

RxODE user manual

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Introduction

Welcome to the RxODE user guide; **RxODE** is an R package for solving and simulating from ode-based models. These models are convert the RxODE mini-language to C and create a compiled dll for fast solving. ODE solving using RxODE has a few key parts:

- RxODE() which creates the C code for fast ODE solving based on a simple syntax (Chapter 6) related to Leibnitz notation.
- The event data, which can be:
 - a NONMEM or deSolve compatible data frame (Chapter 7), or
 - created with et() or EventTable() for easy simulation of events (Chapter 11)
 - The data frame can be augmented by adding time varying or adding individual covariates (iCov= as needed)
- rxSolve() which solves the system of equations using initial conditions and parameters to make predictions
 - With multiple subject data, this may be parallelized.
 - With single subject the output data frame is adaptive
 - Covariances and other metrics of uncertainty can be used to simulate while solving.

While this is the user guide, there are other places that you can visit for help:

- RxODE github pkgdown page
- RxODE tutorial (accessible in tutorials in Rstudio 1.3+)
- RxODE github discussions

This book was assembled on Tue Mar 9 20:57:10 2021 with RxODE version 1.0.5 automatically by github actions.

Authors and Acknowledgments

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- Drew Schmidt Drew Schmidt author of edits for exponential matrix utility taken from R package expm
- Arun Srinivasan forder secondary author (version modified by Matthew Fidler to allow different type of threading, indexing and exclude grouping)

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- Sherwin Sy Weight based dosing example
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- Ross Ihaka R author
- Robert Gentleman R author
- R core team R authors

Related R packages

3.1 ODE solving

This is a brief comparison of pharmacometric ODE solving R packages to RxODE.

There are several R packages for differential equations. The most popular is deSolve.

However for pharmacometrics-specific ODE solving, there are only 2 packages other than RxODE released on CRAN. Each uses compiled code to have faster ODE solving.

- mrgsolve, which uses C++ lsoda solver to solve ODE systems. The user is required to write hybrid R/C++ code to create a mrgsolve model which is translated to C++ for solving.
 - In contrast, RxODE has a R-like mini-language that is parsed into C code that solves the ODE system.
 - Unlike RxODE, mrgsolve does not currently support symbolic manipulation of ODE systems, like automatic Jacobian calculation or forward sensitivity calculation (RxODE currently supports this and this is the basis of nlmixr's FOCEi algorithm)
- dMod, which uses a unique syntax to create "reactions". These reactions create the underlying ODEs and then created c code for a compiled deSolve model.
 - In contrast RxODE defines ODE systems at a lower level. RxODE's parsing of the mini-language comes from C, whereas dMod's parsing comes from R.
 - Like RxODE, dMod supports symbolic manipulation of ODE systems and calculates forward sensitivities and adjoint sensitivities of systems.

Unlike RxODE, dMod is not thread-safe since deSolve is not yet thread-safe.

And there is one package that is not released on CRAN:

• PKPDsim which defines models in an R-like syntax and converts the system to compiled code.

Like mrgsolve, PKPDsim does not currently support symbolic manipulation of ODE systems.

PKPDsim is not thread-safe.

The open pharmacometrics open source community is fairly friendly, and the Rx-ODE maintainers has had positive interactions with all of the ODE-solving pharmacometric projects listed.

3.2 PK Solved systems

RxODE supports 1-3 compartment models with gradients (using stan math's auto-differentiation). This currently uses the same equations as PKADVAN to allow time-varying covariates.

RxODE can mix ODEs and solved systems.

3.2.1 The following packages for solved PK systems are on CRAN

- mrgsolve currently has 1-2 compartment (poly-exponential models) models built-in. The solved systems and ODEs cannot currently be mixed.
- pmxTools currently have 1-3 compartment (super-positioning) models built-in. This is a R-only implementation.
- PKPDmodels has a one-compartment model with gradients.

3.2.2 Non-CRAN libraries:

• PKADVAN Provides 1-3 compartment models using non-superpositioning. This allows time-varying covariates.

Installation

You can install the released version of RxODE from CRAN with:

```
install.packages("RxODE")
```

You can install the development version of RxODE with

```
devtools::install_github("nlmixrdevelopment/RxODE")
```

To build models with RxODE, you need a working c compiler. To use parallel threaded solving in RxODE, this c compiler needs to support open-mp.

You can check to see if R has working c compiler you can check with:

```
## install.packages("pkgbuild")
pkgbuild::has_build_tools(debug = TRUE)
```

If you do not have the toolchain, you can set it up as described by the platform information below:

4.0.1 Windows

In windows you may simply use installr to install rtools:

```
install.packages("installr")
library(installr)
install.rtools()
```

Alternatively you can download and install rtools directly.

4.0.2 Mac OSX

To get the most speed you need OpenMP enabled and compile RxODE with that compiler. There are various options and the most up to date discussion about this is likely the data.table installation faq for MacOS. The last thing to keep in mind is that RxODE uses the code very similar to the original lsoda which requires the gfortran compiler to be setup as well as the OpenMP compilers.

If you are going to be using RxODE and nlmixr together and have an older mac computer, I would suggest trying the following:

```
library(symengine)
```

If this crashes your R session then the binary does not work with your Mac machine. To be able to run nlmixr, you will need to compile this package manually. I will proceed assuming you have homebrew installed on your system.

On your system terminal you will need to install the dependencies to compile symengine:

```
brew install cmake gmp mpfr libmpc
```

After installing the dependencies, you need to reinstall symengine:

```
install.packages("symengine", type="source")
library(symengine)
```

4.0.3 Linux

To install on linux make sure you install gcc (with openmp support) and gfortran using your distribution's package manager.

4.1 Development Version

Since the development version of RxODE uses StanHeaders, you will need to make sure your compiler is setup to support C++14, as described in the rstan setup page. For R 4.0, I do not believe this requires modifying the windows toolchain any longer (so it is much easier to setup).

Once the C++ toolchain is setup appropriately, you can install the development version from GitHub with:

```
# install.packages("devtools")
devtools::install_github("nlmixrdevelopment/RxODE")
```

Getting Started

The model equations can be specified through a text string, a model file or an R expression. Both differential and algebraic equations are permitted. Differential equations are specified by d/dt(var_name) =. Each equation can be separated by a semicolon.

To load RxODE package and compile the model:

```
library(RxODE)

#> RxODE 1.0.5 using 4 threads (see ?getRxThreads)

mod1 <-RxODE({
    C2 = centr/V2;
    C3 = peri/V3;
    d/dt(depot) =-KA*depot;
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) = Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})</pre>
```

5.1 Specify ODE parameters and initial conditions

Model parameters can be defined as named vectors. Names of parameters in the vector must be a superset of parameters in the ODE model, and the order of parameters within the vector is not important.

```
theta <-
c(KA=2.94E-01, CL=1.86E+01, V2=4.02E+01, # central
Q=1.05E+01, V3=2.97E+02, # peripheral
Kin=1, Kout=1, EC50=200) # effects
```

Initial conditions (ICs) can be defined through a vector as well. If the elements are not specified, the initial condition for the compartment is assumed to be zero.

```
inits <- c(eff=1);</pre>
```

If you want to specify the initial conditions in the model you can add:

```
eff(0) = 1
```

5.2 Specify Dosing and sampling in RxODE

RxODE provides a simple and very flexible way to specify dosing and sampling through functions that generate an event table. First, an empty event table is generated through the "eventTable()" function:

```
ev <- eventTable(amount.units='mg', time.units='hours')</pre>
```

Next, use the add.dosing() and add.sampling() functions of the EventTable object to specify the dosing (amounts, frequency and/or times, etc.) and observation times at which to sample the state of the system. These functions can be called multiple times to specify more complex dosing or sampling regiments. Here, these functions are used to specify 10mg BID dosing for 5 days, followed by 20mg QD dosing for 5 days:

If you wish you can also do this with the mattigr pipe operator %>%

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The functions get.dosing() and get.sampling() can be used to retrieve information from the event table.

```
head(ev$get.dosing())
                               amt rate ii addl evid ss dur
    id low time high
                         \mathtt{cmt}
             0 NA (default) 10000
                                     0 12
                                                 1 0
#> 1 1 NA
               NA (default) 20000
                                     0 24
                                                 1 0
#> 2 1 NA 120
head(ev$get.sampling())
    id low time high
                      cmt amt rate ii addl evid ss dur
             0 NA (obs)
#> 1 1 NA
                          NA NA NA
                                      NA
                                           O NA NA
#> 2 1 NA
             1
                NA (obs) NA
                              NA NA
                                      NA
                                           O NA
                                                 NA
#> 3 1 NA
             2 NA (obs) NA NA NA
                                      NA
                                           O NA NA
#> 4 1 NA
             3 NA (obs) NA NA NA
                                      NA
                                            O NA NA
             4 NA (obs) NA
#> 5 1 NA
                             NA NA
                                      NA
                                            O NA
                                                 NA
#> 6 1 NA
                NA (obs)
                          NA
                              NA NA
                                      NA
                                            O NA
                                                 NA
```

You may notice that these are similar to NONMEM event tables; If you are more familiar with NONMEM data and events you could use them directly with the event table function et

```
ev <- et(amountUnits="mg", timeUnits="hours") %>%
  et(amt=10000, addl=9,ii=12,cmt="depot") %>%
  et(time=120, amt=2000, addl=4, ii=14, cmt="depot") %>%
  et(0:240) # Add sampling
```

You can see from the above code, you can dose to the compartment named in the RxODE model. This slight deviation from NONMEM can reduce the need for compartment renumbering.

These events can also be combined and expanded (to multi-subject events and complex regimens) with rbind, c, seq, and rep. For more information about creating complex dosing regimens using RxODE see the RxODE events section.

5.3 Solving ODEs

The ODE can now be solved by calling the model object's run or solve function. Simulation results for all variables in the model are stored in the output matrix x.

```
x <- mod1$solve(theta, ev, inits);
knitr::kable(head(x))
```

time	C2	C3	depot	centr	peri	eff
0	0.00000	0.0000000	10000.000	0.000	0.0000	1.000000
1	44.37555	0.9198298	7452.765	1783.897	273.1895	1.084664
2	54.88296	2.6729825	5554.370	2206.295	793.8758	1.180825
3	51.90343	4.4564927	4139.542	2086.518	1323.5783	1.228914
4	44.49738	5.9807076	3085.103	1788.795	1776.2702	1.234610
5	36.48434	7.1774981	2299.255	1466.670	2131.7169	1.214742

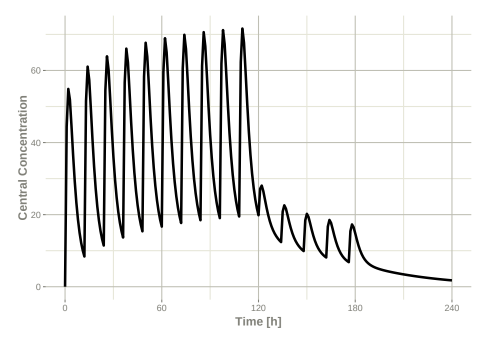
You can also solve this and create a RxODE data frame:

```
x <- mod1 %>% rxSolve(theta, ev, inits);
x
```

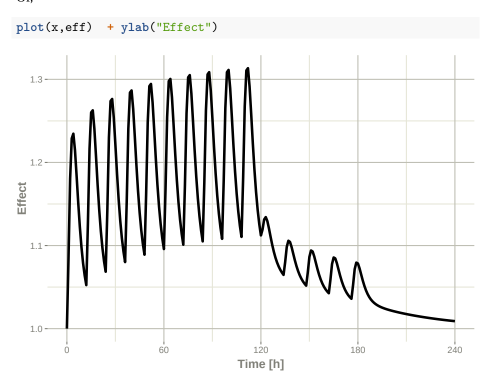
```
#> _____ Solved RxODE object _____
#> -- Parameters (x$params): -----
     V2 V3 KA CL Q
#>
                                 Kin
                                        Kout
#> 40.200 297.000   0.294   18.600   10.500   1.000
                                       1.000 200.000
#> -- Initial Conditions (x$inits): ------
#> depot centr peri eff
#> 0 0 0
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
#> time C2 C3 depot centr peri
#>
     [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
     0 0 0 10000 0
#> 1
                          0 1
#> 2
     1 44.4 0.920 7453. 1784. 273. 1.08
#> 3
     2 54.9 2.67 5554. 2206. 794. 1.18
#> 4
     3 51.9 4.46 4140. 2087. 1324. 1.23
    4 44.5 5.98 3085. 1789. 1776. 1.23
#> 5
#> 6
     5 36.5 7.18 2299. 1467. 2132. 1.21
#> # ... with 235 more rows
```

This returns a modified data frame. You can see the compartment values in the plot below:

```
library(ggplot2)
plot(x,C2) + ylab("Central Concentration")
```



Or,



Note that the labels are automatically labeled with the units from the initial event

table. RxODE extracts units to label the plot (if they are present).

RxODE syntax

This briefly describes the syntax used to define models that RxODE will translate into R-callable compiled code. It also describes the communication of variables between R and the RxODE modeling specification.

6.1 Example

6.2 Syntax

An RxODE model specification consists of one or more statements optionally terminated by semi-colons; and optional comments (comments are delimited by # and an end-of-line).

A block of statements is a set of statements delimited by curly braces, { . . . }.

Statements can be either assignments, conditional if/else if/else, while loops (can be exited by break), special statements, or printing statements (for debugging/testing)

Assignment statements can be:

- **simple** assignments, where the left hand is an identifier (i.e., variable)
- special **time-derivative** assignments, where the left hand specifies the change of the amount in the corresponding state variable (compartment) with respect to time e.g., d/dt(depot):
- special **initial-condition** assignments where the left hand specifies the compartment of the initial condition being specified, e.g. depot (0) = 0
- special model event changes including **bioavailability** (f(depot)=1), **lag time** (alag(depot)=0), **modeled rate** (rate(depot)=2) and **modeled duration** (dur(depot)=2). An example of these model features and the event specification for the modeled infusions the RxODE data specification is found in RxODE events section.
- special **change point syntax, or model times**. These model times are specified by mtime (var)=time
- special **Jacobian-derivative** assignments, where the left hand specifies the change in the compartment ode with respect to a variable. For example, if d/dt(y) = dy, then a Jacobian for this compartment can be specified as df(y)/dy(dy) = 1. There may be some advantage to obtaining the solution or specifying the Jacobian for very stiff ODE systems. However, for the few stiff systems we tried with LSODA, this actually slightly slowed down the solving.

Note that assignment can be done by =, <- or \sim .

When assigning with the ~ operator, the **simple assignments** and **time-derivative** assignments will not be output.

Special statements can be:

• Compartment declaration statements, which can change the default dosing compartment and the assumed compartment number(s) as well as add extra compartment names at the end (useful for multiple-endpoint nlmixr models); These are specified by cmt(compartmentName)

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• Parameter declaration statements, which can make sure the input parameters are in a certain order instead of ordering the parameters by the order they are parsed. This is useful for keeping the parameter order the same when using 2 different ODE models. These are specified by param(par1, par2,...)

An example model is shown below:

```
# simple assignment
C2 = centr/V2;
# time-derivative assignment
d/dt(centr) = F*KA*depot - CL*C2 - Q*C2 + Q*C3;
```

Expressions in assignment and if statements can be numeric or logical, however, no character nor integer expressions are currently supported.

Numeric expressions can include the following numeric operators +, -, *, /, and those mathematical functions defined in the C or the R math libraries (e.g., fabs, exp, log, sin, abs).

You may also access the R's functions in the R math libraries, like lgammafn for the log gamma function.

The RxODE syntax is case-sensitive, i.e., ABC is different than abc, Abc, ABc, etc.

6.2.1 Identifiers

Like R, Identifiers (variable names) may consist of one or more alphanumeric, underscore _ or period . characters, but the first character cannot be a digit or underscore _.

Identifiers in a model specification can refer to:

- State variables in the dynamic system (e.g., compartments in a pharmacokinetics model).
- Implied input variable, t (time), tlast (last time point), and podo (oral dose, in the undocumented case of absorption transit models).
- Special constants like pi or R's predefined constants.
- Model parameters (e.g., ka rate of absorption, CL clearance, etc.)
- Others, as created by assignments as part of the model specification; these are referred as *LHS* (left-hand side) variable.

Currently, the RxODE modeling language only recognizes system state variables and "parameters", thus, any values that need to be passed from R to the ODE model (e.g., age) should be either passed in the params argument of the integrator function rxSolve() or be in the supplied event data-set.

There are certain variable names that are in the RxODE event tables. To avoid confusion, the following event table-related items cannot be assigned, or used as a state but can be accessed in the RxODE code:

- cmt
- dvid
- addl
- ss
- rate
- id

However the following variables are cannot be used in a model specification:

- evid
- ii

Sometimes RxODE generates variables that are fed back to RxODE. Similarly, nlmixr generates some variables that are used in nlmixr estimation and simulation. These variables start with the either the rx or nlmixr prefixes. To avoid any problems, it is suggested to not use these variables starting with either the rx or nlmixr prefixes.

6.3 Logical Operators

Logical operators support the standard R operators ==, !=>= <= > and <. Like R these can be in if() or while() statements, ifelse() expressions. Additionally they can be in a standard assignment. For instance, the following is valid:

```
cov1 = covm*(sexf == "female") + covm*(sexf != "female")
```

Notice that you can also use character expressions in comparisons. This convenience comes at a cost since character comparisons are slower than numeric expressions. Unlike R, as.numeric or as.integer for these logical statements is not only not needed, but will cause an syntax error if you try to use the function.

6.4 cmt() changing compartment numbers for states

The compartment order can be changed with the $\mathtt{cmt}()$ syntax in the model. To understand what the $\mathtt{cmt}()$ can do you need to understand how RxODE numbers the compartments.

Below is an example of how RxODE numbers compartments

6.4.1 How RxODE numbers compartments

RXODE automatically assigns compartment numbers when parsing. For example, with the Mavoglurant PBPK model the following model may be used:

```
library(RxODE)
pbpk <- RxODE({</pre>
    KbBR = exp(1KbBR)
    KbMU = exp(1KbMU)
    KbAD = exp(1KbAD)
    CLint= exp(lCLint + eta.LClint)
   KbBO = exp(1KbBO)
   KbRB = exp(1KbRB)
    ## Regional blood flows
    # Cardiac output (L/h) from White et al (1968)
    CO = (187.00*WT^0.81)*60/1000;
    QHT = 4.0 *CO/100;
    QBR = 12.0*C0/100;
    QMU = 17.0*C0/100;
    QAD = 5.0 *CO/100;
    QSK = 5.0 *CO/100;
    QSP = 3.0 *CO/100;
    QPA = 1.0 *CO/100;
    QLI = 25.5*CO/100;
    QST = 1.0 *CO/100;
    QGU = 14.0*CO/100;
    # Hepatic artery blood flow
    QHA = QLI - (QSP + QPA + QST + QGU);
    QBO = 5.0 *CO/100;
    QKI = 19.0*CO/100;
    QRB = CO - (QHT + QBR + QMU + QAD + QSK + QLI + QBO + QKI);
    QLU = QHT + QBR + QMU + QAD + QSK + QLI + QBO + QKI + QRB;
    ## Organs' volumes = organs' weights / organs' density
   VLU = (0.76 *WT/100)/1.051;
   VHT = (0.47 *WT/100)/1.030;
    VBR = (2.00 *WT/100)/1.036;
   VMU = (40.00*WT/100)/1.041;
   VAD = (21.42*WT/100)/0.916;
   VSK = (3.71 *WT/100)/1.116;
    VSP = (0.26 *WT/100)/1.054;
   VPA = (0.14 *WT/100)/1.045;
    VLI = (2.57 *WT/100)/1.040;
   VST = (0.21 *WT/100)/1.050;
```

```
VGU = (1.44 *WT/100)/1.043;
VBO = (14.29*WT/100)/1.990;
VKI = (0.44 *WT/100)/1.050;
VAB = (2.81 *WT/100)/1.040;
VVB = (5.62 *WT/100)/1.040;
VRB = (3.86 *WT/100)/1.040;
## Fixed parameters
BP = 0.61; # Blood:plasma partition coefficient
fup = 0.028;  # Fraction unbound in plasma
fub = fup/BP; # Fraction unbound in blood
KbLU = exp(0.8334);
KbHT = exp(1.1205);
KbSK = exp(-.5238);
KbSP = exp(0.3224);
KbPA = exp(0.3224);
KbLI = exp(1.7604);
KbST = exp(0.3224);
KbGU = exp(1.2026);
KbKI = exp(1.3171);
##-----
S15 = VVB*BP/1000;
C15 = Venous Blood/S15
##-----
d/dt(Lungs) = QLU*(Venous Blood/VVB - Lungs/KbLU/VLU);
d/dt(Heart) = QHT*(Arterial_Blood/VAB - Heart/KbHT/VHT);
d/dt(Brain) = QBR*(Arterial_Blood/VAB - Brain/KbBR/VBR);
d/dt(Muscles) = QMU*(Arterial_Blood/VAB - Muscles/KbMU/VMU);
d/dt(Adipose) = QAD*(Arterial_Blood/VAB - Adipose/KbAD/VAD);
d/dt(Skin) = QSK*(Arterial_Blood/VAB - Skin/KbSK/VSK);
d/dt(Spleen) = QSP*(Arterial_Blood/VAB - Spleen/KbSP/VSP);
d/dt(Pancreas) = QPA*(Arterial_Blood/VAB - Pancreas/KbPA/VPA);
d/dt(Liver) = QHA*Arterial_Blood/VAB + QSP*Spleen/KbSP/VSP +
  QPA*Pancreas/KbPA/VPA + QST*Stomach/KbST/VST +
  QGU*Gut/KbGU/VGU - CLint*fub*Liver/KbLI/VLI - QLI*Liver/KbLI/VLI;
d/dt(Stomach) = QST*(Arterial_Blood/VAB - Stomach/KbST/VST);
d/dt(Gut) = QGU*(Arterial Blood/VAB - Gut/KbGU/VGU);
d/dt(Bones) = QBO*(Arterial_Blood/VAB - Bones/KbBO/VBO);
d/dt(Kidneys) = QKI*(Arterial Blood/VAB - Kidneys/KbKI/VKI);
d/dt(Arterial_Blood) = QLU*(Lungs/KbLU/VLU - Arterial_Blood/VAB);
d/dt(Venous Blood) = QHT*Heart/KbHT/VHT + QBR*Brain/KbBR/VBR +
  QMU*Muscles/KbMU/VMU + QAD*Adipose/KbAD/VAD + QSK*Skin/KbSK/VSK +
```

```
QLI*Liver/KbLI/VLI + QBO*Bones/KbBO/VBO + QKI*Kidneys/KbKI/VKI +
QRB*Rest_of_Body/KbRB/VRB - QLU*Venous_Blood/VVB;
d/dt(Rest_of_Body) = QRB*(Arterial_Blood/VAB - Rest_of_Body/KbRB/VRB);
})
```

