
  nonmemData = FALSE,
  strictLst = FALSE,
  unintFixed = FALSE,
  extended = getOption("nonmem2rx.extended", FALSE),
  nLinesPro = 20L,
  delta = 1e-04,
  usePhi = TRUE,
  useExt = TRUE,
  useCov = TRUE,
  useXml = TRUE,
  useLst = TRUE,
  mod = ".mod",
  cov = ".cov",
  phi = ".phi",
  lst = getOption("nonmem2rx.lst", ".lst"),
  xml = ".xml",
  ext = ".ext",
  scanLines = getOption("nonmem2rx.scanLines", 50L),
  save = getOption("nonmem2rx.save", NA),
```

```
saveTime = getOption("nonmem2rx.saveTime", 15),
overwrite = getOption("nonmem2rx.overwrite", TRUE),
load = getOption("nonmem2rx.load", TRUE),
compress = getOption("nonmem2rx.compress", TRUE),
keep = getOption("nonmem2rx.keep", c("dfSub", "dfObs", "thetaMat", "sigma"))
)
```

Arguments

file NONMEM run file, like an .xml or .1st file or even a control stream

inputData this is a path to the input dataset (or NULL to determine from the dataset). Often

the input dataset may be different from the place it points to in the control stream because directories can be created to run NONMEM from a script. Because of this, when this is specified the input data will be assumed to be from here instead.

nonmemOutputDir

This is a path the the nonmem output directory. When not NULL it will assume that the directory for the output files is located here instead of where the control

stream currently exists.

rename When not NULL this should be a named character vector that contains the param-

eters that should be renamed. For example, if the model uses the variable YTYPE and has CMT it isn't compatible with rxode2/nlmixr2. You can change this for the input dataset and the model to create a new model that still reproduces the

NONMEM output by specifying rename=c(dvid="YTYPE")

tolowerLhs Boolean to change the lhs to lower case (default: TRUE)

thetaNames this could be a boolean indicating that the theta names should be changed to the

comment-labeled names (default: TRUE). This could also be a character vector

of the theta names (in order) to be replaced.

etaNames this could be a boolean indicating that the eta names should be changed to the

comment-labeled names (default: TRUE). This could also be a character vector

of the theta names (in order) to be replaced.

cmtNames this could be a boolean indicating that the compartment names should be changed

to the named compartments in the MODEL by COMP = (name) (default: TRUE). This could also be a character vector of the compartment names (in order) to be

replaced.

updateFinal Update the parsed model with the model estimates from the .1st output file.

determineError Boolean to try to determine the nlmixr2-style residual error model (like ipred

~ add(add.sd)), otherwise endpoints are not defined in the rxode2/nlmixr2

model (default: TRUE)

validate Boolean that this tool will attempt to "validate" the model by solving the derived

model under pred conditions (etas are zero and eps values are zero)

nonmemData Boolean that tells nonmem2rx to read in the nonmem data (if possible) even if

the model will not be validated (like if it is a simulation run or missing final parameter estimates). By default this is FALSE, nonmem data will not be integrated

into the nonmem2rx ui.

strictLst The list parsing needs to be correct for a successful load (default FALSE).

Treat uninteresting values as fixed parameters (default FALSE) unintFixed

Translate extended control streams from tools like wings for NONMEM extended

nLinesPro The number of lines to check for the \$PROBLEM statement.

delta this is the offset for NONMEM times that are tied

usePhi if present, use the NONMEM phi file to extract etas (default TRUE), otherwise

defaults to etas in the tables (if present)

useExt if present, use the NONMEM ext file to extract parameter estimates (default

TRUE), otherwise defaults to parameter estimates extracted in the NONMEM

output

if present, use the NONMEM cov file to import the covariance, otherwise import useCov

the covariance with list file

useXml if present, use the NONMEM xml file to import much of the NONMEM infor-

mation

useLst if present, use the NONMEM 1st file to extract NONMEM information

the NONMEM output extension, defaults to .mod mod

the NONMEM covariance file extension, defaults to .cov cov phi the NONMEM eta/phi file extension, defaults to .phi lst the NONMEM output extension, defaults to .1st xmlthe NONMEM xml file extension, defaults to .xml the NONMEM ext file extension, defaults to .ext ext

number of lines to scan for comment chars when IGNORE=@. default is 50 scanLines

This can be: save

• a NULL (meaning don't save),

• a logical (default FALSE, don't save) that when TRUE will use the base name of the control stream, append .qs and save the file using qs::qsave()

• A path to a file to write

Note that this file will be saved with qs::qsave() and can be loaded with

• A NA value which means save if the whole process (including validation)

takes too much time

saveTime The time that the translation/validation needs (in secs) before it will save to avoid having to rerun the model (default 15 for 15 seconds)

is a boolean to allow overwriting the save file (see load for more information).

a boolean that says to load the save file (if it exists) instead of re-running the translation and validation. Note if overwrite=TRUE and load=TRUE then this will overwrite based on time stamp of the files. If the save file is newer than the input file, then load that file, otherwise regenerate and overwrite. This works best if you point to an output file, like a .xml or listing file instead of the control

stream

a boolean indicating if the UI should be a compressed UI. If you are using this compress

for simulation with old versions of rxode2, the compressed ui is not supported, so this should be FALSE. Otherwise use TRUE if you are using a newer rxode2.

overwrite

load

keep

is a character vector of imported model items that are kept in the model itself; The defaults is "sigma" which keeps the sigma matrix in the model itself. You can add rxode2 solving options that are imported from NONMEM to keep in the model.

Details

Since some of these options you may want to set per project, the following options are queried:

- nonmem2rx.validate boolean to validate the model (default: TRUE)
- nonmem2rx.lst default extension for output (default: .lst)
- nonmem2rx.save should nonmem2rx save the model output?
- nonmem2rx.overwrite should nonmem2rx save output be overwritten (default TRUE)
- nonmem2rx.load should nonmem2rx load a saved model instead of translating and validating again? (default TRUE)
- nonmem2rx.extended should nonmem2rx support extended control streams? (default FALSE)
- nonmem2rx.compress should the ui be compressed or uncompressed (default: TRUE)

Value

rxode2 function

```
plot(mod, nrow=2, ncol=2, page=c(1,3), log="y")

#' or even all the individuals with
# plot(page=TRUE)

plot(mod, nrow=5, ncol=5, page=TRUE, log="y")

# you can also convert to a nlmixr2 object, but need babelmixr2 for
# that conversion
```

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