If you look at the summary, you can see where RxODE assigned the compartment number(s)

```
summary(pbpk)
```

```
#> RxODE 1.0.5 model named rx_74372d99e4c72628e9dee8939b90cb49 model (ready).
#> DLL: /home/matt/.cache/R/RxODE/rx_74372d99e4c72628e9dee8939b90cb49__.rxd/rx_74372d99e4c72628e9
#> NULL
#>
#> Calculated Variables:
#> [1] "KbBR" "KbMU" "KbAD"
                               "CLint" "KbBO"
                                               "KbRB"
                                                       "CO"
                                                               "QHT"
                                                                       "QBR"
#> [10] "QMU"
               "QAD"
                       "QSK"
                               "QSP"
                                       "QPA"
                                               "QLI"
                                                       "QST"
                                                               "QGU"
                                                                       "QHA"
                                                "VHT"
#> [19] "QBO"
               "QKI"
                       "QRB"
                               "QLU"
                                       "VLU"
                                                       "VBR"
                                                               "VMU"
                                                                       "VAD"
#> [28] "VSK"
               "VSP"
                       "VPA"
                                       "VST"
                                                "VGU"
                                                       "VBO"
                                                               "VKI"
                                                                       "VAB"
                               "VLI"
               "VRB"
#> [37] "VVB"
                       "fub"
                               "KbLU"
                                       "KbHT"
                                               "KbSK" "KbSP"
                                                               "KbPA" "KbLI"
#> [46] "KbST" "KbGU" "KbKI" "S15"
                                       "C15"
#> _____ RxODE Model Syntax _____
#> RxODE({
#>
      KbBR = exp(1KbBR)
#>
      KbMU = exp(1KbMU)
#>
      KbAD = exp(1KbAD)
#>
      CLint = exp(lCLint + eta.LClint)
#>
      KbBO = exp(1KbBO)
#>
      KbRB = exp(1KbRB)
#>
      CO = (187 * WT^{0.81}) * 60/1000
#>
      QHT = 4 * CO/100
#>
      QBR = 12 * CO/100
#>
      QMU = 17 * CO/100
      QAD = 5 * CO/100
#>
#>
      QSK = 5 * CO/100
#>
      QSP = 3 * CO/100
#>
      QPA = 1 * CO/100
#>
      QLI = 25.5 * CO/100
#>
      QST = 1 * CO/100
#>
      QGU = 14 * CO/100
#>
      QHA = QLI - (QSP + QPA + QST + QGU)
#>
      QBO = 5 * CO/100
#>
      QKI = 19 * CO/100
      QRB = CO - (QHT + QBR + QMU + QAD + QSK + QLI + QBO + QKI)
#>
```

```
#>
       QLU = QHT + QBR + QMU + QAD + QSK + QLI + QBO + QKI + QRB
#>
       VLU = (0.76 * WT/100)/1.051
#>
       VHT = (0.47 * WT/100)/1.03
#>
       VBR = (2 * WT/100)/1.036
#>
       VMU = (40 * WT/100)/1.041
#>
       VAD = (21.42 * WT/100)/0.916
       VSK = (3.71 * WT/100)/1.116
#>
#>
       VSP = (0.26 * WT/100)/1.054
#>
       VPA = (0.14 * WT/100)/1.045
#>
       VLI = (2.57 * WT/100)/1.04
#>
       VST = (0.21 * WT/100)/1.05
#>
       VGU = (1.44 * WT/100)/1.043
#>
       VBO = (14.29 * WT/100)/1.99
#>
       VKI = (0.44 * WT/100)/1.05
#>
       VAB = (2.81 * WT/100)/1.04
       VVB = (5.62 * WT/100)/1.04
#>
#>
       VRB = (3.86 * WT/100)/1.04
#>
       BP = 0.61
#>
       fup = 0.028
#>
       fub = fup/BP
#>
       KbLU = exp(0.8334)
#>
       KbHT = exp(1.1205)
#>
       KbSK = exp(-0.5238)
#>
       KbSP = exp(0.3224)
#>
       KbPA = exp(0.3224)
#>
       KbLI = exp(1.7604)
#>
       KbST = exp(0.3224)
#>
       KbGU = exp(1.2026)
#>
       KbKI = exp(1.3171)
#>
       S15 = VVB * BP/1000
#>
       C15 = Venous_Blood/S15
#>
       d/dt(Lungs) = QLU * (Venous_Blood/VVB - Lungs/KbLU/VLU)
#>
       d/dt(Heart) = QHT * (Arterial_Blood/VAB - Heart/KbHT/VHT)
#>
       d/dt(Brain) = QBR * (Arterial_Blood/VAB - Brain/KbBR/VBR)
#>
       d/dt(Muscles) = QMU * (Arterial_Blood/VAB - Muscles/KbMU/VMU)
#>
       d/dt(Adipose) = QAD * (Arterial_Blood/VAB - Adipose/KbAD/VAD)
#>
       d/dt(Skin) = QSK * (Arterial_Blood/VAB - Skin/KbSK/VSK)
#>
       d/dt(Spleen) = QSP * (Arterial_Blood/VAB - Spleen/KbSP/VSP)
#>
       d/dt(Pancreas) = QPA * (Arterial_Blood/VAB - Pancreas/KbPA/VPA)
#>
       d/dt(Liver) = QHA * Arterial_Blood/VAB + QSP * Spleen/KbSP/VSP +
#>
           QPA * Pancreas/KbPA/VPA + QST * Stomach/KbST/VST + QGU *
#>
           Gut/KbGU/VGU - CLint * fub * Liver/KbLI/VLI - QLI * Liver/KbLI/VLI
#>
       d/dt(Stomach) = QST * (Arterial_Blood/VAB - Stomach/KbST/VST)
#>
       d/dt(Gut) = QGU * (Arterial_Blood/VAB - Gut/KbGU/VGU)
#>
       d/dt(Bones) = QBO * (Arterial Blood/VAB - Bones/KbBO/VBO)
#>
       d/dt(Kidneys) = QKI * (Arterial_Blood/VAB - Kidneys/KbKI/VKI)
```

In this case, Venous_Blood is assigned to compartment 15. Figuring this out can be inconvenient and also lead to re-numbering compartment in simulation or estimation datasets. While it is easy and probably clearer to specify the compartment by name, other tools only support compartment numbers. Therefore, having a way to number compartment easily can lead to less data modification between multiple tools.

6.4.2 Changing compartments by pre-declaring with cmt()

To add the compartments to the RxODE model in the order you desire you simply need to pre-declare the compartments with cmt. For example specifying is Venous_Blood and Skin to be the 1st and 2nd compartments, respectively, is simple:

```
pbpk2 <- RxODE({</pre>
  ## Now this is the first compartment, ie cmt=1
  cmt(Venous_Blood)
  ## Skin may be a compartment you wish to dose to as well,
  ## so it is now cmt=2
  cmt(Skin)
  KbBR = exp(1KbBR)
  KbMU = exp(1KbMU)
  KbAD = exp(1KbAD)
  CLint= exp(lCLint + eta.LClint)
  KbBO = exp(1KbBO)
  KbRB = exp(1KbRB)
  ## Regional blood flows
  # Cardiac output (L/h) from White et al (1968)m
  CO = (187.00*WT^0.81)*60/1000;
  QHT = 4.0 *CO/100;
  QBR = 12.0*C0/100;
  QMU = 17.0*CO/100;
  QAD = 5.0 *CO/100;
```

```
QSK = 5.0 *CO/100;
QSP = 3.0 *CO/100;
QPA = 1.0 *CO/100;
QLI = 25.5*CO/100;
QST = 1.0 *CO/100;
QGU = 14.0*CO/100;
QHA = QLI - (QSP + QPA + QST + QGU); # Hepatic artery blood flow
QBO = 5.0 *CO/100;
QKI = 19.0*CO/100;
QRB = CO - (QHT + QBR + QMU + QAD + QSK + QLI + QBO + QKI);
QLU = QHT + QBR + QMU + QAD + QSK + QLI + QBO + QKI + QRB;
## Organs' volumes = organs' weights / organs' density
VLU = (0.76 *WT/100)/1.051;
VHT = (0.47 *WT/100)/1.030;
VBR = (2.00 *WT/100)/1.036;
VMU = (40.00*WT/100)/1.041;
VAD = (21.42*WT/100)/0.916;
VSK = (3.71 *WT/100)/1.116;
VSP = (0.26 *WT/100)/1.054;
VPA = (0.14 *WT/100)/1.045;
VLI = (2.57 *WT/100)/1.040;
VST = (0.21 *WT/100)/1.050;
VGU = (1.44 *WT/100)/1.043;
VBO = (14.29*WT/100)/1.990;
VKI = (0.44 *WT/100)/1.050;
VAB = (2.81 *WT/100)/1.040;
VVB = (5.62 *WT/100)/1.040;
VRB = (3.86 *WT/100)/1.040;
## Fixed parameters
BP = 0.61; # Blood:plasma partition coefficient
fup = 0.028;
              # Fraction unbound in plasma
fub = fup/BP; # Fraction unbound in blood
KbLU = \exp(0.8334);
KbHT = exp(1.1205);
KbSK = exp(-.5238);
KbSP = exp(0.3224);
KbPA = exp(0.3224);
KbLI = exp(1.7604);
KbST = \exp(0.3224);
KbGU = \exp(1.2026);
KbKI = exp(1.3171);
```

```
S15 = VVB*BP/1000;
C15 = Venous_Blood/S15
d/dt(Lungs) = QLU*(Venous Blood/VVB - Lungs/KbLU/VLU);
d/dt(Heart) = QHT*(Arterial_Blood/VAB - Heart/KbHT/VHT);
d/dt(Brain) = QBR*(Arterial_Blood/VAB - Brain/KbBR/VBR);
d/dt(Muscles) = QMU*(Arterial_Blood/VAB - Muscles/KbMU/VMU);
d/dt(Adipose) = QAD*(Arterial_Blood/VAB - Adipose/KbAD/VAD);
d/dt(Skin) = QSK*(Arterial_Blood/VAB - Skin/KbSK/VSK);
d/dt(Spleen) = QSP*(Arterial_Blood/VAB - Spleen/KbSP/VSP);
d/dt(Pancreas) = QPA*(Arterial_Blood/VAB - Pancreas/KbPA/VPA);
d/dt(Liver) = QHA*Arterial_Blood/VAB + QSP*Spleen/KbSP/VSP +
  QPA*Pancreas/KbPA/VPA + QST*Stomach/KbST/VST + QGU*Gut/KbGU/VGU -
  CLint*fub*Liver/KbLI/VLI - QLI*Liver/KbLI/VLI;
d/dt(Stomach) = QST*(Arterial_Blood/VAB - Stomach/KbST/VST);
d/dt(Gut) = QGU*(Arterial_Blood/VAB - Gut/KbGU/VGU);
d/dt(Bones) = QBO*(Arterial_Blood/VAB - Bones/KbBO/VBO);
d/dt(Kidneys) = QKI*(Arterial_Blood/VAB - Kidneys/KbKI/VKI);
d/dt(Arterial_Blood) = QLU*(Lungs/KbLU/VLU - Arterial_Blood/VAB);
d/dt(Venous_Blood) = QHT*Heart/KbHT/VHT + QBR*Brain/KbBR/VBR +
  QMU*Muscles/KbMU/VMU + QAD*Adipose/KbAD/VAD + QSK*Skin/KbSK/VSK +
  QLI*Liver/KbLI/VLI + QBO*Bones/KbBO/VBO + QKI*Kidneys/KbKI/VKI +
  QRB*Rest_of_Body/KbRB/VRB - QLU*Venous_Blood/VVB;
d/dt(Rest_of_Body) = QRB*(Arterial_Blood/VAB - Rest_of_Body/KbRB/VRB);
```

You can see this change in the simple printout

```
pbpk2
```

```
#> RxODE 1.0.5 model named rx_a65bdb529b0485f601cec6187b5faaf5 model (ready).
#> x$state: Venous_Blood, Skin, Lungs, Heart, Brain, Muscles, Adipose, Spleen, Pancreas, Liver, Sparams: 1KbBR, 1KbMU, 1KbAD, 1CLint, eta.LClint, 1KbBO, 1KbRB, WT, BP, fup
#> x$lhs: KbBR, KbMU, KbAD, CLint, KbBO, KbRB, CO, QHT, QBR, QMU, QAD, QSK, QSP, QPA, QLI, QST, Q
```

The first two compartments are Venous_Blood followed by Skin.

6.4.3 Appending compartments to the model with cmt()

You can also append "compartments" to the model. Because of the ODE solving internals, you cannot add fake compartments to the model until after all the differential equations are defined.

For example this is legal:

```
ode.1c.ka <- RxODE({
    C2 = center/V;
    d / dt(depot) = -KA * depot
    d/dt(center) = KA * depot - CL*C2
    cmt(eff);
})
print(ode.1c.ka)
#> RxODE 1.0.5 model named rx_47c1eb3facce5268a288e5652999299e model (ready).
#> $state: depot, center
#> $stateExtra: eff
#> $params: V, KA, CL
#> $1hs: C2
But compartments defined before all the differential equations is not supported;
So the model below:
ode.1c.ka <- RxODE({
    cmt(eff);
    C2 = center/V;
    d / dt(depot) = -KA * depot
    d/dt(center) = KA * depot - CL*C2
})
will give an error:
Error in rxModelVars_(obj) :
  Evaluation error: Compartment 'eff' needs differential equations defined.
```

RxODE events

7.1 RxODE event tables

In general, RxODE event tables follow NONMEM dataset convention with the exceptions:

- The compartment data item (cmt) can be a string/factor with compartment names
 - You may turn off a compartment with a negative compartment number or "-cmt" where cmt is the compartment name.
 - The compartment data item (cmt) can still be a number, the number of the compartment is defined by the appearance of the compartment name in the model. This can be tedious to count, so you can specify compartment numbers easier by using the cmt(cmtName) at the beginning of the model.
- An additional column, dur can specify the duration of infusions;
 - Bioavailability changes will change the rate of infusion since dur/amt are fixed in the input data.
 - Similarly, when specifying rate/amt for an infusion, the bioavailability will change the infusion duration since rate/amt are fixed in the input data.
- Some infrequent NONMEM columns are not supported: pcmt, call.
- Additional events are supported:
 - evid=5 or replace event; This replaces the value of a compartment with the value specified in the amt column. This is equivalent to deSolve=replace.

evid=6 or multiply event; This multiplies the value in the compartment with the value specified by the amt column. This is equivalent to deSolve=multiply.

Here are the legal entries to a data table:

Data		
Item	Meaning	Notes
id	Individual identifier	Can be a integer, factor, character, or numeric
time	Individual time	Numeric for each time.
amt	dose amount	Positive for doses zero/NA for observations
rate	infusion rate	When specified the infusion duration will be dur=amt/rate
		rate = -1, rate modeled; rate = -2, duration modeled
dur	infusion	When specified the infusion rate will be rate =
	duration	amt/dur
evid	event ID	0=Observation; 1=Dose; 2=Other; 3=Reset;
		4=Reset+Dose; 5=Replace; 6=Multiply
cmt	Compartment	Represents compartment #/name for
	-	dose/observation
SS	Steady State	0 = non-steady-state; 1=steady state; 2=steady state
	Flag	+prior states
ii	Inter-dose	Time between doses.
	Interval	
addl	# of additional doses	Number of doses like the current dose.

Other notes:

- The evid can be the classic RxODE (described here) or the NONMEM-style evid described above.
- NONMEM's DV is not required; RxODE is a ODE solving framework.
- NONMEM's MDV is not required, since it is captured in EVID.
- Instead of NONMEM-compatible data, it can accept deSolve compatible data-frames.

When returning the RxODE solved data-set there are a few additional event ids (EVID) that you may see depending on the solving options:

- EVID = -1 is when a modeled rate ends (corresponds to rate = -1)
- EVID = -2 is when a modeled duration ends (corresponds to rate=-2)
- EVID = -10 when a rate specified zero-order infusion ends (corresponds to rate > 0)

- EVID = -20 when a duration specified zero-order infusion ends (corresponds to dur > 0)
- EVID = 101, 102, 103,... These correspond to the 1, 2, 3,... modeled time (mtime).

These can only be accessed when solving with the option combination addDosing=TRUE and subsetNonmem=FALSE. If you want to see the classic EVID equivalents you can use addDosing=NA.

To illustrate the event types we will use the model from the original RxODE tutorial.

```
library(RxODE)
### Model from RxODE tutorial
m1 <-RxODE({
   KA=2.94E-01;
    CL=1.86E+01;
   V2=4.02E+01;
    Q=1.05E+01;
   V3=2.97E+02;
   Kin=1;
   Kout=1;
   EC50=200;
    ## Added modeled bioavaiblity, duration and rate
   fdepot = 1;
   durDepot = 8;
   rateDepot = 1250;
    C2 = centr/V2;
   C3 = peri/V3;
   d/dt(depot) =-KA*depot;
   f(depot) = fdepot
   dur(depot) = durDepot
   rate(depot) = rateDepot
   d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
   d/dt(peri) =
                                     Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
    eff(0) = 1
});
```

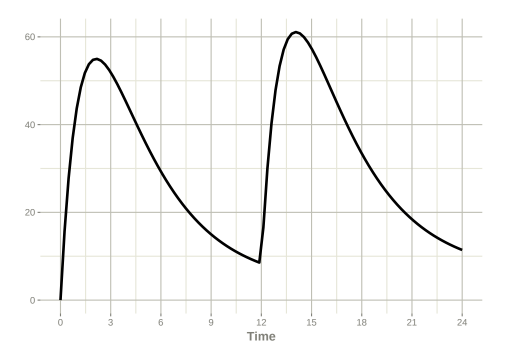
7.2 Bolus/Additive Doses

A bolus dose is the default type of dose in RxODE and only requires the amt/dose. Note that this uses the convenience function et() described in the RxODE event tables

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12,until=24) %>%
    et(seq(0, 24, length.out=100))
```

```
#> ------ EventTable with 101 records ------
#>
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
     100 observation times (see x$get.sampling(); add with add.sampling or et)
#>
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: ------
#> # A tibble: 101 x 5
#>
         time amt
                    ii addl evid
          [h] <dbl> [h] <int> <evid>
#>
#> 1 0.0000000 NA NA NA 0:Observation
#> 2 0.0000000 10000 12
                          2 1:Dose (Add)
#> 3 0.2424242 NA NA NA 0:Observation
#> 4 0.4848485 NA NA NA 0:Observation
#> 5 0.7272727 NA NA NA 0:Observation
#> 6 0.9696970 NA NA NA 0:Observation
#> 7 1.2121212 NA NA NA 0:Observation
#> 8 1.4545455 NA NA NA 0:Observation
#> 9 1.6969697
                    NA NA 0:Observation
               NA
#> 10 1.9393939
              NA
                   NA NA 0:Observation
#> # ... with 91 more rows
```

```
rxSolve(m1, ev) %>% plot(C2) +
    xlab("Time")
```



7.3 Infusion Doses

There are a few different type of infusions that RxODE supports:

- Constant Rate Infusion (rate)
- Constant Duration Infusion (dur)
- Estimated Rate of Infusion
- Estimated Duration of Infusion

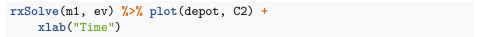
7.3.1 Constant Infusion (in terms of duration and rate)

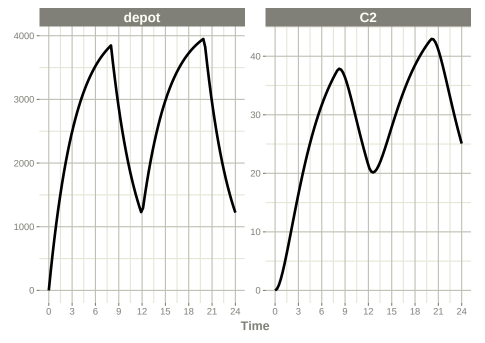
The next type of event is an infusion; There are two ways to specify an infusion; The first is the dur keyword.

An example of this is:

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12,until=24, dur=8) %>%
    et(seq(0, 24, length.out=100))
```

```
----- EventTable with 101 records ------
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
     100 observation times (see x$get.sampling(); add with add.sampling or et)
#>
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#>
#> -- First part of x: ------
#> # A tibble: 101 x 6
#>
          time
                 amt
                       ii addl evid
                                                dur
           [h] <dbl>
                       [h] <int> <evid>
                                                [h]
#>
#>
   1 0.0000000
                  NA
                       NA
                             NA 0:Observation
                                                NA
   2 0.0000000 10000
                             2 1:Dose (Add)
                                                 8
#>
   3 0.2424242
                       NA
                             NA 0:Observation
                                                NA
                             NA 0:Observation
   4 0.4848485
                       NA
   5 0.7272727
                  NA
                       NA
                             NA 0:Observation
#>
                                                NA
   6 0.9696970
                 NA
                       NA
                             NA 0:Observation
   7 1.2121212
                             NA 0:Observation
#>
                  NA
                       NA
                                                NA
   8 1.4545455
                  NA
                       NA
                             NA 0:Observation
                                                NA
  9 1.6969697
                  NA
                       NA
                             NA 0:Observation
                                                NA
#> 10 1.9393939
                  NA
                             NA 0:Observation
                       NA
                                                NA
#> # ... with 91 more rows
```



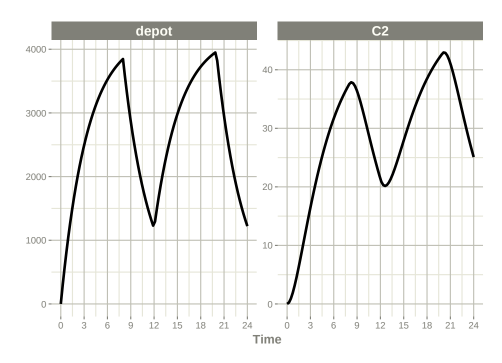


It can be also specified by the rate component:

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12,until=24, rate=10000/8) %>%
    et(seq(0, 24, length.out=100))
ev
```

```
#> ----- EventTable with 101 records -----
#>
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
     100 observation times (see x$get.sampling(); add with add.sampling or et)
#>
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: -----
#> # A tibble: 101 x 6
#>
         time amt rate
                             ii addl evid
          [h] <dbl> <rate/dur> [h] <int> <evid>
#>
#> 1 0.000000 NA NA
                             NA NA 0:Observation
#> 2 0.0000000 10000 1250
                             12
                                   2 1:Dose (Add)
#> 3 0.2424242 NA NA
                             NA
                                   NA 0:Observation
#> 4 0.4848485 NA NA
                             NA NA 0:Observation
#> 5 0.7272727 NA NA
                              NA
                                   NA 0:Observation
#> 6 0.9696970 NA NA
                             NA
                                   NA 0:Observation
#> 7 1.2121212 NA NA
                             NA NA 0:Observation
#> 8 1.4545455 NA NA
                             NA NA 0:Observation
#> 9 1.6969697
              NA NA
                              NA
                                   NA 0:Observation
#> 10 1.9393939
              NA NA
                              NA NA 0:Observation
#> # ... with 91 more rows
```

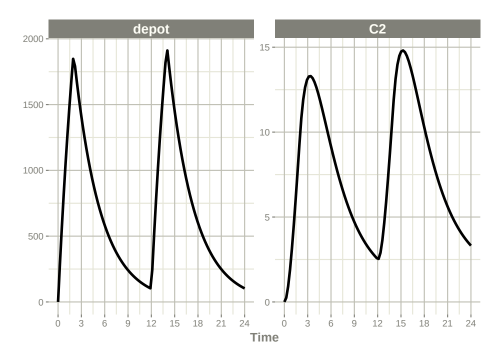
```
rxSolve(m1, ev) %>% plot(depot, C2) +
    xlab("Time")
```



These are the same with the exception of how bioavailability changes the infusion.

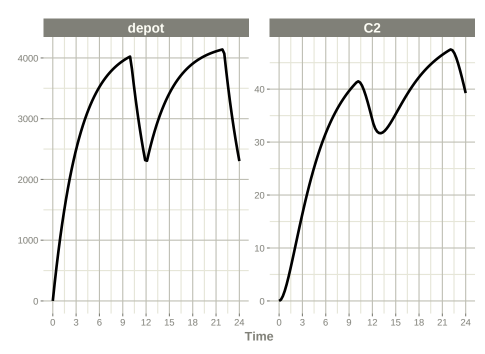
In the case of modeling rate, a bioavailability decrease, decreases the infusion duration, as in NONMEM. For example:

```
rxSolve(m1, ev, c(fdepot=0.25)) %>% plot(depot, C2) +
    xlab("Time")
```



Similarly increasing the bioavailability increases the infusion duration.

```
rxSolve(m1, ev, c(fdepot=1.25)) %>% plot(depot, C2) +
    xlab("Time")
```

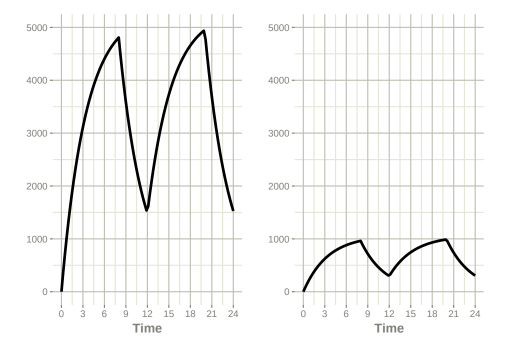


The rationale for this behavior is that the rate and amt are specified by the event table, so the only thing that can change with a bioavailability increase is the duration of the infusion.

If you specify the amt and dur components in the event table, bioavailability changes affect the rate of infusion.

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12,until=24, dur=8) %>%
    et(seq(0, 24, length.out=100))
```

You can see the side-by-side comparison of bioavailability changes affecting rate instead of duration with these records in the following plots:



7.3.2 Modeled Rate and Duration of Infusion

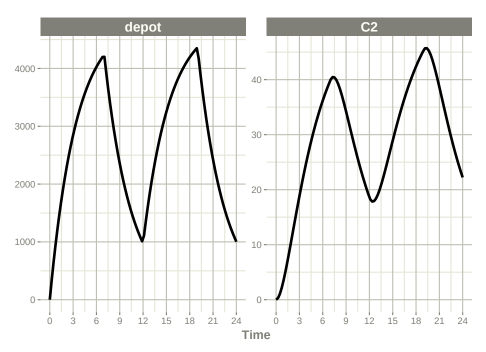
You can model the duration, which is equivalent to NONMEM's rate=-2. As a mnemonic you can use the dur=model instead of rate=-2

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12,until=24, dur=model) %>%
    et(seq(0, 24, length.out=100))
```

```
----- EventTable with 101 records -----
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
     100 observation times (see x$get.sampling(); add with add.sampling or et)
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#>
#> -- First part of x: ------
#> # A tibble: 101 x 6
                              ii addl evid
#>
         time
              amt rate
          [h] <dbl> <rate/dur>
                              [h] <int> <evid>
#>
#> 1 0.0000000
                NA NA
                             NA NA 0:Observation
#> 2 0.0000000 10000 -2:dur
                              12
                                    2 1:Dose (Add)
#> 3 0.2424242 NA NA
                               NA
                                    NA 0:Observation
```

```
#>
   4 0.4848485
                   NA NA
                                    NA
                                          NA 0:Observation
#>
   5 0.7272727
                  NA NA
                                    NA
                                          NA 0:Observation
                  NA NA
                                          NA 0:Observation
#> 6 0.9696970
                                    NA
#> 7 1.2121212
                  NA NA
                                          NA 0:Observation
                                   NA
#> 8 1.4545455
                  NA NA
                                          NA 0:Observation
                                   NA
#> 9 1.6969697
                  NA NA
                                   NA
                                          NA 0:Observation
#> 10 1.9393939
                  NA NA
                                   NA
                                          NA 0:Observation
#> # ... with 91 more rows
```

```
rxSolve(m1, ev, c(durDepot=7)) %>% plot(depot, C2) +
    xlab("Time")
```

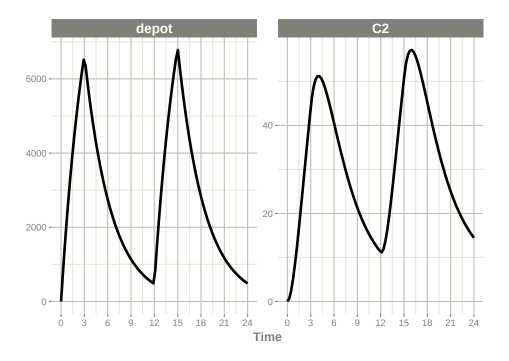


Similarly, you may also model rate. This is equivalent to NONMEM's rate=-1 and is how RxODE's event table specifies the data item as well. You can also use rate=model as a mnemonic:

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12,until=24, rate=model) %>%
    et(seq(0, 24, length.out=100))
```

```
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
     100 observation times (see x$get.sampling(); add with add.sampling or et)
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: -----
#> # A tibble: 101 x 6
                                ii addl evid
#>
          time
                amt rate
#>
           [h] <dbl> <rate/dur>
                                [h] <int> <evid>
   1 0.0000000
                                      NA 0:Observation
                 NA NA
                               NA
   2 0.0000000 10000 -1:rate
                              12
                                       2 1:Dose (Add)
#>
#>
  3 0.2424242
                 NA NA
                                NA
                                      NA 0:Observation
  4 0.4848485
                 NA NA
                                NA
                                      NA 0:Observation
#> 5 0.7272727
                 NA NA
                                NA
                                      NA 0:Observation
  6 0.9696970
                 NA NA
                                 NA
                                      NA 0:Observation
  7 1.2121212
               NA NA
                                 NA
                                      NA 0:Observation
#> 8 1.4545455
                 NA NA
                                 NA
                                      NA 0:Observation
#> 9 1.6969697
               NA NA
                                NA
                                      NA 0:Observation
#> 10 1.9393939
                 NA NA
                                 NA
                                      NA 0:Observation
#> # ... with 91 more rows
```

```
rxSolve(m1, ev, c(rateDepot=10000/3)) %>% plot(depot, C2) +
    xlab("Time")
```



7.4 Steady State

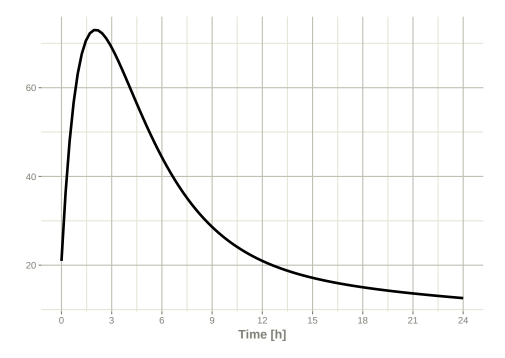
These doses are solved until a steady state is reached with a constant inter-dose interval.

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12, ss=1) %>%
    et(seq(0, 24, length.out=100))
ev
```

```
#> ----- EventTable with 101 records -----
#>
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
     100 observation times (see x$get.sampling(); add with add.sampling or et)
#> -- First part of x: -----
#> # A tibble: 101 x 5
#>
         time amt
                    ii evid
                                     SS
          [h] <dbl> [h] <evid> <int>
#>
#> 1 0.0000000 NA NA 0:Observation
                                     NA
#> 2 0.0000000 10000 12 1:Dose (Add)
                                      1
#> 3 0.2424242 NA NA 0:Observation
                                     NA
#> 4 0.4848485 NA NA 0:Observation
                                    NA
#> 5 0.7272727 NA NA 0:Observation NA
#> 6 0.9696970 NA NA 0:Observation NA
#> 7 1.2121212 NA NA 0:Observation NA
#> 8 1.4545455 NA NA 0:Observation NA
#> 9 1.6969697 NA NA 0:Observation
                                     NA
             NA
#> 10 1.9393939
                    NA 0:Observation
                                     NA
#> # ... with 91 more rows
```

```
rxSolve(m1, ev) %>% plot(C2)
```

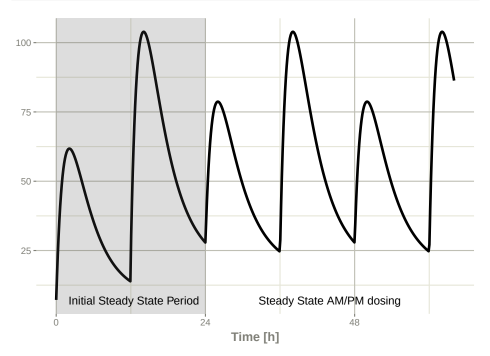
49



7.4.1 Steady state for complex dosing

By using the ss=2 flag, you can use the super-positioning principle in linear kinetics to get steady state nonstandard dosing (i.e. morning 100 mg vs evening 150 mg). This is done by:

- Saving all the state values
- Resetting all the states and solving the system to steady state
- · Adding back all the prior state values



You can see that it takes a full dose cycle to reach the true complex steady state dosing.

7.4.2 Steady state for constant infusion or zero order processes

The last type of steady state that RxODE supports is steady-state constant infusion rate. This can be specified the same way as NONMEM, that is:

- No inter-dose interval ii=0
- A steady state dose, ie ss=1
- Either a positive rate (rate>0) or a estimated rate rate=-1.
- A zero dose, ie amt=0
- Once the steady-state constant infusion is achieved, the infusion is turned off when using this record, just like NONMEM.

Note that rate=-2 where we model the duration of infusion doesn't make much sense since we are solving the infusion until steady state. The duration is specified by the steady state solution.

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Also note that bioavailability changes on this steady state infusion also do not make sense because they neither change the rate or the duration of the steady state infusion. Hence modeled bioavailability on this type of dosing event is ignored.

Here is an example:

```
ev <- et(timeUnits="hr") %>%
    et(amt=0, ss=1,rate=10000/8)

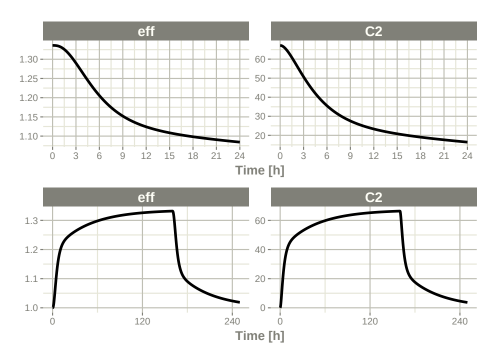
p1 <- rxSolve(m1, ev) %>% plot(C2, eff)

ev <- et(timeUnits="hr") %>%
    et(amt=200000, rate=10000/8) %>%
    et(0, 250, length.out=1000)

p2 <- rxSolve(m1, ev) %>% plot(C2, eff)

library(patchwork)

p1 / p2
```



Not only can this be used for PK, it can be used for steady-state disease processes.

7.5 Reset Events

Reset events are implemented by evid=3 or evid=reset, for reset and evid=4 for reset and dose.

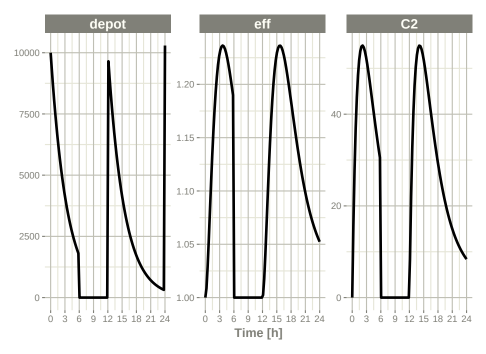
```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12, addl=3) %>%
    et(time=6, evid=reset) %>%
    et(seq(0, 24, length.out=100))
```

```
#> ----- EventTable with 102 records -----
#>
#>
      2 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
      100 observation times (see x$get.sampling(); add with add.sampling or et)
      multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: ------
#> # A tibble: 102 x 5
#>
           time amt ii addl evid
            [h] <dbl> [h] <int> <evid>
#>
#> 1 0.0000000 NA NA NA 0:Observation
#> 2 0.0000000 10000 12 3 1:Dose (Add)
#> 3 0.2424242 NA NA NA 0:Observation
#> 4 0.4848485 NA NA NA 0:Observation
#> 5 0.7272727 NA NA NA 0:Observation
#> 6 0.9696970 NA NA NA 0:Observation
#> 7 1.2121212     NA     NA     O:Observation
#> 8 1.4545455     NA     NA     NA     O:Observation
#> 9 1.6969697     NA     NA     NA     O:Observation
#> 10 1.9393939 NA NA NA 0:Observation
#> # ... with 92 more rows
```

The solving show what happens in this system when the system is reset at 6 hours post-dose.

```
rxSolve(m1, ev) %>% plot(depot,C2, eff)
```

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You can see all the compartments are reset to their initial values. The next dose start the dosing cycle over.

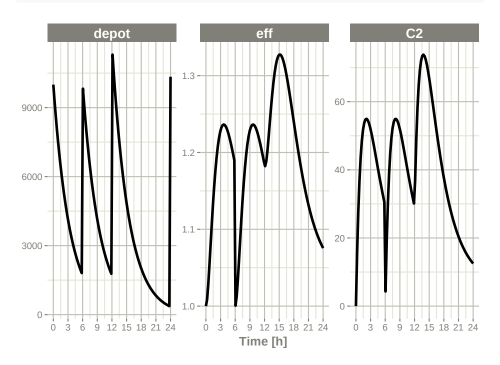
```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12, addl=3) %>%
    et(time=6, amt=10000, evid=4) %>%
    et(seq(0, 24, length.out=100))
```

```
----- EventTable with 102 records ------
#>
     2 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
     100 observation times (see x$get.sampling(); add with add.sampling or et)
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: -----
#> # A tibble: 102 x 5
#>
         time
               amt
                     ii addl evid
          [h] <dbl>
#>
                     [h] <int> <evid>
  1 0.0000000
                          NA 0:Observation
#>
                NA
                     NA
  2 0.0000000 10000
                     12
                           3 1:Dose (Add)
  3 0.2424242
              NA
                     NA
                         NA 0:Observation
              NA NA NA 0:Observation
  4 0.4848485
                   NA NA 0:Observation
#> 5 0.7272727
              NA
```

```
#>
   6 0.9696970
                   NA
                        NA
                              NA 0:Observation
#>
   7 1.2121212
                  NA
                        NA
                              NA 0:Observation
                  NA
#> 8 1.4545455
                        NA
                              NA 0:Observation
#> 9 1.6969697
                  NA
                              NA 0:Observation
                        NA
#> 10 1.9393939
                              NA 0:Observation
                  NA
                        NA
#> # ... with 92 more rows
```

In this case, the whole system is reset and the dose is given

```
rxSolve(m1, ev) %>% plot(depot,C2, eff)
```



7.6 Turning off compartments

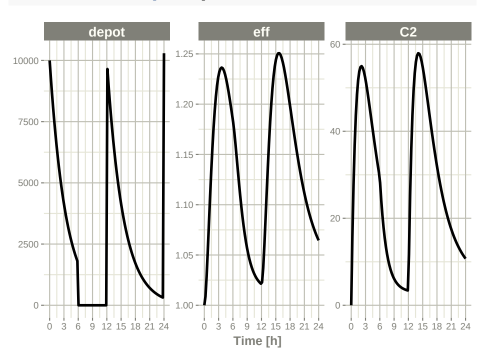
You may also turn off a compartment, which is similar to a reset event.

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12, addl=3) %>%
    et(time=6, cmt="-depot", evid=2) %>%
    et(seq(0, 24, length.out=100))
```

```
----- EventTable with 102 records -----
#>
#>
     2 dosing records (see x$get.dosing(); add with add.dosing or et)
     100 observation times (see x$get.sampling(); add with add.sampling or et)
#>
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#>
  -- First part of x: -----
#> # A tibble: 102 x 6
                              ii addl evid
#>
         time cmt
                       \mathtt{amt}
          [h] <chr>
                       <dbl>
                              [h] <int> <evid>
#>
   1 0.0000000 (obs)
                    NA NA
                                    NA 0:Observation
  2 0.0000000 (default) 10000 12
                                    3 1:Dose (Add)
#> 3 0.2424242 (obs)
                     NA NA NA 0:Observation
  4 0.4848485 (obs)
                        NA NA
                                    NA 0:Observation
                                  NA 0:Observation
#> 5 0.7272727 (obs)
                        NA NA
  6 0.9696970 (obs)
                        NA NA NA 0:Observation
  7 1.2121212 (obs)
                        NA NA
                                    NA 0:Observation
   8 1.4545455 (obs)
                         NA
                               NA
                                    NA 0:Observation
#> 9 1.6969697 (obs)
                         NA
                               NA
                                    NA 0:Observation
#> 10 1.9393939 (obs)
                         NA
                             NA
                                    NA 0:Observation
#> # ... with 92 more rows
```

Solving shows what this does in the system:

```
rxSolve(m1, ev) %>% plot(depot,C2, eff)
```

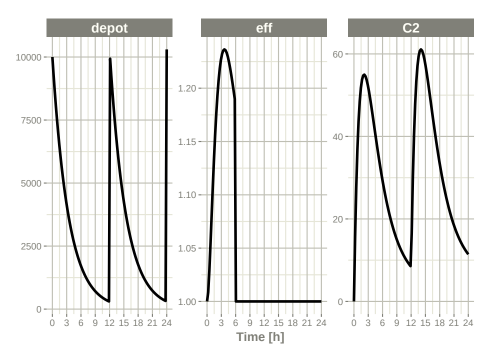


In this case, the depot is turned off, and the depot compartment concentrations are set to the initial values but the other compartment concentrations/levels are not reset. When another dose to the depot is administered the depot compartment is turned back on.

Note that a dose to a compartment only turns back on the compartment that was dosed. Hence if you turn off the effect compartment, it continues to be off after another dose to the depot.

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12, addl=3) %>%
    et(time=6, cmt="-eff", evid=2) %>%
    et(seq(0, 24, length.out=100))

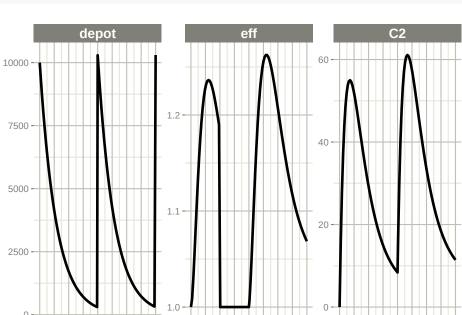
rxSolve(m1, ev) %>% plot(depot,C2, eff)
```



To turn back on the compartment, a zero-dose to the compartment or a evid=2 with the compartment would be needed.

```
ev <- et(timeUnits="hr") %>%
    et(amt=10000, ii=12, addl=3) %>%
    et(time=6, cmt="-eff", evid=2) %>%
    et(time=12,cmt="eff",evid=2) %>%
    et(seq(0, 24, length.out=100))
```

0 3 6 9 12 15 18 21 24



rxSolve(m1, ev) %>% plot(depot,C2, eff)

7.7 Classic RxODE events

0 3 6 9 12 15 18 21 24

Originally RxODE supported compound event IDs; RxODE still supports these parameters, but it is often more useful to use the the normal NONMEM dataset standard that is used by many modeling tools like NONMEM, Monolix and nlmixr, described in the RxODE event types article.

0 3 6 9 12 15 18 21 24 **Time [h]**

Classically, RxODE supported event coding in a single event id evid described in the following table.

100+		<99	
cmt	Infusion/Event Flag	Cmt	SS flag & Turning of Compartment
100+ cmt	0 = bolus dose	< 99 cmt	1 = dose
	1 = infusion (rate)		10 = Steady state 1 (equivalent to SS=1)
	2 = infusion (dur)		20 = Steady state 2 (equivalent to SS=2)
	6 = turn off modeled duration		30 = Turn off a compartment (equivalent to -CMT w/EVID=2)

100 +		<99	
cmt	Infusion/Event Flag	Cmt	SS flag & Turning of Compartment
	7 = turn off modeled rate 8 = turn on modeled duration 9 = turn on modeled rate 4 = replace event 5 = multiply event		

The classic EVID concatenate the numbers in the above table, so an infusion would to compartment 1 would be 10101 and an infusion to compartment 199 would be 119901.

EVID = 0 (observations), EVID=2 (other type event) and EVID=3 are all supported. Internally an EVID=9 is a non-observation event and makes sure the system is initialized to zero; EVID=9 should not be manually set. EVID 10-99 represents modeled time interventions, similar to NONMEM's MTIME. This along with amount (amt) and time columns specify the events in the ODE system.

For infusions specified with EVIDs > 100 the amt column represents the rate value.

For Infusion flags 1 and 2 +amt turn on the infusion to a specific compartment -amt turn off the infusion to a specific compartment. To specify a dose/duration you place the dosing records at the time the duration starts or stops.

For modeled rate/duration infusion flags the on infusion flag must be followed by an off infusion record.

These number are concatenated together to form a full RxODE event ID, as shown in the following examples:

7.7.1 Bolus Dose Examples

A 100 bolus dose to compartment #1 at time 0

time	evid	amt
0	101	100
0.5	0	0
1	0	0

A 100 bolus dose to compartment #99 at time 0

59

time	evid	amt
0	9901	100
0.5	0	0
1	0	0

A 100 bolus dose to compartment #199 at time 0

time	evid	amt
0	109901	100
0.5	0	0
1	0	0

7.7.2 Infusion Event Examples

Bolus infusion with rate 50 to compartment 1 for $1.5~\mathrm{hr}$, (modeled bioavailability changes duration of infusion)

time	evid	amt
0	10101	50
0.5	0	0
1	0	0
1.5	10101	-50

Bolus infusion with rate 50 to compartment 1 for 1.5 hr (modeled bioavailability changes rate of infusion)

time	evid	amt
0	20101	50
0.5	0	0
1	0	0
1.5	20101	-50

Modeled rate with amount of 50

time	evid	am
0	90101	50
0	70101	50
0.5	0	0

time	evid	amt
1	0	0

Modeled duration with amount of 50

time	evid	amt
0	80101	50
0	60101	50
0.5	0	0
1	0	0

7.7.3 Steady State for classic RxODE EVID example

Steady state dose to cmt 1

time	evid	amt
0	110	50

Steady State with super-positioning principle for am 50 and pm 100 dose

time	evid	amt
0	110	50
12	120	100

7.7.4 Turning off a compartment with classic RxODE EVID

Turn off the first compartment at time 12

evid	amt
110	50
130	NA
	110

Event coding in RxODE is encoded in a single event number evid. For compartments under 100, this is coded as:

- This event is 0 for observation events.
- For a specified compartment a bolus dose is defined as:

- 100*(Compartment Number) + 1
- The dose is then captured in the amt
- For IV bolus doses the event is defined as:
 - -10000 + 100*(Compartment Number) + 1
 - The infusion rate is captured in the amt column
 - The infusion is turned off by subtracting amt with the same evid at the stop of the infusion.

For compartments greater or equal to 100, the 100s place and above digits are transferred to the 100,000th place digit. For doses to the 99th compartment the evid for a bolus dose would be 9901 and the evid for an infusion would be 19901. For a bolus dose to the 199th compartment the evid for the bolus dose would be 109901. An infusion dosing record for the 199th compartment would be 119901.

7.8 Datasets for RxODE & nlmixr

Data for input into nlmixr is the same type of data input for RxODE, and it is similar to data for NONMEM (most NONMEM-ready datasets can be used directly in nlmixr).

7.9 Columns Described by Type of Use

7.9.1 Subject Identification Columns

The subject identification column separates subjects for identification of random effects.

• ID: A subject identifier that may be an integer, character, or factor.

7.9.2 Observation Columns

Observation columns are used to indicate the dependent variable and how to use or measure it.

- DV: A numeric column with the measurement
- CENS: A numeric column for indication of censoring, such as below the limit of quantification for an assay.
- LIMIT: A numeric column for helping indicate the type of censoring, such as below the limit of quantification for an assay.

- MDV: An indicator for missing DV values
- CMT: The name or number of the compartment
- DVID: The dependent variable identifier
- EVID The event identifier

7.9.3 Dosing Columns

- AMT: The amount of the dose
- CMT: The name or number of the compartment
- EVID: The event identifier
- ADDL: The number of additional doses
- RATE or DUR: The rate or duration of a dose

7.9.4 Covariate Columns

7.10 Details for Specific Dataset Columns

The details below are sorted alphabetically by column name. For grouping by use, see the documentation above.

7.10.1 AMT Column

The AMT column defines the amount of a dose.

For observation rows, it should be 0 or NA.

For dosing rows, it is the amount of the dose administered to the CMT. If the dose has a zero-order rate (such as a constant infusion), the infusion may be setup using the RATE or DUR column.

7.10.2 CENS/LIMIT Columns

The CENS column is an indicator column indicating if censoring occurred. For pharmacokinetic modeling, censoring is typically when a sample is below the limit of quantification. Internally RxODE saves these values so that nlmixr can use them in likelihood calculations.

CENS = 0 indicates that the value in DV is measured without censoring.

CENS = 1 indicates that a value is left censored (or below the limit of quantitation) and that the value in DV is censoring/quantitation limit.

CENS = -1 indicates that a value is right censored (or above limit of quantitation) and that the value in DV is censoring/quantitation limit.

The LIMIT is additional information about how censoring is handled with nlmixr and is stored in RxODE's data structure as well. When a value is left censored, like below a limit of 1 you may also believe that the value is above a certain threshold, like zero. In this case, a limit of 0 indicates that the censored value is between 0 and 1.

In short when:

CENS = 0 a LIMIT is ignored because the observation is not censored

CENS = 1 the value is censored between (LIMIT, DV)

CENS = -1 the value is censored between (DV, LIMIT)

7.10.3 CMT Column

The CMT column indicates the compartment where an event occurs. When given as a character string or factor (the preferred method), it is matched by name in the model. When given as an integer, it is matched by the order that compartments appear in the model.

7.10.4 DUR Column

The DUR column defines the duration of an infusion. It is used to set the duration of a zero-order rate of infusion.

7.10.5 DV Column

The DV column indicates the current measurement in the current compartment (see CMT) with the current measurement identifier (see DVID) which may be missing (see MDV) or censored (see CENS).

7.10.6 DVID Column

TODO

7.10.7 EVID Column

The EVID column is the event identifier for a row of data.

For observation records, it will be 0. For normal dosing records, it will be 1. Many more EVID values are detailed in the RxODE Event Types and Classic RxODE Events vignettes.

7.10.8 ID Column

The ID column is a subject identifier. This column is used to separate one individual (usually a single person or animal) from another.

In the model, the ID column is used to separate individuals. The numerical integrator re-initializes with each new individual, and new values for all random effects are selected.

7.10.9 RATE Column

TODO

Chapter 8

Easily creating RxODE events

An event table in RxODE is a specialized data frame that acts as a container for all of RxODE's events and observation times.

To create an RxODE event table you may use the code eventTable(), et(), or even create your own data frame with the right event information contained in it. This is closely related to the types of events that RxODE supports.

```
library(RxODE)
library(units)
#> udunits database from /usr/share/xml/udunits/udunits2.xml
(ev <- eventTable())</pre>
#> ------ EventTable with 0 records ------
#>
#>
     O dosing records (see x$get.dosing(); add with add.dosing or et)
     O observation times (see x$get.sampling(); add with add.sampling or et)
or
(ev <- et())
#> ------ EventTable with 0 records ------
     O dosing records (see x$get.dosing(); add with add.dosing or et)
#>
#>
     O observation times (see x$get.sampling(); add with add.sampling or et)
```

With this event table you can add sampling/observations or doses by piping or direct access.

This is a short table of the two main functions to create dosing

add.dosing()	et()	Description
dose nbr.doses dosing.interval dosing.to rate start.time	amt addl ii cmt rate time dur	Dose/Rate/Duration amount Additional doses or number of doses Dosing Interval Dosing Compartment Infusion rate Dosing start time Infusion Duration

Sampling times can be added with add.sampling (sampling times) or et(sampling times). Dosing intervals and sampling windows are also supported.

For these models, we can illustrate by using the model shared in the RxODE tutorial:

```
## Model from RxODE tutorial
m1 <-RxODE({
    KA=2.94E-01;
   CL=1.86E+01;
   V2=4.02E+01;
    Q=1.05E+01;
    V3=2.97E+02;
   Kin=1;
    Kout=1;
    EC50=200;
    ## Added modeled bioavaiblity, duration and rate
    fdepot = 1;
    durDepot = 8;
    rateDepot = 1250;
    C2 = centr/V2;
    C3 = peri/V3;
    d/dt(depot) =-KA*depot;
    f(depot) = fdepot
    dur(depot) = durDepot
    rate(depot) = rateDepot
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) =
                                      Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
    eff(0) = 1
})
```

8.1 Adding doses to the event table

Once created you can add dosing to the event table by the $\mathtt{add.dosing}()$, and $\mathtt{et}()$ functions.

Using the add.dosing() function you have:

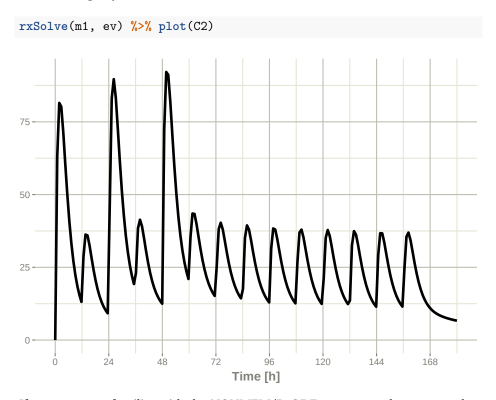
argument	meaning
dose	dose amount
nbr.doses	Number of doses; Should be at least 1.
dosing.interval	Dosing interval; By default this is 24.
dosing.to	Compartment where dose is administered.
rate	Infusion rate
start.time	The start time of the dose

```
#> ------ EventTable with 2 records ------
#>
     2 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
     O observation times (see x$get.sampling(); add with add.sampling or et)
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: -----
#> # A tibble: 2 x 5
                ii addl evid
#>
     time amt
      [h] [mg] [h] <int> <evid>
#>
     0 10000 24 2 1:Dose (Add)
#> 1
#> 2
    0 5000 12 13 1:Dose (Add)
```

Notice that the units were specified in the table. When specified, the units use the units package to keep track of the units and convert them if needed. Additionally,

ggforce uses them to label the ggplot axes. The set_units and drop_units are useful to set and drop the RxODE event table units.

In this example, you can see the time axes is labeled:



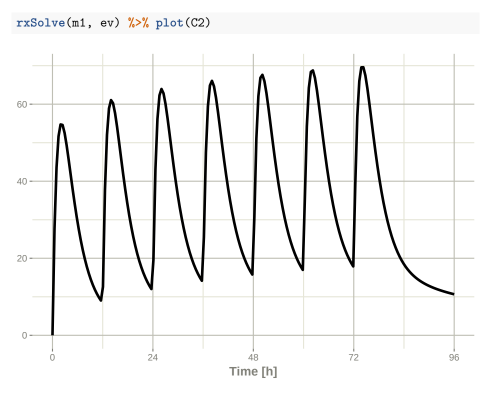
If you are more familiar with the NONMEM/RxODE event records, you can also specify dosing using et with the dose elements directly:

ev <- et(timeUnits="hr") %>%

```
et(amt=10000, until = set_units(3, days),
    ii=12) # loading doses
ev
#> ----- EventTable with 1 records -----
#>
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
     O observation times (see x$get.sampling(); add with add.sampling or et)
#>
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: ------
#> # A tibble: 1 x 5
#>
     time
           amt
                 ii addl evid
```

```
#> [h] <dbl> [h] <int> <evid>
#> 1     0 10000     12     6 1:Dose (Add)
```

Which gives:



This shows how easy creating event tables can be.

8.2 Adding sampling to an event table

If you notice in the above examples, RxODE generated some default sampling times since there was not any sampling times. If you wish more control over the sampling time, you should add the samples to the RxODE event table by add.sampling or et

```
ev <- eventTable(amount.units="mg", time.units="hr")

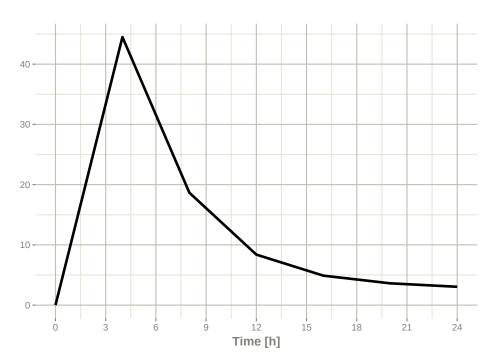
## The methods ar attached to the event table, so you can use them
## directly
ev$add.dosing(dose=10000, nbr.doses = 3)# loading doses</pre>
```

```
ev$add.sampling(seq(0,24,by=4))
ev
```

```
#> ------ EventTable with 8 records ------
#>
    1 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
#>
    7 observation times (see x$get.sampling(); add with add.sampling or et)
    multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: ------
#> # A tibble: 8 x 5
    time amt
               ii addl evid
     [h] [mg] [h] <int> <evid>
#>
#> 1
      O NA NA NA O:Observation
#> 2
      0 10000 24
                    2 1:Dose (Add)
         NA NA NA 0:Observation
#> 3
      4
#> 4
           NA NA NA 0:Observation
      8
#> 5
     12 NA NA NA 0:Observation
      16
#> 6
           NA
                NA NA 0:Observation
#> 7
      20
               NA NA 0:Observation
           NA
#> 8 24
           NA NA NA O:Observation
```

Which gives:

```
solve(m1, ev) %>% plot(C2)
```



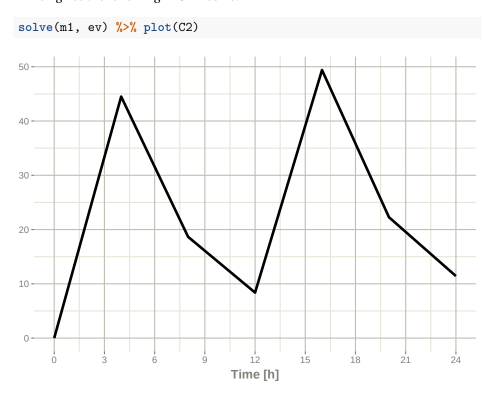
Or if you use et you can simply add them in a similar way to add.sampling:

```
ev <- et(timeUnits="hr") %>%
  et(amt=10000, until = set_units(3, days),
        ii=12) %>% # loading doses
  et(seq(0,24,by=4))
```

```
#> ----- EventTable with 8 records -----
#>
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
     7 observation times (see x$get.sampling(); add with add.sampling or et)
#>
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: -----
#> # A tibble: 8 x 5
#>
     time amt
                 ii addl evid
#>
      [h] <dbl>
                [h] <int> <evid>
#> 1
       0
            NA
               NA
                      NA 0:Observation
#> 2
       0 10000
               12
                       6 1:Dose (Add)
#> 3
       4
           NA
               NA
                      NA 0:Observation
#> 4
       8
           NA
               NA NA 0:Observation
      12 NA NA NA 0:Observation
#> 5
           NA NA NA O:Observation
#> 6
      16
```

```
\#> 7 20 NA NA NA 0:Observation \#> 8 24 NA NA NA 0:Observation
```

which gives the following RxODE solve:



Note the jagged nature of these plots since there was only a few sample times.

8.3 Expand the event table to a multi-subject event table.

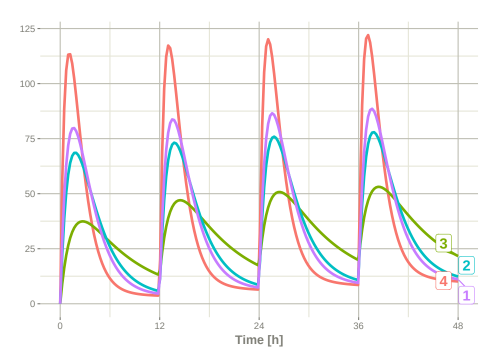
The only thing that is needed to expand an event table is a list of IDs that you want to expand;

```
#> ----- EventTable with 804 records -----
#>
    4 individuals
#>
    4 dosing records (see x$get.dosing(); add with add.dosing or et)
    800 observation times (see x$get.sampling(); add with add.sampling or et)
#>
    multiple doses in `addl` columns, expand with x=expand(); or etExpand(x)
#> -- First part of x: -----
#> # A tibble: 804 x 6
#>
      id
            time amt ii addl evid
            [h] <dbl> [h] <int> <evid>
#>
    <int>
\#> 1 1 0.0000000 NA NA NA 0:Observation
      1 0.0000000 10000 12 6 1:Dose (Add)
#> 3
      1 0.2412060 NA NA NA 0:Observation
      1 0.4824121 NA NA NA 0:Observation
#> 4
#> 5
     1 0.7236181 NA NA NA 0:Observation
#> 6 1 0.9648241 NA NA NA 0:Observation
      1 1.2060302 NA NA NA 0:Observation
#> 7
     1 1.4472362 NA NA NA 0:Observation
#> 9 1 1.6884422 NA NA NA 0:Observation
#> # ... with 794 more rows
```

You can see in the following simulation there are 4 individuals that are solved for:

```
#> Warning: 'ID' missing in 'parameters' dataset
```

^{#&}gt; individual parameters are assumed to have the same order as the event dataset



8.4 Add doses and samples within a sampling window

In addition to adding fixed doses and fixed sampling times, you can have windows where you sample and draw doses from. For dosing windows you specify the time as an ordered numerical vector with the lowest dosing time and the highest dosing time inside a list.

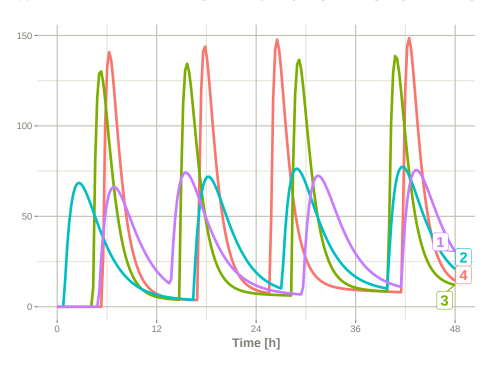
In this example, you start with a dosing time with a 6 hour dosing window:

```
#> # A tibble: 16 x 6
#>
       id
            low
                     time high
                                amt evid
#>
     <int>
            [h]
                      [h] [h] <dbl> <evid>
             0 5.4888363
                           6 10000 1:Dose (Add)
#>
   1
        1
            #>
  2
        1
#>
   3
        1
            24 25.7168372 30 10000 1:Dose (Add)
#>
  4
        1
           36 41.6224525 42 10000 1:Dose (Add)
#> 5
        2
            0 4.3146735 6 10000 1:Dose (Add)
           12 14.7464507 18 10000 1:Dose (Add)
24 28.2303887 30 10000 1:Dose (Add)
        2
#> 6
#> 7
        2
#> 8
        2
           36 39.9419537 42 10000 1:Dose (Add)
#> 9
        3
             0 0.8079996
                           6 10000 1:Dose (Add)
        3
           12 16.4195299 18 10000 1:Dose (Add)
#> 10
#> 11
        3
           24 27.1145757 30 10000 1:Dose (Add)
        3
           36 39.8504731 42 10000 1:Dose (Add)
#> 12
#> 13
            0 4.9826858
        4
                            6 10000 1:Dose (Add)
           12 13.7168372
#> 14
                            18 10000 1:Dose (Add)
#> 15
        4 24 29.6224525 30 10000 1:Dose (Add)
#> 16
        4 36 41.4888363 42 10000 1:Dose (Add)
```

You can clearly see different dosing times in the following simulation:

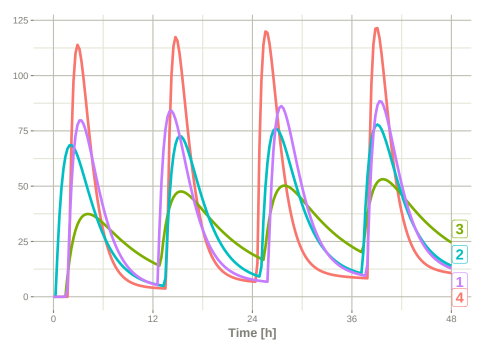
```
#> Warning: 'ID' missing in 'parameters' dataset
```

^{#&}gt; individual parameters are assumed to have the same order as the event dataset



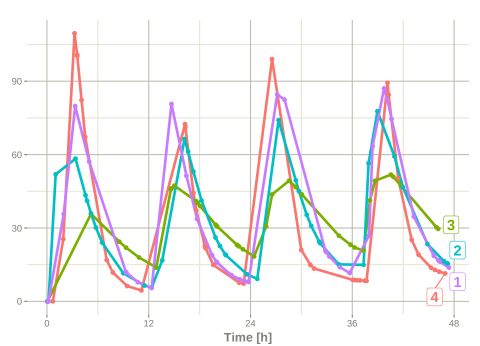
Of course in reality the dosing interval may only be 2 hours:

- #> Warning: 'ID' missing in 'parameters' dataset
- #> individual parameters are assumed to have the same order as the event dataset



The same sort of thing can be specified with sampling times. To specify the sampling times in terms of a sampling window, you can create a list of the sampling times. Each sampling time will be a two element ordered numeric vector.

#> Warning: 'ID' missing in 'parameters' dataset



#> individual parameters are assumed to have the same order as the event dataset

This shows the flexibility in dosing and sampling that the RxODE event tables allow.

8.5 Combining event tables

Since you can create dosing records and sampling records, you can create any complex dosing regimen you wish. In addition, RxODE allows you to combine event tables by c, seq, rep, and rbind.

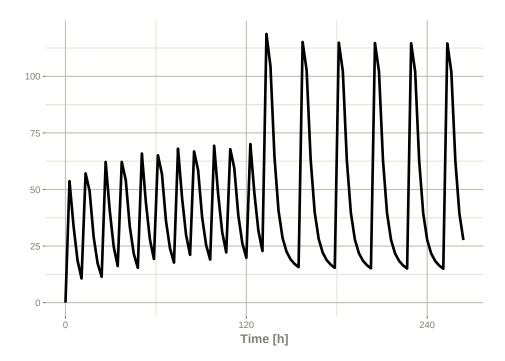
8.6 Sequencing event tables

One way to combine event table is to sequence them by c, seq or etSeq. This takes the two dosing groups and adds at least one inter-dose interval between them:

```
## bid for 5 days
bid <- et(timeUnits="hr") %>%
            et(amt=10000,ii=12,until=set_units(5, "days"))

## qd for 5 days
qd <- et(timeUnits="hr") %>%
```

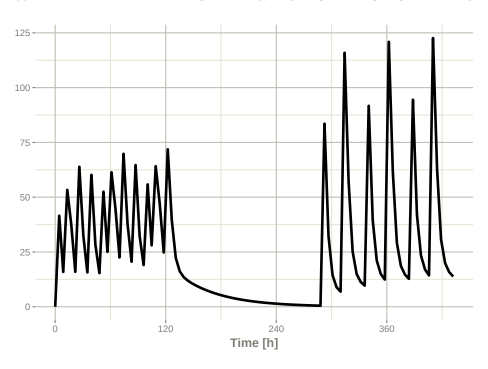
```
et(amt=20000,ii=24,until=set_units(5, "days"))
## bid for 5 days followed by qd for 5 days
et <- seq(bid,qd) %>% et(seq(0,11*24,length.out=100));
rxSolve(m1, et) %>% plot(C2)
```



When sequencing events, you can also separate this sequence by a period of time; For example if you wanted to separate this by a week, you could easily do that with the following sequence of event tables:

```
## bid for 5 days followed by qd for 5 days
et <- seq(bid,set_units(1, "week"), qd) %>%
    et(seq(0,18*24,length.out=100));

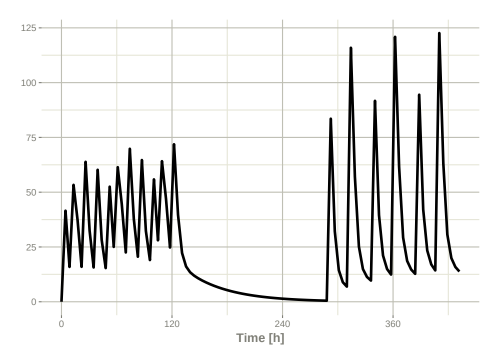
rxSolve(m1, et) %>% plot(C2)
```



Note that in this example the time between the bid and the qd event tables is exactly one week, not 1 week plus 24 hours because of the inter-dose interval. If you want that behavior, you can sequence it using the wait="+ii".

```
## bid for 5 days followed by qd for 5 days
et <- seq(bid,set_units(1, "week"), qd,wait="+ii") %>%
    et(seq(0,18*24,length.out=100));

rxSolve(m1, et) %>% plot(C2)
```



Also note, that RxODE assumes that the dosing is what you want to space the event tables by, and clears out any sampling records when you combine the event tables. If that is not true, you can also use the option samples="use"

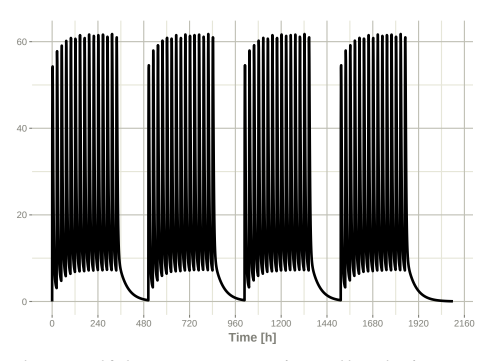
8.7 Repeating event tables

You can have an event table that you can repeat with etRep or rep. For example 4 rounds of 2 weeks on QD therapy and 1 week off of therapy can be simply specified:

```
qd <-et(timeUnits = "hr") %>%
  et(amt=10000, ii=24, until=set_units(2, "weeks"), cmt="depot")

et <- rep(qd, times=4, wait=set_units(1, "weeks")) %>%
        add.sampling(set_units(seq(0, 12.5,by=0.005),weeks))

rxSolve(m1, et) %>% plot(C2)
```



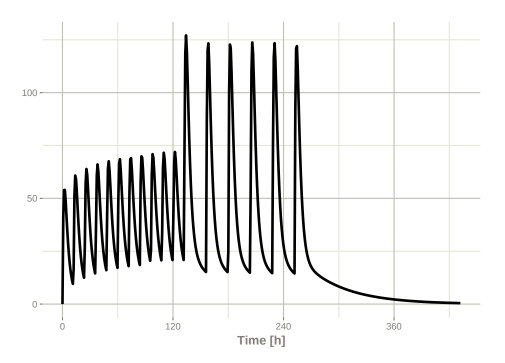
This is a simplified way to use a sequence of event tables. Therefore, many of the same options still apply; That is samples are cleared unless you use samples="use", and the time between event tables is at least the inter-dose interval. You can adjust the timing by the wait option.

8.8 Combining event tables with rbind

You may combine event tables with rbind. This does not consider the event times when combining the event tables, but keeps them the same times. If you space the event tables by a waiting period, it also does not consider the inter-dose interval.

Using the previous seq you can clearly see the difference. Here was the sequence:

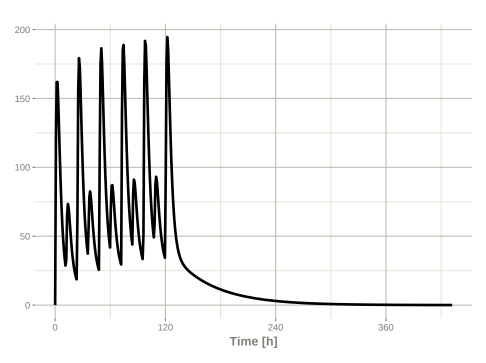
```
rxSolve(m1, et) %>% plot(C2)
```



But if you bind them together with rbind

```
## bid for 5 days
et <- rbind(bid,qd) %>%
    et(seq(0,18*24,length.out=500));

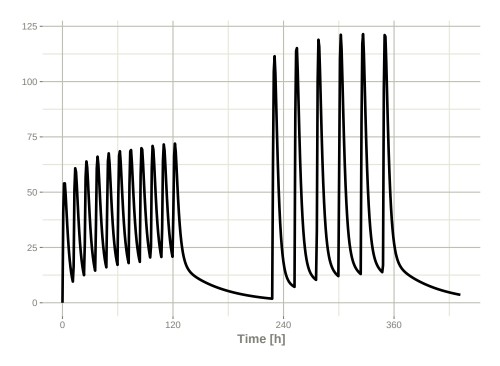
rxSolve(m1, et) %>% plot(C2)
```



Still the waiting period applies (but does not consider the inter-dose interval)

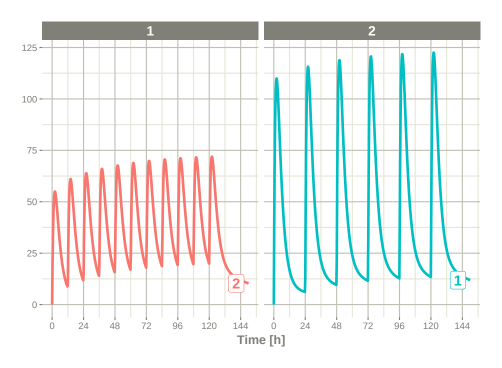
```
et <- rbind(bid,wait=set_units(10,days),qd) %>%
    et(seq(0,18*24,length.out=500));

rxSolve(m1, et) %>% plot(C2)
```



You can also bind the tables together and make each ID in the event table unique; This can be good to combine cohorts with different expected dosing and sampling times. This requires the id="unique" option; Using the first example shows how this is different in this case:

```
## bid for 5 days
et <- etRbind(bid,qd, id="unique") %>%
    et(seq(0,150,length.out=500));
library(ggplot2)
rxSolve(m1, et) %>% plot(C2) + facet_wrap( ~ id)
```



8.9 Expanding events

Event tables can be expanded so they contain an addl data item, like the following example:

```
ev <- et() %>%
 et(dose=50, ii=8, until=48)
  ----- EventTable with 1 records -----
#>
     1 dosing records (see x$get.dosing(); add with add.dosing or et)
#>
#>
     O observation times (see x$get.sampling(); add with add.sampling or et)
     multiple doses in `addl` columns, expand with x$expand(); or etExpand(x)
#> -- First part of x: ------
#> # A tibble: 1 x 5
     time
           amt
                 ii addl evid
    <dbl> <dbl> <int> <evid>
#> 1
            50
                  8
                       6 1:Dose (Add)
```

You can expand the events so they do not have the addl items by \$expand() or etExpand(ev):

print(ev)

The first, etExpand(ev) expands the event table without modifying the original data frame:

```
etExpand(ev)
#> ------ EventTable with 7 records ------
#>
#>
    7 dosing records (see x$get.dosing(); add with add.dosing or et)
    O observation times (see x$get.sampling(); add with add.sampling or et)
#> -- First part of x: -----
#> # A tibble: 7 x 4
    time amt ii evid
#> <dbl> <dbl> <evid>
#> 1
    0 50 0 1:Dose (Add)
#> 2
     8 50 0 1:Dose (Add)
#> 3 16 50 0 1:Dose (Add)
#> 4 24 50 0 1:Dose (Add)
#> 5 32 50 0 1:Dose (Add)
#> 6 40 50 0 1:Dose (Add)
#> 7
      48
          50 0 1:Dose (Add)
```

You can see the addl events were expanded, however the original data frame remained intact:

If you use ev\$expand() it will modify the ev object. This is similar to an object-oriented method:

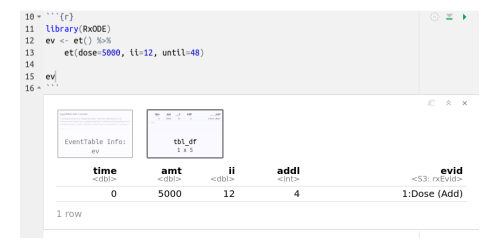
```
ev$expand()
ev
```

```
#>
  ----- EventTable with 7 records -----
#>
#>
     7 dosing records (see x$get.dosing(); add with add.dosing or et)
     O observation times (see x$get.sampling(); add with add.sampling or et)
#>
#> -- First part of x: ------
#> # A tibble: 7 x 4
#>
     time
           \mathtt{amt}
                  ii evid
    <dbl> <dbl> <dbl> <evid>
#>
#> 1
                  0 1:Dose (Add)
        0
            50
#> 2
        8
            50
                   0 1:Dose (Add)
#> 3
       16
            50
                  0 1:Dose (Add)
#> 4
       24
            50
                   0 1:Dose (Add)
#> 5
       32
            50
                   0 1:Dose (Add)
#> 6
       40
            50
                   0 1:Dose (Add)
#> 7
       48
            50
                   0 1:Dose (Add)
```

8.10 Event tables in Rstudio Notebooks

In addition to the output in the console which has been shown in the above examples, Rstudio notebook output is different and can be seen in the following screenshots;

The first screenshot shows how the event table looks after evaluating it in the Rstduio notebook



This is a simple dataframe that allows you to page through the contents. If you click on the first box in the Rstudio notebook output, it will have the notes about the event table:



Chapter 9

Solving and solving options

In general, ODEs are solved using a combination of:

- A compiled model specification from RxODE(), specified with object=
- Input parameters, specified with params= (and could be blank)
- Input data or event table, specified with events=
- Initial conditions, specified by inits= (and possibly in the model itself by state(0)=)

The solving options are given in the sections below:

9.1 General Solving Options

9.1.1 object

object is a either a RxODE family of objects, or a file-name with a RxODE model specification, or a string with a RxODE model specification.

9.1.2 params

params a numeric named vector with values for every parameter in the ODE system; the names must correspond to the parameter identifiers used in the ODE specification;

9.1.3 events

events an eventTable object describing the input (e.g., doses) to the dynamic system and observation sampling time points (see [eventTable()]);

9.1.4 inits

inits a vector of initial values of the state variables (e.g., amounts in each compartment), and the order in this vector must be the same as the state variables (e.g., PK/PD compartments);

9.1.5 method

method The method for solving ODEs. Currently this supports:

- "liblsoda" thread safe Isoda. This supports parallel thread-based solving, and ignores user Jacobian specification.
- "lsoda" LSODA solver. Does not support parallel thread-based solving, but allows user Jacobian specification.
- "dop853" DOP853 solver. Does not support parallel thread-based solving nor user Jacobain specification
- "indLin" Solving through inductive linearization. The RxODE dll must be setup specially to use this solving routine.

9.1.6 stiff

stiff a logical (TRUE by default) indicating whether the ODE system is stiff or not.

For stiff ODE systems (stiff = TRUE), RxODE uses the LSODA (Livermore Solver for Ordinary Differential Equations) Fortran package, which implements an automatic method switching for stiff and non-stiff problems along the integration interval, authored by Hindmarsh and Petzold (2003).

For non-stiff systems (stiff = FALSE), RxODE uses DOP853, an explicit Runge-Kutta method of order 8(5, 3) of Dormand and Prince as implemented in C by Hairer and Wanner (1993).

If stiff is not specified, the method argument is used instead.

9.2 lsoda/dop solving options

9.2.1 atol

atol a numeric absolute tolerance (1e-8 by default) used by the ODE solver to determine if a good solution has been achieved; This is also used in the solved linear model to check if prior doses do not add anything to the solution.

9.2.2 rtol

rtol a numeric relative tolerance (1e-6 by default) used by the ODE solver to determine if a good solution has been achieved. This is also used in the solved linear model to check if prior doses do not add anything to the solution.

9.2.3 maxsteps

maxsteps maximum number of (internally defined) steps allowed during one call to the solver. (5000 by default)

9.2.4 hmin

hmin The minimum absolute step size allowed. The default value is 0.

9.2.5 hmax

hmax The maximum absolute step size allowed. When hmax=NA (default), uses the average difference + hmaxSd*sd in times and sampling events. The hmaxSd is a user specified parameter and which defaults to zero. When hmax=NULL RXODE uses the maximum difference in times in your sampling and events. The value 0 is equivalent to infinite maximum absolute step size.

9.2.6 hmaxSd

 $\verb|hmaxSd|$ The number of standard deviations of the time difference to add to hmax. The default is 0

9.2.7 hini

hini The step size to be attempted on the first step. The default value is determined by the solver (when hini = 0)

9.2.8 maxordn

maxordn The maximum order to be allowed for the nonstiff (Adams) method. The default is 12. It can be between 1 and 12.

9.2.9 maxords

maxords The maximum order to be allowed for the stiff (BDF) method. The default value is 5. This can be between 1 and 5.

9.2.10 mxhnil

mxhnil maximum number of messages printed (per problem) warning that T + H = T on a step (H = step size). This must be positive to result in a non-default value. The default value is 0 (or infinite).

9.2.11 hmxi

hmxi inverse of the maximum absolute value of H to are used. hmxi = 0.0 is allowed and corresponds to an infinite hmax1 (default).hminandhmximay be changed at any time, but will not take effect until the next change of H is considered. This option is only considered withmethod="liblsoda".

9.2.12 istateReset

istateReset When TRUE, reset the ISTATE variable to 1 for Isoda and libIsoda with doses, like deSolve; When FALSE, do not reset the ISTATE variable with doses.

9.3 Inductive Linerization Options

9.3.1 indLinMatExpType

indLinMatExpType This is them matrix exponential type that is use for RxODE. Currently the following are supported:

- Al-Mohy Uses the exponential matrix method of Al-Mohy Higham (2009)
- arma Use the exponential matrix from RcppArmadillo
- expokit Use the exponential matrix from Roger B. Sidje (1998)

9.3.2 indLinMatExpOrder

indLinMatExpOrder an integer, the order of approximation to be used, for the Al-Mohy and expokit values. The best value for this depends on machine precision (and slightly on the matrix). We use 6 as a default.

9.3.3 indLinPhiTol

indLinPhiTol the requested accuracy tolerance on exponential matrix.

9.3.4 indLinPhiM

indLinPhiM the maximum size for the Krylov basis

9.4 Steady State Solving Options

9.4.1 minSS

minSS Minimum number of iterations for a steady-state dose

9.4.2 maxSS

maxSS Maximum number of iterations for a steady-state dose

9.4.3 strictSS

strictSS Boolean indicating if a strict steady-state is required. If a strict steady-state is (TRUE) required then at least minSS doses are administered and the total number of steady states doses will continue until maxSS is reached, or atol and rtol for every compartment have been reached. However, if ODE solving problems occur after the minSS has been reached the whole subject is considered an invalid solve. If strictSS is FALSE then as long as minSS has been reached the last good solve before ODE solving problems occur is considered the steady state, even though either atol, rtol or maxSS have not been achieved.

9.4.4 infSSstep

infSSstep Step size for determining if a constant infusion has reached steady state. By default this is large value, 420.

9.4.5 ssAtol

ssAtol Steady state atol convergence factor. Can be a vector based on each state.

9.4.6 ssRtol

ssRtol Steady state rtol convergence factor. Can be a vector based on each state.

9.5 RxODE numeric stability options

9.5.1 maxAtolRtolFactor

 ${\tt maxAtolRtolFactor}$ The maximum atol/rtol that FOCEi and other routines may adjust to. By default 0.1

9.5.2 stateTrim

stateTrim When amounts/concentrations in one of the states are above this value, trim them to be this value. By default Inf. Also trims to -stateTrim for large negative amounts/concentrations. If you want to trim between a range say c(0, 2000000) you may specify 2 values with a lower and upper range to make sure all state values are in the reasonable range.

9.5.3 safeZero

safeZero Use safe zero divide and log routines. By default this is turned on but you may turn it off if you wish.

9.5.4 sumType

sumType Sum type to use for sum() in RxODE code blocks.

pairwise uses the pairwise sum (fast, default)

fsum uses Python's fsum function (most accurate)

kahan uses Kahan correction

neumaier uses Neumaier correction

c uses no correction: default/native summing

9.5.5 prodType

prodType Product to use for prod() in RxODE blocks

long double converts to long double, performs the multiplication and then converts back.

double uses the standard double scale for multiplication.

9.5.6 maxwhile

maxwhile represents the maximum times a while loop is evaluated before exiting. By default this is 100000

9.5.7 transitAbs

transitAbs boolean indicating if this is a transit compartment absorption

9.6 Linear compartment model sensitivity options

9.6.1 sensType

sensType Sensitivity type for linCmt() model: advan Use the direct advan solutions autodiff Use the autodiff advan solutions forward Use forward difference solutions central Use central differences

9.6.2 linDiff

linDiff This gives the linear difference amount for all the types of linear compartment model parameters where sensitivities are not calculated. The named components of this numeric vector are:

- "lag" Central compartment lag
- "f" Central compartment bioavailability
- "rate" Central compartment modeled rate
- "dur" Central compartment modeled duration
- "lag2" Depot compartment lag
- "f2" Depot compartment bioavailability

- "rate2" Depot compartment modeled rate
- "dur2" Depot compartment modeled duration

9.6.3 linDiffCentral

linDiffCentral This gives the which parameters use central differences for the linear compartment model parameters. The are the same components as linDiff

9.7 Covariate Solving Options

9.7.1 iCov

iCov A data frame of individual non-time varying covariates to combine with the params to form a parameter data.frame.

9.7.2 covsInterpolation

covsInterpolation specifies the interpolation method for time-varying covariates. When solving ODEs it often samples times outside the sampling time specified in events. When this happens, the time varying covariates are interpolated. Currently this can be:

- "linear" interpolation, which interpolates the covariate by solving the line between the observed covariates and extrapolating the new covariate value.
- "constant" Last observation carried forward (the default).
- "NOCB" Next Observation Carried Backward. This is the same method that NONMEM uses.
- "midpoint" Last observation carried forward to midpoint; Next observation carried backward to midpoint.

9.7.3 addCov

addCov A boolean indicating if covariates should be added to the output matrix or data frame. By default this is disabled.

9.8 Simulation options

9.8.1 seed

seed an object specifying if and how the random number generator should be initialized

9.8.2 nsim

nsim represents the number of simulations. For RxODE, if you supply single subject event tables (created with [eventTable()])

9.8.3 thetaMat

thetaMat Named theta matrix.

9.8.4 thetaLower

thetaLower Lower bounds for simulated population parameter variability (by default -Inf)

9.8.5 thetaUpper

thetaUpper Upper bounds for simulated population unexplained variability (by default Inf)

9.8.6 thetaDf

thetaDf The degrees of freedom of a t-distribution for simulation. By default this is NULL which is equivalent to Inf degrees, or to simulate from a normal distribution instead of a t-distribution.

9.8.7 thetaIsChol

thetaIsChol Indicates if the theta supplied is a Cholesky decomposed matrix instead of the traditional symmetric matrix.

9.8.8 nStud

nStud Number virtual studies to characterize uncertainty in estimated parameters.

9.8.9 omega

omega Estimate of Covariance matrix. When omega is a list, assume it is a block matrix and convert it to a full matrix for simulations.

9.8.10 omegalsChol

omegaIsChol Indicates if the omega supplied is a Cholesky decomposed matrix instead of the traditional symmetric matrix.

9.8.11 omegaSeparation

omegaSeparation Omega separation strategy

Tells the type of separation strategy when simulating covariance with parameter uncertainty with standard deviations modeled in the thetaMat matrix.

- "lkj" simulates the correlation matrix from the rLKJ1 matrix with the distribution parameter eta equal to the degrees of freedom nu by (nu-1)/2
- "separation" simulates from the identity inverse Wishart covariance matrix with nu degrees of freedom. This is then converted to a covariance matrix and augmented with the modeled standard deviations. While computationally more complex than the "lkj" prior, it performs better when the covariance matrix size is greater or equal to 10
- "auto" chooses "lkj" when the dimension of the matrix is less than 10 and "separation" when greater than equal to 10.

9.8.12 omegaXform

omegaXform When taking omega values from the thetaMat simulations (using the separation strategy for covariance simulation), how should the thetaMat values be turned int standard deviation values:

• identity This is when standard deviation values are directly modeled by the params and thetaMat matrix

- variance This is when the params and thetaMat simulates the variance that are directly modeled by the thetaMat matrix
- log This is when the params and thetaMat simulates log(sd)
- nlmixrSqrt This is when the params and thetaMat simulates the inverse cholesky decomposed matrix with the x^2 modeled along the diagonal. This only works with a diagonal matrix.
- nlmixrLog This is when the params and thetaMat simulates the inverse cholesky decomposed matrix with the exp(x^2) along the diagonal. This only works with a diagonal matrix.
- nlmixrIdentity This is when the params and thetaMat simulates the inverse cholesky decomposed matrix. This only works with a diagonal matrix.

9.8.13 omegaLower

omegaLower Lower bounds for simulated ETAs (by default -Inf)

9.8.14 omegaUpper

omegaUpper Upper bounds for simulated ETAs (by default Inf)

9.8.15 omegaDf

omegaDf The degrees of freedom of a t-distribution for simulation. By default this is NULL which is equivalent to Inf degrees, or to simulate from a normal distribution instead of a t-distribution.

9.8.16 nSub

nSub Number between subject variabilities (ETAs) simulated for every realization of the parameters.

9.8.17 dfSub

dfSub Degrees of freedom to sample the between subject variability matrix from the inverse Wishart distribution (scaled) or scaled inverse chi squared distribution.

9.8.18 sigma

sigma Named sigma covariance or Cholesky decomposition of a covariance matrix. The names of the columns indicate parameters that are simulated. These are simulated for every observation in the solved system.

9.8.19 sigmaLower

sigmaLower Lower bounds for simulated unexplained variability (by default -Inf)

9.8.20 sigmaUpper

sigmaUpper Upper bounds for simulated unexplained variability (by default Inf)

9.8.21 sigmaXform

sigmaXform When taking sigma values from the thetaMat simulations (using the separation strategy for covariance simulation), how should the thetaMat values be turned int standard deviation values:

- identity This is when standard deviation values are directly modeled by the params and thetaMat matrix
- variance This is when the params and thetaMat simulates the variance that are directly modeled by the thetaMat matrix
- log This is when the params and thetaMat simulates log(sd)
- nlmixrSqrt This is when the params and thetaMat simulates the inverse cholesky decomposed matrix with the x^2 modeled along the diagonal. This only works with a diagonal matrix.
- nlmixrLog This is when the params and thetaMat simulates the inverse cholesky decomposed matrix with the $\exp(x^2)$ along the diagonal. This only works with a diagonal matrix.
- nlmixrIdentity This is when the params and thetaMat simulates the inverse cholesky decomposed matrix. This only works with a diagonal matrix.

9.8.22 sigmaDf

sigmaDf Degrees of freedom of the sigma t-distribution. By default it is equivalent to Inf, or a normal distribution.

9.8.23 sigmaIsChol

sigmaIsChol Boolean indicating if the sigma is in the Cholesky decomposition instead of a symmetric covariance

9.8.24 sigmaSeparation

sigmaSeparation separation strategy for sigma;

Tells the type of separation strategy when simulating covariance with parameter uncertainty with standard deviations modeled in the thetaMat matrix.

- "lkj" simulates the correlation matrix from the rLKJ1 matrix with the distribution parameter eta equal to the degrees of freedom nu by (nu-1)/2
- "separation" simulates from the identity inverse Wishart covariance matrix with nu degrees of freedom. This is then converted to a covariance matrix and augmented with the modeled standard deviations. While computationally more complex than the "lkj" prior, it performs better when the covariance matrix size is greater or equal to 10
- "auto" chooses "lkj" when the dimension of the matrix is less than 10 and "separation" when greater than equal to 10.

9.8.25 dfObs

dfObs Degrees of freedom to sample the unexplained variability matrix from the inverse Wishart distribution (scaled) or scaled inverse chi squared distribution.

9.8.26 resample

resample A character vector of model variables to resample from the input dataset; This sampling is done with replacement. When NULL or FALSE no resampling is done. When TRUE resampling is done on all covariates in the input dataset

9.8.27 resampleID

resampleID boolean representing if the resampling should be done on an individual basis TRUE (ie. a whole patient is selected) or each covariate is resampled independent of the subject identifier FALSE. When resampleID=TRUE correlations of parameters are retained, where as when resampleID=FALSE ignores patient covariate correaltions. Hence the default is resampleID=TRUE.

9.9 RxODE output options

9.9.1 returnType

returnType This tells what type of object is returned. The currently supported types are:

- "rxSolve" (default) will return a reactive data frame that can change easily change different pieces of the solve and update the data frame. This is the currently standard solving method in RxODE, is used for rxSolve(object, ...), solve(object,...),
- "data.frame" returns a plain, non-reactive data frame; Currently very slightly faster than returnType="matrix"
- "matrix" returns a plain matrix with column names attached to the solved object. This is what is used object\$run as well as object\$solve
- "data.table" returns a data.table; The data.table is created by reference (ie setDt()), which should be fast.
- "tbl" or "tibble" returns a tibble format.

9.9.2 addDosing

addDosing Boolean indicating if the solve should add RxODE EVID and related columns. This will also include dosing information and estimates at the doses. Be default, RxODE only includes estimates at the observations. (default FALSE). When addDosing is NULL, only include EVID=0 on solve and exclude any modeltimes or EVID=2. If addDosing is NA the classic RxODE EVID events are returned. When addDosing is TRUE add the event information in NONMEM-style format; If subsetNonmem=FALSE RxODE will also include extra event types (EVID) for ending infusion and modeled times:

- EVID=-1 when the modeled rate infusions are turned off (matches rate=-1)
- EVID=-2 When the modeled duration infusions are turned off (matches rate=-2)
- EVID=-10 When the specified rate infusions are turned off (matches rate>0)
- EVID=-20 When the specified dur infusions are turned off (matches dur>0)
- EVID=101,102,103,... Modeled time where 101 is the first model time, 102 is the second etc.

9.9.3 keep

keep Columns to keep from either the input dataset or the iCov dataset. With the iCov dataset, the column is kept once per line. For the input dataset, if any records are added to the data LOCF (Last Observation Carried forward) imputation is performed.

9.9.4 drop

drop Columns to drop from the output

9.9.5 idFactor

idFactor This boolean indicates if original ID values should be maintained. This changes the default sequentially ordered ID to a factor with the original ID values in the original dataset. By default this is enabled.

9.9.6 subsetNonmem

subsetNonmem subset to NONMEM compatible EVIDs only. By default TRUE.

9.9.7 matrix

 $\verb|matrix| A boolean indicating if a matrix should be returned instead of the RxODE's solved object.$

9.9.8 scale

scale a numeric named vector with scaling for ode parameters of the system. The names must correspond to the parameter identifiers in the ODE specification. Each of the ODE variables will be divided by the scaling factor. For example scale=c(center=2) will divide the center ODE variable by 2.

9.9.9 amountUnits

amountUnits This supplies the dose units of a data frame supplied instead of an event table. This is for importing the data as an RxODE event table.

9.9.10 timeUnits

timeUnits This supplies the time units of a data frame supplied instead of an event table. This is for importing the data as an RxODE event table.

9.9.11 theta

theta A vector of parameters that will be named THETA \ [#\] and added to parameters

9.9.12 eta

eta A vector of parameters that will be named ETA\[#\] and added to parameters

9.9.13 from

from When there is no observations in the event table, start observations at this value. By default this is zero.

9.9.14 to

to When there is no observations in the event table, end observations at this value. By default this is 24 + maximum dose time.

9.9.15 length.out

length.out The number of observations to create if there isn't any observations in the event table. By default this is 200.

9.9.16 by

by When there are no observations in the event table, this is the amount to increment for the observations between from and to.

9.9.17 warnIdSort

warnIdSort Warn if the ID is not present and RxODE assumes the order of the parameters/iCov are the same as the order of the parameters in the input dataset.

9.9.18 warnDrop

warnDrop Warn if column(s) were supposed to be dropped, but were not present.

9.10 Internal RxODE options

9.10.1 nDisplayProgress

nDisplayProgress An integer indicating the minimum number of c-based solves before a progress bar is shown. By default this is 10,000.

9.10.2 ...

... Other arguments including scaling factors for each compartment. This includes S# = numeric will scale a compartment # by a dividing the compartment amount by the scale factor, like NONMEM.

9.10.3 a

a when using solve(), this is equivalent to the object argument. If you specify object later in the argument list it overwrites this parameter.

9.10.4 b

b when using solve(), this is equivalent to the params argument. If you specify params as a named argument, this overwrites the output

9.10.5 updateObject

updateObject This is an internally used flag to update the RxODE solved object (when supplying an RxODE solved object) as well as returning a new object. You probably should not modify it's FALSE default unless you are willing to have unexpected results.

9.11 Parallel/Threaded Solve

9.11.1 cores

cores Number of cores used in parallel ODE solving. This is equivalent to calling [setRxThreads()]

9.11.2 nCoresRV

nCoresRV Number of cores used for the simulation of the sigma variables. By default this is 1. To reproduce the results you need to run on the same platform with the same number of cores. This is the reason this is set to be one, regardless of what the number of cores are used in threaded ODE solving.

Chapter 10

RxODE output

10.1 Using RxODE data frames

10.1.1 Creating an interactive data frame

RxODE supports returning a solved object that is a modified data-frame. This is done by the predict(), solve(), or rxSolve() methods.

```
library(RxODE)
library(units)
### Setup example model
mod1 <-RxODE({</pre>
 C2 = centr/V2;
 C3 = peri/V3;
 d/dt(depot) =-KA*depot;
 d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
 d/dt(peri) =
                                  Q*C2 - Q*C3;
 d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})
### Seup parameters and initial conditions
theta <-
 c(KA=2.94E-01, CL=1.86E+01, V2=4.02E+01, # central
   Q=1.05E+01, V3=2.97E+02, # peripheral
   Kin=1, Kout=1, EC50=200)
                                         # effects
inits <- c(eff=1)</pre>
```

```
### Setup dosing event information
ev <- eventTable(amount.units="mg", time.units="hours") %>%
 add.dosing(dose=10000, nbr.doses=10, dosing.interval=12) %>%
 add.dosing(dose=20000, nbr.doses=5, start.time=120,
         dosing.interval=24) %>%
 add.sampling(0:240);
### Now solve
x <- predict(mod1, theta, ev, inits)
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): ------
     V2 V3 KA CL Q Kin Kout
#>
#> 40.200 297.000 0.294 18.600 10.500 1.000 1.000 200.000
#> -- Initial Conditions ($inits): ------
#> depot centr peri eff
#> 0 0 0 1
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
   time C2 C3 depot centr peri
    [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#>
#> 1
     0 0 0 10000 0 0 1
     1 44.4 0.920 7453. 1784. 273. 1.08
#> 2
#> 3
      2 54.9 2.67 5554. 2206. 794. 1.18
#> 4 3 51.9 4.46 4140. 2087. 1324. 1.23
5 36.5 7.18 2299. 1467. 2132. 1.21
#> 6
#> # ... with 235 more rows
or
x <- solve(mod1, theta, ev, inits)
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
#>
     V2 V3 KA CL
                             Q
                                  Kin
                                        Kout EC50
#> 40.200 297.000  0.294  18.600  10.500  1.000  1.000  200.000
#> -- Initial Conditions ($inits): ------
#> depot centr peri eff
```

```
#> -- First part of data (object): ------
#> # A tibble: 241 x 7
    time C2 C3 depot centr peri eff
     [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#>
      0 0 0 10000
                      0
                           0 1
#> 2
     1 44.4 0.920 7453. 1784. 273. 1.08
     2 54.9 2.67 5554. 2206. 794. 1.18
#> 3
     3 51.9 4.46 4140. 2087. 1324. 1.23
#> 4
#> 5 4 44.5 5.98 3085. 1789. 1776. 1.23
#> 6 5 36.5 7.18 2299. 1467. 2132. 1.21
#> # ... with 235 more rows
Or with mattigr
x <- mod1 %>% solve(theta, ev, inits)
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
                      CL Q
     V2
         V3 KA
                                  Kin Kout
#> 40.200 297.000 0.294 18.600 10.500 1.000 1.000 200.000
#> -- Initial Conditions ($inits): ------
#> depot centr peri eff
     0 0 0
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
    time C2 C3 depot centr peri
     [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#>
#> 1
     0 0 0 10000
                      0 0 1
      1 44.4 0.920 7453. 1784. 273. 1.08
#> 3
     2 54.9 2.67 5554. 2206. 794. 1.18
     3 51.9 4.46 4140. 2087. 1324. 1.23
      4 44.5 5.98
#> 5
                  3085. 1789. 1776. 1.23
                 2299. 1467. 2132. 1.21
      5 36.5 7.18
#> # ... with 235 more rows
#> ______
```

10.1.2 RxODE solved object properties

10.1.3 Using the solved object as a simple data frame

The solved object acts as a data.frame or tbl that can be filtered by dpylr. For example you could filter it easily.

```
library(dplyr)
#> Attaching package: 'dplyr'
#> The following objects are masked from 'package:stats':
#>
#>
      filter, lag
#> The following objects are masked from 'package:base':
#>
#>
       intersect, setdiff, setequal, union
### You can drop units for comparisons and filtering
x <- mod1 %>% solve(theta,ev,inits) %>%
    drop_units %>% filter(time <= 3) %>% as.tbl
#> Warning: `as.tbl()` was deprecated in dplyr 1.0.0.
#> Please use `tibble::as_tibble()` instead.
### or keep them and compare with the proper units.
x <- mod1 %>% solve(theta, ev, inits) %>%
    filter(time <= set_units(3, hr)) %>% as.tbl
#> # A tibble: 4 x 7
     time C2 C3 depot centr peri
       [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
           0 0
#> 1
        0
                     10000
                                0
                                     0
        1 44.4 0.920 7453. 1784. 273. 1.08
#> 3
       2 54.9 2.67 5554. 2206. 794. 1.18
#> 4
       3 51.9 4.46 4140. 2087. 1324. 1.23
```

10.2 Updating the data-set interactively

However it isn't just a simple data object. You can use the solved object to update parameters on the fly, or even change the sampling time.

First we need to recreate the original solved system:

```
x <- mod1 %>% solve(theta,ev,inits);
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
         V3
                            Q Kin Kout
#>
      V2
                  ΚA
                         CL
               0.294 18.600 10.500 1.000
                                        1.000 200.000
#> 40.200 297.000
#> -- Initial Conditions ($inits): ------
#> depot centr peri
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
    time
         C2 C3 depot centr peri eff
     [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#>
                  10000
#> 1
      0 0 0
                          0
                              0 1
      1 44.4 0.920 7453. 1784. 273. 1.08
#> 2
#> 3
      2 54.9 2.67
                  5554. 2206. 794. 1.18
#> 4
      3 51.9 4.46
                  4140. 2087. 1324. 1.23
#> 5
      4 44.5 5.98
                  3085. 1789. 1776. 1.23
#> 6 5 36.5 7.18 2299. 1467. 2132. 1.21
#> # ... with 235 more rows
```

10.2.1 Modifying initial conditions

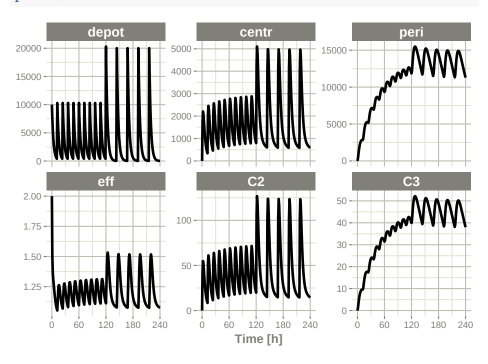
To examine or change initial conditions, you can use the syntax cmt.0, cmt0, or cmt_0. In the case of the eff compartment defined by the model, this is:

```
x$eff0
#> [1] 1
```

which shows the initial condition of the effect compartment. If you wished to change this initial condition to 2, this can be done easily by:

```
#> -- Initial Conditions ($inits): -----
#> depot centr peri
                    eff
                      2
           0
                 0
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
     time
            C2
                  C3
                     depot centr peri
                                        eff
#>
      [h] <dbl> <dbl>
                     <dbl> <dbl> <dbl> <dbl> <
        0
           0
               0
                     10000
                              0
                                   0
                                       2
#> 1
#> 2
        1
          44.4 0.920 7453. 1784.
                                 273. 1.50
                     5554. 2206. 794.
                                      1.37
        2
          54.9 2.67
        3 51.9 4.46
                     4140. 2087. 1324. 1.31
#> 5
        4 44.5 5.98
                     3085. 1789. 1776. 1.27
          36.5 7.18
                     2299. 1467. 2132. 1.23
#> # ... with 235 more rows
```

plot(x)



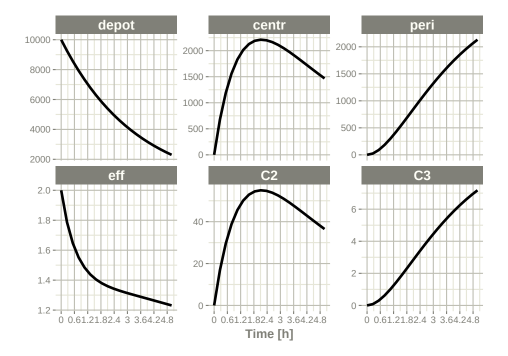
10.2.2 Modifying observation times for RxODE

Notice that the initial effect is now 2.

You can also change the sampling times easily by this method by changing t or time. For example:

```
x$t <- seq(0,5,length.out=20)
print(x)</pre>
```

```
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
\#> V2 V3 KA CL Q Kin Kout EC50
#> 40.200 297.000 0.294 18.600 10.500 1.000 1.000 200.000
#> -- Initial Conditions ($inits): ------
#> depot centr peri eff
#> 0 0 0 2
#> -- First part of data (object): -----
#> # A tibble: 20 x 7
       time C2 C3 depot centr peri eff
#>
        [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#> 1 0.0000000 0 0 10000 0 0
#> 2 0.2631579 16.8 0.0817 9255. 677. 24.3 1.79
#> 3 0.5263158 29.5 0.299 8566. 1187. 88.7 1.65
#> 4 0.7894737 38.9 0.615 7929. 1562. 183. 1.55
#> 5 1.0526316 45.5 1.00 7338. 1830. 298. 1.49
#> 6 1.3157895 50.1 1.44 6792. 2013. 427. 1.44
#> # ... with 14 more rows
```



10.2.3 Modifying simulation parameters

You can also access or change parameters by the \$ operator. For example, accessing KA can be done by:

```
x$KA
```

#> [1] 0.294

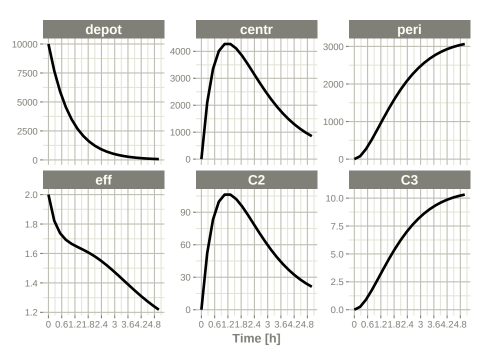
And you may change it by assigning it to a new value.

```
x$KA <- 1
print(x)
```

```
Solved RxODE object _____
#> -- Parameters ($params): --
#>
    ٧2
         VЗ
              KA
                   CL
                            Kin Kout EC50
  40.2 297.0
              1.0
                 18.6 10.5
                            1.0
                                 1.0 200.0
#> -- Initial Conditions ($inits): ------
                  eff
#> depot centr peri
     0
#> -- First part of data (object): ------
```

```
#> # A tibble: 20 x 7
#>
          time
                  C2
                         СЗ
                            depot centr
                                           peri
                                                   eff
#>
           [h] <dbl> <dbl>
                             <dbl> <dbl>
                                          <dbl> <dbl>
#> 1 0.0000000
                 0
                     0
                            10000
                                      0
                                            0
                                                  2
#> 2 0.2631579 52.2 0.261
                             7686. 2098.
                                           77.6
                                                 1.82
#> 3 0.5263158
                83.3 0.900
                            5908. 3348.
                                          267.
                                                  1.74
#> 4 0.7894737
                99.8 1.75
                             4541. 4010.
                                          519.
                                                  1.69
#> 5 1.0526316 106.
                     2.69
                             3490. 4273.
                                          800.
                                                  1.67
#> 6 1.3157895 106. 3.66
                             2683. 4272. 1086.
                                                  1.64
#> # ... with 14 more rows
```

plot(x)



You can access/change all the parameters, initialization(s) or events with the \$params, \$inits, \$events accessor syntax, similar to what is used above.

This syntax makes it easy to update and explore the effect of various parameters on the solved object.

Chapter 11

Simulation

11.1 Single Subject solving

Originally, RxODE was only created to solve ODEs for one individual. That is a single system without any changes in individual parameters.

Of course this is still supported, the classic examples are found in RxODE intro.

This article discusses the differences between multiple subject and single subject solving. There are three differences:

- Single solving does not solve each ID in parallel
- Single solving lacks the id column in parameters(\$params) as well as in the actual dataset.
- Single solving allows parameter exploration easier because each parameter can be modified. With multiple subject solves, you have to make sure to update each individual parameter.

The first obvious difference is in speed; With multiple subjects you can run each subject ID in parallel. For more information and examples of the speed gains with multiple subject solving see the Speeding up RxODE vignette.

The next difference is the amount of information output in the final data.

Taking the 2 compartment indirect response model originally in the tutorial:

```
library(RxODE)
mod1 <-RxODE({
    KA=2.94E-01
    CL=1.86E+01
    V2=4.02E+01
```

```
Q=1.05E+01
   V3=2.97E+02
   Kin=1
   Kout=1
   EC50=200
   C2 = centr/V2
   C3 = peri/V3
   d/dt(depot) =-KA*depot
   d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3
   d/dt(peri) = Q*C2 - Q*C3
   d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff
   eff(0) = 1
})
et <- et(amount.units='mg', time.units='hours') %>%
   et(dose=10000, addl=9, ii=12) %>%
   et(amt=20000, nbr.doses=5, start.time=120, dosing.interval=24) %>%
   et(0:240) # sampling
```

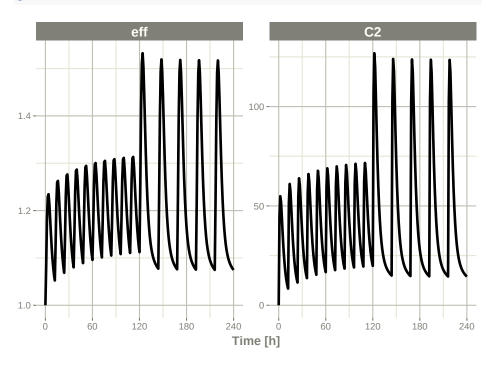
Now a simple solve

```
x <- rxSolve(mod1, et)
#> _____ Solved RxODE object _____
#> -- Parameters (x$params): ------
    KA CL V2 Q V3 Kin Kout EC50
#>
#> 0.294 18.600 40.200 10.500 297.000 1.000 1.000 200.000
#> -- Initial Conditions (x$inits): ------
#> depot centr peri eff
#> 0 0 0
                1
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
  time C2 C3 depot centr peri
#>
#>
    [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#> 1 0 0 0 10000 0 0 1
     1 44.4 0.920 7453. 1784. 273. 1.08
#> 3
     2 54.9 2.67 5554. 2206. 794. 1.18
#> 4 3 51.9 4.46 4140. 2087. 1324. 1.23
#> 5 4 44.5 5.98 3085. 1789. 1776. 1.23
#> 6
     5 36.5 7.18 2299. 1467. 2132. 1.21
#> # ... with 235 more rows
```

print(x)

```
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
                     Q V3
  KA CL V2
                                  Kin
                                       Kout EC50
#> 0.294 18.600 40.200 10.500 297.000 1.000 1.000 200.000
#> -- Initial Conditions ($inits): ------
#> depot centr peri eff
#> 0 0 0 1
#> -- First part of data (object): -----
#> # A tibble: 241 x 7
    time C2 C3 depot centr peri eff
    [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#>
    0 0 0 10000
#> 1
                      0 0 1
     1 44.4 0.920 7453. 1784. 273. 1.08
     2 54.9 2.67 5554. 2206. 794. 1.18
#> 3
#> 4
     3 51.9 4.46 4140. 2087. 1324. 1.23
#> 5
    4 44.5 5.98 3085. 1789. 1776. 1.23
    5 36.5 7.18 2299. 1467. 2132. 1.21
#> # ... with 235 more rows
```

plot(x, C2, eff)



To better see the differences between the single solve, you can solve for 2 individuals

```
x2 <- rxSolve(mod1, et %>% et(id=1:2), params=data.frame(CL=c(18.6, 7.6)))
```

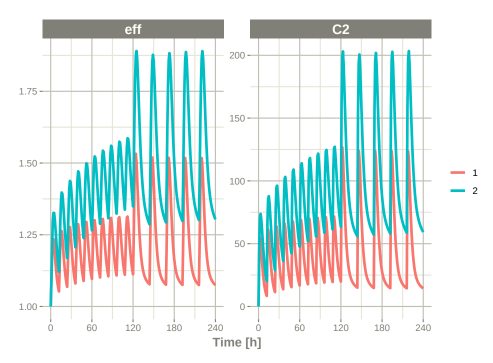
```
#> Warning: 'ID' missing in 'parameters' dataset
```

#> individual parameters are assumed to have the same order as the event dataset

print(x2)

```
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
#> # A tibble: 2 x 9
#> id KA CL V2 Q V3 Kin Kout EC50
#> <fct> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> 
1
                                1
       0.294 7.6 40.2 10.5 297
                              1
                                  1
#> -- Initial Conditions ($inits): ------
#> depot centr peri eff
    0
        0
          0
#> -- First part of data (object): -----
#> # A tibble: 482 x 8
#>
     id time C2 C3 depot centr peri eff
   <int> [h] <dbl> <dbl> <dbl> <dbl> <dbl> <dbl><</pre>
#>
#> 1
                          0 0 1
       0 0 0 10000
     1
         1 44.4 0.920 7453. 1784. 273. 1.08
#> 3
     1 2 54.9 2.67 5554. 2206. 794. 1.18
#> 4
      1 3 51.9 4.46
                   4140. 2087. 1324. 1.23
#> 5
     1 4 44.5 5.98 3085. 1789. 1776. 1.23
#> 6
     1 5 36.5 7.18 2299. 1467. 2132. 1.21
#> # ... with 476 more rows
#> _____
```

```
plot(x2, C2, eff)
```



By observing the two solves, you can see:

• A multiple subject solve contains the id column both in the data frame and then data frame of parameters for each subject.

The last feature that is not as obvious, modifying the individual parameters. For single subject data, you can modify the RxODE data frame changing initial conditions and parameter values as if they were part of the data frame, as described in the RxODE Data Frames.

For multiple subject solving, this feature still works, but requires care when supplying each individual's parameter value, otherwise you may change the solve and drop parameter for key individuals.

11.1.1 Summary of Single solve vs Multiple subject solving

Feature	Single Subject Solve	Multiple Subject Solve
Parallel	None	Each Subject
\$params	data.frame with one	data.frame with one parameter per
	parameter value	subject (w/ID column)
solved	Can modify individual	Have to modify all the parameters to
data	parameters with \$ syntax	update solved object

11.2 Population Simulations with RxODE

11.2.1 Simulation of Variability with RxODE

In pharmacometrics the nonlinear-mixed effect modeling software (like nlmixr) characterizes the between-subject variability. With this between subject variability you can simulate new subjects.

Assuming that you have a 2-compartment, indirect response model, you can set create an RxODE model describing this system below:

11.2.1.1 Setting up the RxODE model

11.2.1.2 Adding the parameter estimates

The next step is to get the parameters into R so that you can start the simulation:

```
theta <- c(KA=2.94E-01, TCl=1.86E+01, V2=4.02E+01, # central Q=1.05E+01, V3=2.97E+02, # peripheral Kin=1, Kout=1, EC50=200, prop.err=0) # effects
```

In this case, I use lotri to specify the omega since it uses similar lower-triangular matrix specification as nlmixr (also similar to NONMEM):

```
### the column names of the omega matrix need to match the parameters specified by RxO omega <- lotri(eta.Cl ~ 0.4^2) omega
```

```
#> eta.Cl
#> eta.Cl 0.16
```

11.2.1.3 Simulating

The next step to simulate is to create the dosing regimen for overall simulation:

```
ev <- et(amount.units="mg", time.units="hours") %>%
  et(amt=10000, cmt="centr")
```

If you wish, you can also add sampling times (though now RxODE can fill these in for you):

```
ev <- ev %>% et(0,48, length.out=100)
```

Note the et takes similar arguments as seq when adding sampling times. There are more methods to adding sampling times and events to make complex dosing regimens (See the event vignette). This includes ways to add variability to the both the sampling and dosing times).

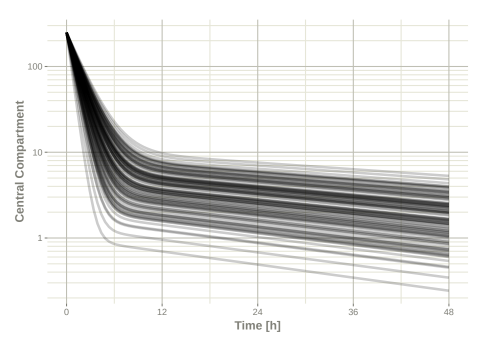
Once this is complete you can simulate using the rxSolve routine:

```
sim <- rxSolve(mod,theta,ev,omega=omega,nSub=100)</pre>
```

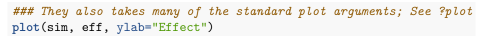
To quickly look and customize your simulation you use the default plot routine. Since this is an RxODE object, it will create a ggplot2 object that you can modify as you wish. The extra parameter to the plot tells RxODE/R what piece of information you are interested in plotting. In this case, we are interested in looking at the derived parameter C2:

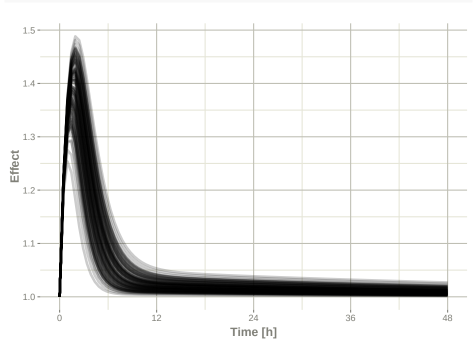
11.2.1.4 Checking the simulation with plot

```
library(ggplot2)
### The plots from RxODE are ggplots so they can be modified with
### standard ggplot commands.
plot(sim, C2, log="y") +
    ylab("Central Compartment")
```



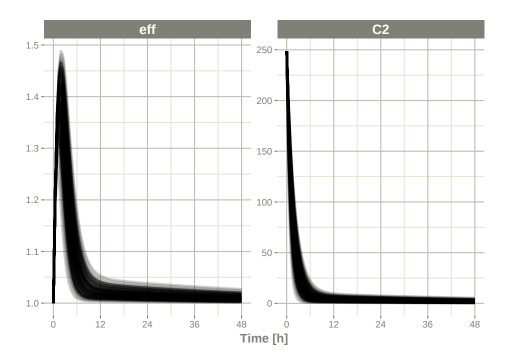
Of course this additional parameter could also be a state value, like eff:





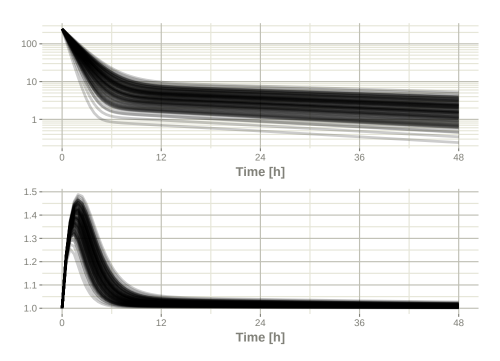
Or you could even look at the two side-by-side:

```
plot(sim, C2, eff)
```



Or stack them with ${\tt patchwork}$

```
library(patchwork)
plot(sim, C2, log="y") / plot(sim, eff)
```



11.2.1.5 Processing the data to create summary plots

Usually in pharmacometric simulations it is not enough to simply simulate the system. We have to do something easier to digest, like look at the central and extreme tendencies of the simulation.

Since the RxODE solve object is a type of data frame

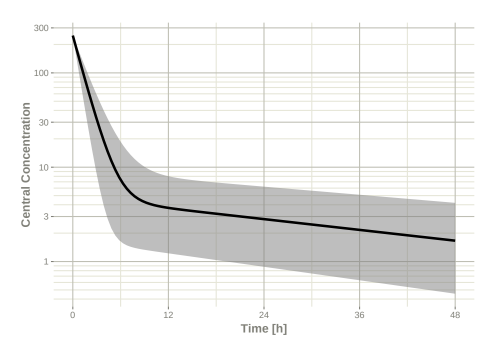
It is now straightforward to perform calculations and generate plots with the simulated data. You can

Below, the 5th, 50th, and 95th percentiles of the simulated data are plotted.

```
confint(sim, "C2", level=0.95) %>%
   plot(ylab="Central Concentration", log="y")
```

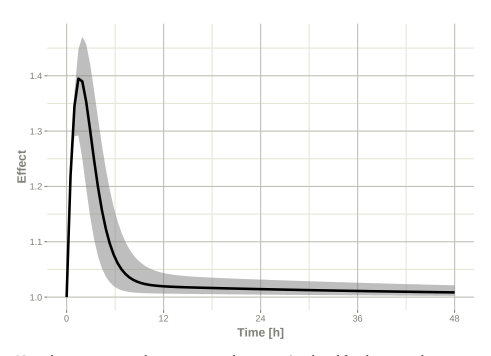
#> ! in order to put confidence bands around the intervals, you need at least 2500 sim

```
#> summarizing data...done
```



! in order to put confidence bands around the intervals, you need at least 2500 simulations

#> summarizing data...done



Note that you can see the parameters that were simulated for the example

```
head(sim$param)
              V2 prop.err V3 TCl
                                       eta.Cl
                                                 ΚA
                                                        Q Kin Kout EC50
#> 1
          1 40.2
                                    0.2368417 0.294 10.5
                                                                    200
                        0 297 18.6
                                                            1
                                                                 1
#> 2
          2 40.2
                        0 297 18.6
                                    0.5454099 0.294 10.5
                                                            1
                                                                 1
                                                                    200
#> 3
          3 40.2
                        0 297 18.6 0.1828379 0.294 10.5
                                                                 1
                                                                    200
#> 4
          4 40.2
                        0 297 18.6 -0.2237885 0.294 10.5
                                                            1
                                                                1
                                                                    200
#> 5
          5 40.2
                        0 297 18.6
                                   0.4640872 0.294 10.5
                                                                 1
                                                                    200
                                                            1
#> 6
          6 40.2
                        0 297 18.6 -0.2748536 0.294 10.5
                                                                    200
```

11.2.1.6 Simulation of unexplained variability (sigma)

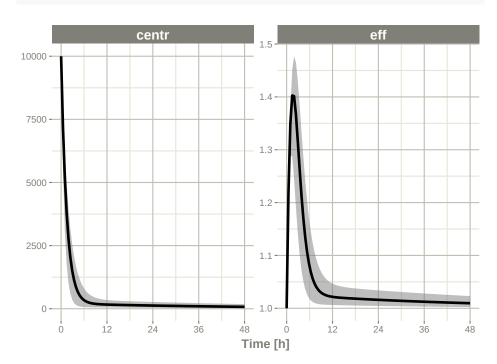
In addition to conveniently simulating between subject variability, you can also easily simulate unexplained variability.

```
mod <- RxODE({
  eff(0) = 1
  C2 = centr/V2;
  C3 = peri/V3;
  CL = TCl*exp(eta.Cl) ## This is coded as a variable in the model
  d/dt(depot) =-KA*depot;</pre>
```

#> ! in order to put confidence bands around the intervals, you need at least 2500 simulations

#> summarizing data...done

plot(s)



11.2.1.7 Simulation of Individuals

Sometimes you may want to match the dosing and observations of individuals in a clinical trial. To do this you will have to create a data.frame using the RxODE event specification as well as an ID column to indicate an individual. The RxODE event vignette talks more about how these datasets should be created.

```
#>
      id
                 time
                                         ср
#> 1
      1 0.000000 [h] 1.0563542 11329.59098
      1 5.333333 [h] 1.4003578
                                  376.07820
#> 3
      1 10.666667 [h] 0.0510544
                                  117.09167
#> 4
      1 16.000000 [h] 1.4589483
                                  141.30089
#> 5
      1 21.333333 [h] 1.1416624
                                   84.85403
      1 26.666667 [h] 1.2504412
                                   95.93320
#> 7
      1 32.000000 [h] 0.9425509
                                  144.84771
      1 37.333333 [h] 1.5173332
#> 8
                                  148.73731
      1 42.666667 [h] 1.2391798
#> 9
                                   60.69626
#> 10 1 48.000000 [h] 1.3173971
                                   53.60546
#> 11 2 0.000000 [h] 0.7351683 5471.03043
#> 12 2 6.857143 [h] 0.7138482
                                  109.19130
#> 13 2 13.714286 [h] 1.2041123
                                  137.81498
#> 14 2 20.571429 [h] 1.1766657
                                   81.08167
#> 15 2 27.428571 [h] 1.7274978
                                   57.74205
#> 16 2 34.285714 [h] 0.4546936
                                   60.74535
#> 17 2 41.142857 [h] 0.7159257
                                   44.59950
#> 18 2 48.000000 [h] 1.3206859
                                   42.03860
```

11.3 Simulation of Clinical Trials

By either using a simple single event table, or data from a clinical trial as described above, a complete clinical trial simulation can be performed.

Typically in clinical trial simulations you want to account for the uncertainty in the fixed parameter estimates, and even the uncertainty in both your between subject variability as well as the unexplained variability.

RxODE allows you to account for these uncertainties by simulating multiple virtual "studies," specified by the parameter nStud. Each of these studies samples a realization of fixed effect parameters and covariance matrices for the between subject variability(omega) and unexplained variabilities (sigma). Depending on the information you have from the models, there are a few strategies for simulating a realization of the omega and sigma matrices.

The first strategy occurs when either there is not any standard errors for standard deviations (or related parameters), or there is a modeled correlation in the model you are simulating from. In that case the suggested strategy is to use the inverse Wishart (parameterized to scale to the conjugate prior)/scaled inverse chi distribution. this approach uses a single parameter to inform the variability of the covariance matrix sampled (the degrees of freedom).

The second strategy occurs if you have standard errors on the variance/standard deviation with no modeled correlations in the covariance matrix. In this approach you perform separate simulations for the standard deviations and the correlation matrix. First you simulate the variance/standard deviation components in the thetaMat multivariate normal simulation. After simulation and transformation to standard deviations, a correlation matrix is simulated using the degrees of freedom of your covariance matrix. Combining the simulated standard deviation with the simulated correlation matrix will give a simulated covariance matrix. For smaller dimension covariance matrices (dimension < 10x10) it is recommended you use the lkj distribution to simulate the correlation matrix. For higher dimension covariance matrices it is suggested you use the inverse wishart distribution (transformed to a correlation matrix) for the simulations.

The covariance/variance prior is simulated from RxODEs cvPost() function.

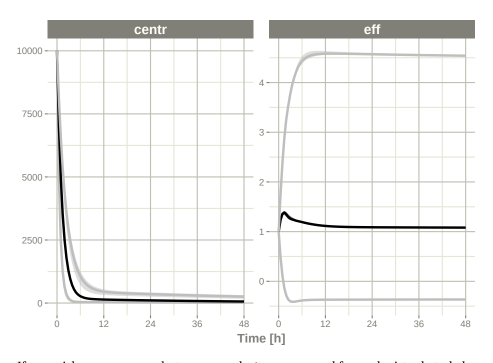
11.3.1 Simulation from inverse Wishart correlations

An example of this simulation is below:

```
### Creating covariance matrix
tmp <- matrix(rnorm(8^2), 8, 8)
tMat <- tcrossprod(tmp, tmp) / (8 ^ 2)
dimnames(tMat) <- list(NULL, names(theta))</pre>
```

#> summarizing data...done

plot(s)



If you wish you can see what omega and sigma was used for each virtual study by accessing them in the solved data object with omega.list and sigma.list:

head(sim\$omega.list)

```
#> [[1]]
#> [,1]
#> [1,] 0.5728809
#>
#> [[2]]
#> [,1]
#> [1,] 0.3465021
#>
```

#>

#>

#>

#> [[6]]

[,1]

[,1] [,2]

#> [1,] 0.098080929 -0.006730568 #> [2,] -0.006730568 0.112366768

```
#> [[3]]
#>
             [,1]
#> [1,] 0.1386869
#>
#> [[4]]
             [,1]
#>
#> [1,] 0.1570577
#>
#> [[5]]
#>
             [,1]
#> [1,] 0.1677731
#>
#> [[6]]
#>
             [,1]
#> [1,] 0.3184372
head(sim$sigma.list)
#> [[1]]
#>
              [,1]
                          [,2]
#> [1,] 0.093539238 0.007270049
#> [2,] 0.007270049 0.098648424
#>
#> [[2]]
#>
                [,1]
                             [,2]
#> [1,] 0.109020277 -0.004127612
#> [2,] -0.004127612  0.087054268
#>
#> [[3]]
#>
              [,1]
                         [,2]
#> [1,] 0.10606530 0.01457913
#> [2,] 0.01457913 0.10189653
#>
#> [[4]]
#>
                 [,1]
                               [,2]
#> [1,] 0.1025867133 -0.0007429996
#> [2,] -0.0007429996  0.0962922149
#>
#> [[5]]
```

[,2]

```
#> [1,] 0.1123437 0.0188019
#> [2,] 0.0188019 0.1021367
```

You can also see the parameter realizations from the \$params data frame.

11.3.2 Simulate using variance/standard deviation standard errors

Lets assume we wish to simulate from the nonmem run included in xpose First we setup the model:

```
rx1 <- RxODE({
   cl <- tcl*(1+crcl.cl*(CLCR-65)) * exp(eta.v)
   v <- tv * WT * exp(eta.v)
   ka <- tka * exp(eta.ka)
   ipred <- linCmt()
   obs <- ipred * (1 + prop.sd) + add.sd
})</pre>
```

Next we input the estimated parameters:

```
theta <- c(tcl=2.63E+01, tv=1.35E+00, tka=4.20E+00, tlag=2.08E-01, prop.sd=2.05E-01, add.sd=1.06E-02, crcl.cl=7.17E-03, ## Note that since we are using the separation strategy the ETA variances a eta.cl=7.30E-02, eta.v=3.80E-02, eta.ka=1.91E+00)
```

And also their covariances; To me, the easiest way to create a named covariance matrix is to use lotri():

```
thetaMat <- lotri(
    tcl + tv + tka + tlag + prop.sd + add.sd + crcl.cl + eta.cl + eta.v + eta.ka ~
        c(7.95E-01,
        2.05E-02, 1.92E-03,
        7.22E-02, -8.30E-03, 6.55E-01,
        -3.45E-03, -6.42E-05, 3.22E-03, 2.47E-04,
        8.71E-04, 2.53E-04, -4.71E-03, -5.79E-05, 5.04E-04,
        6.30E-04, -3.17E-06, -6.52E-04, -1.53E-05, -3.14E-05, 1.34E-05,
        -3.30E-04, 5.46E-06, -3.15E-04, 2.46E-06, 3.15E-06, -1.58E-06, 2.88E-06,
        -1.29E-03, -7.97E-05, 1.68E-03, -2.75E-05, -8.26E-05, 1.13E-05, -1.66E-06, 1
        -1.23E-03, -1.27E-05, -1.33E-03, -1.47E-05, -1.03E-04, 1.02E-05, 1.67E-06, 6
        7.69E-02, -7.23E-03, 3.74E-01, 1.79E-03, -2.85E-03, 1.18E-05, -2.54E-04, 1.6

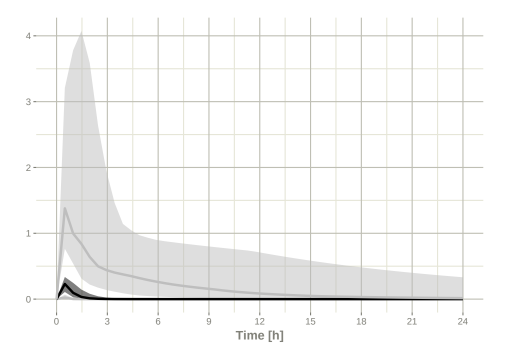
evw <- et(amount.units="mg", time.units="hours") %>%
```

#> Simulation with uncertainty in:

```
et(amt=100) %>%
   ## For this problem we will simulate with sampling windows
   et(list(c(0, 0.5),
      c(0.5, 1),
      c(1, 3),
      c(3, 6),
      c(6, 12))) %>%
   et(id=1:1000)
### From the run we know that:
### total number of observations is: 476
    Total number of individuals: 74
sim <- rxSolve(rx1, theta, evw, nSub=100, nStud=10,
              thetaMat=thetaMat,
              ## Match boundaries of problem
              thetaLower=0,
              sigma=c("prop.sd", "add.sd"), ## Sigmas are standard deviations
              sigmaXform="identity", # default sigma xform="identity"
              omega=c("eta.cl", "eta.v", "eta.ka"), ## etas are variances
              omegaXform="variance", # default omega xform="variance"
              iCov=data.frame(WT=rnorm(1000, 70, 15), CLCR=rnorm(1000, 65, 25)),
              dfSub=74, dfObs=476);
print(sim)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): ------
#> # A tibble: 10,000 x 10
     sim.id id
                 tcl crcl.cl CLCR
                                    eta.v tv WT tka eta.ka
#>
      <int> <fct> <dbl> <dbl> <dbl>
                                    <dbl> <dbl> <dbl> <dbl> <
                                                          <dbl>
                       1.04 54.0 0.907 2.00 71.8 5.69 -0.153
        1 1
#> 1
                 27.0
#> 2
         1 2
                 27.0 1.04 19.7 -0.225 2.00 80.2 5.69 0.249
#> 3
        1 3
                 27.0 1.04 45.7 1.66
                                         2.00 66.3 5.69 0.236
        1 4
                 27.0 1.04 73.9 0.556 2.00 69.4 5.69 -0.156
#> 4
#> 5
        1 5
                 27.0
                      1.04 91.4 0.296 2.00 45.5 5.69 -0.331
#> 6
        1 6
                 27.0
                      1.04 94.9 -0.680 2.00 35.8 5.69 0.372
#> 7
        1 7
               27.0
                      1.04 13.6 -0.327 2.00 95.9 5.69 -0.0760
#> 8
        18
                 27.0 1.04 66.2 0.589 2.00 57.3 5.69 0.688
                 27.0
                        1.04 71.7 -0.611
#> 9
         1 9
                                          2.00 41.0 5.69 0.212
#> 10
         1 10
                 27.0 1.04 76.6 0.00250 2.00 66.5 5.69 0.243
#> # ... with 9,990 more rows
#> -- Initial Conditions ($inits): ------
#> named numeric(0)
#>
```

plot(s)

```
#> * parameters (sim$thetaMat for changes)
#> * omega matrix (sim$omegaList)
#> * sigma matrix (sim$sigmaList)
#> -- First part of data (object): -----
#> # A tibble: 50,000 x 8
#> sim.id
           id
                                        ka ipred
                    time
                           cl v
#> <int> <int>
                     [h] <dbl> <dbl> <dbl> <dbl> <dbl>
       1 1 0.20072222 -696. 356. 4.88
#> 1
                                              NA
             1 0.79938985 -696. 356. 4.88 NA
#> 2
        1
                                                  NA
             1 2.50526151 -696. 356. 4.88 NA NA
#> 3
       1
       1 1 3.38595486 -696. 356. 4.88 NA
1 1 9.28579107 -696. 356. 4.88 NA
1 2 0.04197341 -992 100 - 100
#> 4
             1 3.38595486 -696. 356. 4.88 NA NA
#> 5
                                                   NA
#> 6
              2 0.04197341 -992. 128. 7.30 NA
                                                   NA
#> # ... with 49,994 more rows
#> ______
### Notice that the simulation time-points change for the individual
### If you want the same sampling time-points you can do that as well:
evw <- et(amount.units="mg", time.units="hours") %>%
   et(amt=100) %>%
   et(0, 24, length.out=50) %>%
   et(id=1:100)
sim <- rxSolve(rx1, theta, evw, nSub=100, nStud=10,
              thetaMat=thetaMat,
              ## Match boundaries of problem
              thetaLower=0,
              sigma=c("prop.sd", "add.sd"), ## Sigmas are standard deviations
              sigmaXform="identity", # default sigma xform="identity"
              omega=c("eta.cl", "eta.v", "eta.ka"), ## etas are variances
              omegaXform="variance", # default omega xform="variance"
              iCov=data.frame(WT=rnorm(1000, 70, 15), CLCR=rnorm(1000, 65, 25)),
              dfSub=74, df0bs=476)
s <-sim %>% confint(c("ipred"))
#> summarizing data...done
```



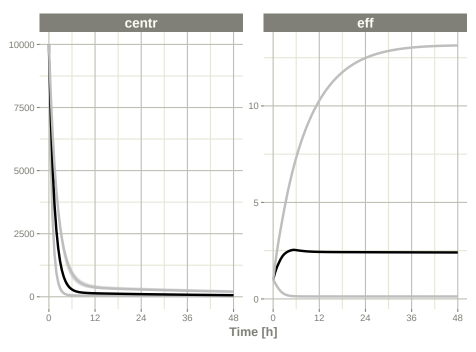
11.3.3 Simulate without uncertainty in omega or sigma parameters

If you do not wish to sample from the prior distributions of either the omega or sigma matrices, you can turn off this feature by specifying the simVariability = FALSE option when solving:

```
mod <- RxODE({</pre>
  eff(0) = 1
  C2 = centr/V2;
  C3 = peri/V3;
  CL = TCl*exp(eta.Cl) ## This is coded as a variable in the model
  d/dt(depot) =-KA*depot;
  d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
  d/dt(peri) =
                                   Q*C2 - Q*C3;
  d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
  e = eff+eff.err
  cp = centr*(1+cp.err)
})
theta <- c(KA=2.94E-01, TCl=1.86E+01, V2=4.02E+01, # central
           Q=1.05E+01, V3=2.97E+02,
                                                    # peripheral
           Kin=1, Kout=1, EC50=200)
                                                    # effects
```

#> summarizing data...done

plot(s)



Note since realizations of omega and sigma were not simulated, $\mathtt{somega.list}$ and $\mathtt{sigma.list}$ both return NULL.

11.3.3.0.1 RxODE multi-threaded solving and simulation RxODE now supports multi-threaded solving on OpenMP supported compilers, including linux and windows. Mac OSX can also be supported By default it uses all your available cores for solving as determined by rxCores(). This may be overkill depending on your system, at a certain point the speed of solving is limited by things other than computing power.

You can also speed up simulation by using the multi-cores to generate random deviates with mvnfast (either mvnfast::rmvn() or mvnfast::rmvt()). This is controlled by the nCoresRV parameter. For example:

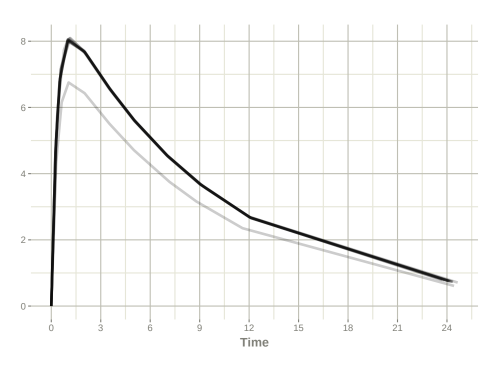
The default for this is 1 core since the result depends on the number of cores and the random seed you use in your simulation as well as the work-load each thread is sharing/architecture. However, you can always speed up this process with more cores if you are sure your collaborators have the same number of cores available to them and have OpenMP thread-capable compile.

11.4 Using prior data for solving

RxODE can use a single subject or multiple subjects with a single event table to solve ODEs. Additionally, RxODE can use an arbitrary data frame with individualized events. For example when using nlmixr, you could use the RxODE/vignettes/theo_sd data frame

```
library(RxODE)
### Load data from nlmixr
d <- qs::qread("RxODE/vignettes/theo_sd.qs")</pre>
### Create RxODE model
theo <- RxODE({
    tka ~ 0.45 # Log Ka
    tcl ~ 1 # Log Cl
    tv ~ 3.45
               # Log V
    eta.ka ~ 0.6
    eta.cl ~ 0.3
    eta.v ~ 0.1
    ka <- exp(tka + eta.ka)
    cl <- exp(tcl + eta.cl)</pre>
    v <- exp(tv + eta.v)
    d/dt(depot) = -ka * depot
    d/dt(center) = ka * depot - cl / v * center
    cp = center / v
})
```

```
### Create parameter dataset
library(dplyr)
parsDf <- tribble(</pre>
 ~ eta.ka, ~ eta.cl, ~ eta.v,
 0.105, -0.487, -0.080,
 0.221, 0.144, 0.021,
 0.368, 0.031, 0.058,
 -0.277, -0.015, -0.007,
 -0.046, -0.155, -0.142,
 -0.382, 0.367, 0.203,
 -0.791, 0.160, 0.047,
 -0.181, 0.168, 0.096,
 1.420, 0.042, 0.012,
 -0.738, -0.391, -0.170,
 0.790, 0.281, 0.146,
 -0.527, -0.126, -0.198) %>%
    mutate(tka = 0.451, tcl = 1.017, tv = 3.449)
### Now solve the dataset
solveData <- rxSolve(theo, parsDf, d)</pre>
plot(solveData, cp)
```



```
print(solveData)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
#> # A tibble: 12 x 1
#>
    id
    <fct>
#>
#> 1 1
#> 2 2
#> 3 3
#> 4 4
#> 5 5
#> 6 6
#> 7 7
#> 8 8
#> 9 9
#> 10 10
#> 11 11
#> 12 12
#> -- Initial Conditions ($inits): ------
#> depot center
     0
#> -- First part of data (object): -----
#> # A tibble: 132 x 8
      id time ka cl v cp depot center
                                 <dbl> <dbl>
#> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <</pre>
      1 0
             2.86 3.67 34.8 0 320.
                                          0
      1 0.25 2.86 3.67 34.8 4.62 157.
#> 3
      1 0.570 2.86 3.67 34.8 7.12 62.8
                                         248.
      1 1.12 2.86 3.67 34.8 8.09 13.0
                                          282.
      1 2.02 2.86 3.67 34.8 7.68 0.996
                                          267.
      1 3.82 2.86 3.67 34.8 6.38 0.00581
#> # ... with 126 more rows
#> ______
### Of course the fasest way to solve if you don't care about the RxODE extra parameters is
solveData <- rxSolve(theo, parsDf, d, returnType="data.frame")</pre>
### solved data
dplyr::as.tbl(solveData)
#> # A tibble: 132 x 8
       id time ka
#>
                    cl v cp depot center
```

```
#>
     <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
                  2.86 3.67 34.8 0
#>
   1
        1 0
                                      3.20e+2
                  2.86 3.67 34.8 4.62 1.57e+2 161.
#> 2
         1 0.25
#>
        1 0.570 2.86 3.67 34.8 7.12 6.28e+1 248.
#> 4
         1 1.12
                  2.86 3.67 34.8 8.09 1.30e+1 282.
#> 5
        1 2.02
                  2.86 3.67
                            34.8 7.68 9.96e-1 267.
#> 6
        1 3.82
                  2.86 3.67
                            34.8 6.38 5.81e-3 222.
#> 7
        1 5.1
                  2.86 3.67 34.8 5.58 1.50e-4 194.
                  2.86 3.67 34.8 4.55 6.02e-7 158.
#> 8
         1 7.03
        1 9.05
#> 9
                  2.86 3.67 34.8 3.68 1.77e-9 128.
#> 10
        1 12.1
                  2.86 3.67 34.8 2.66 9.43e-9
#> # ... with 122 more rows
```

data.table::data.table(solveData)

```
id time
#>
                     ka
                               cl
                                        V
                                                 ср
                                                            depot
                                                                     center
#>
    1: 1 0.00 2.857651 3.669297 34.81332 0.0000000 3.199920e+02
                                                                    0.00000
    2: 1 0.25 2.857651 3.669297 34.81332 4.6240421 1.566295e+02 160.97825
    3: 1 0.57 2.857651 3.669297 34.81332 7.1151647 6.276731e+01 247.70249
    4: 1 1.12 2.857651 3.669297 34.81332 8.0922106 1.303613e+01 281.71670
#>
    5: 1 2.02 2.857651 3.669297 34.81332 7.6837844 9.958446e-01 267.49803
#>
#> ---
#> 128: 12 5.07 2.857651 3.669297 34.81332 5.6044213 1.636210e-04 195.10850
#> 129: 12 7.07 2.857651 3.669297 34.81332 4.5392337 5.385697e-07 158.02579
#> 130: 12 9.03 2.857651 3.669297 34.81332 3.6920276 1.882087e-09 128.53173
#> 131: 12 12.05 2.857651 3.669297 34.81332 2.6855080 8.461424e-09 93.49144
#> 132: 12 24.15 2.857651 3.669297 34.81332 0.7501667 -4.775222e-10 26.11579
```

Chapter 12

Examples

This section is for example models to get you started in common simulation scenarios.

12.1 Prediction only models

Prediction only models are simple to create. You use the RxODE syntax without any ODE systems in them. A very simple example is a one-compartment model.

```
library(RxODE)
mod <- RxODE({
    ipre <- 10 * exp(-ke * t);
})
mod</pre>
```

```
\# RxODE 1.0.5 model named rx_7e31c1ae655db7c06244b075015b3e86 model (ready). \# x$params: ke \# x$lhs: ipre
```

Solving the RxODE models are the same as saving the simple ODE system, but faster of course.

```
et <- et(seq(0,24,length.out=50))
cmt1 <- rxSolve(mod,et,params=c(ke=0.5))
cmt1</pre>
```

#> _____ Solved RxODE object _____

```
#> -- Parameters (x$params): ------
#> ke
#> 0.5
#> -- Initial Conditions (x$inits): ------
#> named numeric(0)
#> -- First part of data (object): -----
#> # A tibble: 50 x 2
#>
    time ipre
   <dbl> <dbl>
#>
#> 1 0
       10
#> 2 0.490 7.83
#> 3 0.980 6.13
#> 4 1.47 4.80
#> 5 1.96 3.75
#> 6 2.45 2.94
#> # ... with 44 more rows
#> _____
```

12.2 Solved compartment models

Solved models are also simple to create. You simply place the linCmt() psuedo-function into your code. The linCmt() function figures out the type of model to use based on the parameter names specified.

Most often, pharmacometric models are parameterized in terms of volume and clearances. Clearances are specified by NONMEM-style names of CL, Q, Q1, Q2, etc. or distributional clearances CLD, CLD2. Volumes are specified by Central (VC or V), Peripheral/Tissue (VP, VT). While more translations are available, some example translations are below:

```
#>
#> Attaching package: 'kableExtra'

#> The following object is masked from 'package:dplyr':
#>
#> group_rows
```

Table 12.1: Clearance Based linCmt() parameterizations

par1	par2	par3	par4	par5	par6	par7	ncmt
ka	cl	q	q2	V	vp	vp2	3
cl	q	q2	v	vp	vp2		3
ka	cl	q	q2	vc	vp	vp2	3

Table 12.1: Clearance Based linCmt() parameterizations (continued)

par1	par2	par3	par4	par5	par6	par7	ncmt
cl	q	q2	vc	vp	vp2		3
ka	cl	q1	q2	V	vp	vp2	3
cl	q1	q2	V	vp	vp2		3
ka	cl	q1	q2	vc	vp	vp2	3
cl	q1	q2	vc	vp	vp2		3
ka	cl	q2	V	vp2			2
cl	q2	V	vp2				2
ka	cl	q2	vc	vp2			2
cl	q2	vc	vp2				2
ka	cl	cld	cld2	V	vp	vp2	3
cl	cld	cld2	V	vp	vp2		3
ka	cl	cld	cld2	vc	vp	vp2	3
cl	cld	cld2	vc	vp	vp2		3
ka	cl	cld2	V	vp2			2
cl	cld2	V	vp2				2
ka	cl	cld2	vc	vp2			2
cl	cld2	vc	vp2				2
ka	cl	q	v	vp2			2
cl	q	V	vp2				2
ka	cl	q	vc	vp2			2
cl	q	vc	vp2	_			2
ka	cl	q1	V	vp2			2
cl	q1	v	vp2				2
ka	cl	q1	vc	vp2			2
cl	q1	vc	vp2				2
ka	cl	cld	V	vp2			2
cl	cld	V	vp2				2
ka	cl	cld	vc	vp2			2
cl	cld	vc	vp2				2
ka	cl	q	q2	V	v2	v3	3
cl	q	q2	v	v2	v3		3
ka	cl	q	q2	v2	v3	vc	3
cl	q	q2	v2	v3	vc		3
ka	cl	q	q2	v1	v2	v3	3
cl	q	q2	v1	v2	v3		3
ka	cl	q1	q2	V	v2	v3	3
cl	q1	q2	V	v2	v3		3
ka	cl	q1	q2	v2	v3	vc	3
cl	q1	q2	v2	v3	vc		3

Table 12.1: Clearance Based linCmt() parameterizations (continued)

par1	par2	par3	par4	par5	par6	par7	ncmt
ka	cl	q1	q2	v1	v2	v3	3
cl	q1	q2	v1	v2	v3		3
ka	cl	q2	v2	v3			2
cl	q2	v2	v3				2
ka	cl	q2	v	v3			2
cl	q2	v	v3				2
ka	cl	q2	v3	vc			2
cl	q2	v3	vc				2
ka	cl	cld	cld2	v	v2	v3	3
cl	cld	cld2	V	v2	v3		3
ka	cl	cld	cld2	v2	v3	vc	3
cl	cld	cld2	v2	v3	vc		3
ka	cl	cld	cld2	v1	v2	v3	3
cl	cld	cld2	v1	v2	v3		3
ka	cl	cld2	v2	v3			2
cl	cld2	v2	v3				2
ka	cl	cld2	V	v3			2
cl	cld2	V	v3				2
ka	cl	cld2	v3	vc			2
cl	cld2	v3	vc				2
ka	cl	q	v2	v3			2
cl	q	v2	v3				2
ka	cl	q1	v2	v3			2
cl	q1	v2	v3				2
ka	cl	cld	v2	v3			2
cl	cld	v2	v3				2
ka	cl	q	V	v3			2
cl	q	V	v3				2
ka	cl	q	v3	vc			2
cl	q	v3	vc				2
ka	cl	q1	V	v3			2
cl	q1	V	v3				2
ka	cl	q1	v3	vc			2
cl	q1	v3	vc				2
ka	cl	cld	V	v3			2
cl	cld	V	v3				2
ka	cl	cld	v3	vc			2
cl	cld	v3	vc				2
ka	cl	q	q2	V	vt	vt2	3

Table 12.1: Clearance Based linCmt() parameterizations (continued)

			4				
par1	par2	par3	par4	par5	par6	par7	ncmt
cl	q	q2	v	vt	vt2		3
ka	cl	q	q2	vc	vt	vt2	3
cl	q	q2	vc	vt	vt2		3
ka	cl	q1	q2	V	vt	vt2	3
cl	q1	q2	v	vt	vt2		3
ka	cl	q1	q2	vc	vt	vt2	3
cl	q1	q2	vc	vt	vt2		3
ka	cl	q2	V	vt2			2
cl	q2	V	vt2				2
ka	cl	q2	vc	vt2			2
cl	q2	vc	vt2				2
ka	cl	cld	cld2	V	vt	vt2	3
cl	cld	cld2	v	vt	vt2		3
ka	cl	cld	cld2	vc	vt	vt2	3
cl	cld	cld2	vc	vt	vt2		3
ka	cl	cld2	v	vt2			2
cl	cld2	v	vt2				2
ka	cl	cld2	vc	vt2			2
cl	cld2	vc	vt2				2
ka	cl	q	V	vt2			2
cl	q	v	vt2				2
ka	cl	q	vc	vt2			2
cl	q	vc	vt2				2
ka	cl	q1	V	vt2			2
cl	q1	v	vt2				2
ka	cl	q1	vc	vt2			2
cl	q1	vc	vt2				2
ka	cl	cld	V	vt2			2
cl	cld	\mathbf{v}	vt2				2
ka	cl	cld	vc	vt2			2
cl	cld	vc	vt2				2
ka	cl	q2	V	v2			2
cl	q2	v	v2				2
ka	cl	q2	v2	vc			2
cl	q2	v2	vc				2
ka	cl	q2	v1	v2			2
cl	q2	v1	v2				2
ka	cl	q2	v	vp			2
cl	q2	v	vp	_			2

Table 12.1: Clearance Based linCmt() parameterizations (continued)

1							
parl	par2	par3	par4	pars	par6	par /	ncmt
ka	cl	q2	vc	vp			2
cl	q2	vc	vp				2
ka	cl	q2	V	vt			2
cl	q2	V	vt				2
ka	cl	q2	vc	vt			2
cl	q2	vc	vt				2
ka	cl	q2	V	vss			2
cl	q2	V	VSS				2
ka	cl	q2	vc	vss			2
cl	q2	vc	VSS				2
ka	cl	cld2	V	v2			2
cl	cld2	v	v2				2
ka	cl	cld2	v2	vc			2
cl	cld2	v2	vc				2
ka	cl	cld2	v1	v2			2
cl	cld2	v1	v2				2
ka	cl	cld2	V	vp			2
cl	cld2	v	vp				2
ka	cl	cld2	vc	vp			2
cl	cld2	vc	vp				2
ka	cl	cld2	v	vt			2
cl	cld2	v	vt				2
ka	cl	cld2	vc	vt			2
cl	cld2	vc	vt				2
ka	cl	cld2	V	vss			2
cl	cld2	v	vss				2
ka	cl	cld2	vc	vss			2
cl	cld2	vc	vss				2
ka	cl	q	V	v2			2
cl	q	V	v2				2
ka	cl	q	v2	vc			2
cl	q	v2	vc				2
ka	cl	q	v1	v2			2
cl	q	v1	v2				2
ka	cl	q1	V	v2			2
cl	q1	V	v2				2
ka	cl	q1	v2	vc			2
cl	q1	v2	vc				2
ka	cl	q1	v1	v2			2

Table 12.1: Clearance Based linCmt() parameterizations (continued)

par1	par2	par3	par4	par5	par6	par7	ncmt
cl	q1	vl	v2				2
ka	cl	cld	V	v2			2
cl	cld	V	v2				2
ka	cl	cld	v2	vc			2
cl	cld	v2	vc				2
ka	cl	cld	v1	v2			2
cl	cld	v1	v2				2
ka	cl	v2					1
cl	v2						1
ka	cl	q	V	vp			2
cl	q	V	vp				2
ka	cl	q	vc	vp			2
cl	q	vc	vp				2
ka	cl	q1	V	vp			2
cl	q1	V	vp				2
ka	cl	q1	vc	vp			2
cl	q1	vc	vp				2
ka	cl	cld	v	vp			2
cl	cld	V	vp				2
ka	cl	cld	vc	vp			2
cl	cld	vc	vp				2
ka	cl	q	V	vt			2
cl	q	V	vt				2
ka	cl	q	vc	vt			2
cl	q	vc	vt				2
ka	cl	q1	V	vt			2
cl	q1	\mathbf{v}	vt				2
ka	cl	q1	vc	vt			2
cl	q1	vc	vt				2
ka	cl	cld	V	vt			2
cl	cld	V	vt				2
ka	cl	cld	vc	vt			2
cl	cld	vc	vt				2
ka	cl	q	V	VSS			2
cl	q	V	VSS				2
ka	cl	q	vc	VSS			2
cl	q	vc	VSS				2
ka	cl	q1	V	VSS			2
cl	q1	\mathbf{v}	VSS				2

Table 12.1: Clearance Based linCmt() parameterizations (continued)

par1	par2	par3	par4	par5	par6	par7	ncmt
ka	cl	ql	vc	VSS			2
cl	q1	vc	vss				2
ka	cl	cld	v	vss			2
cl	cld	V	vss				2
ka	cl	cld	vc	vss			2
cl	cld	vc	vss				2
ka	cl	V					1
cl	v						1
ka	cl	vc					1
cl	vc						1
ka	cl	v1					1
cl	v1						1

Another popular parameterization is in terms of micro-constants. RxODE assumes compartment 1 is the central compartment. The elimination constant would be specified by K, Ke or Ke1. Some example translations are below:

Table 12.2: Kel Based linCmt() parameterizations

par1	par2	par3	par4	par5	par6	par7	ncmt
ka	v	k	k12	k21	k13	k31	3
v	k	k12	k21	k13	k31		3
ka	vc	k	k12	k21	k13	k31	3
vc	k	k12	k21	k13	k31		3
ka	v1	k	k12	k21	k13	k31	3
v1	k	k12	k21	k13	k31		3
ka	V	ke	k12	k21	k13	k31	3
v	ke	k12	k21	k13	k31		3
ka	vc	ke	k12	k21	k13	k31	3
vc	ke	k12	k21	k13	k31		3
ka	v1	ke	k12	k21	k13	k31	3
v1	ke	k12	k21	k13	k31		3
ka	V	kel	k12	k21	k13	k31	3
v	kel	k12	k21	k13	k31		3
ka	vc	kel	k12	k21	k13	k31	3
vc	kel	k12	k21	k13	k31		3
ka	v1	kel	k12	k21	k13	k31	3
v1	kel	k12	k21	k13	k31		3
ka	V	k	k12	k21			2

Table 12.2: Kel Based linCmt() parameterizations (continued)

parl	par2	par3	par4	par5	par6	par7	ncmt
v	k	k12	k21				2
ka	vc	k	k12	k21			2
vc	k	k12	k21				2
ka	v1	k	k12	k21			2
v1	k	k12	k21				2
ka	V	ke	k12	k21			2
\mathbf{v}	ke	k12	k21				2
ka	vc	ke	k12	k21			2
vc	ke	k12	k21				2
ka	v1	ke	k12	k21			2
v1	ke	k12	k21				2
ka	v	kel	k12	k21			2
v	kel	k12	k21				2
ka	vc	kel	k12	k21			2
vc	kel	k12	k21				2
ka	v1	kel	k12	k21			2
v1	kel	k12	k21				2
ka	V	k					1
V	k						1
ka	vc	k					1
vc	k						1
ka	v1	k					1
v1	k						1
ka	v	ke					1
V	ke						1
ka	vc	ke					1
vc	ke						1
ka	v1	ke					1
v1	ke						1
ka	v	kel					1
V	kel						1
ka	vc	kel					1
vc	kel						1
ka	v1	kel					1
v1	kel						1

The last parameterization possible is using alpha and V and/or A/B/C. Some example translations are below:

par1	par2	par3	par4	par5	par6	par7	ncmt
ka	V	alpha	beta	aob			1
v	alpha	beta	aob				1
ka	vc	alpha	beta	aob			1
vc	alpha	beta	aob				1
ka	v1	alpha	beta	aob			1
v1	alpha	beta	aob				1
ka	v	alpha	beta	k21			1
v	alpha	beta	k21				1
ka	vc	alpha	beta	k21			1
vc	alpha	beta	k21				1
ka	v1	alpha	beta	k21			1
v1	alpha	beta	k21				1
ka	V	alpha					2
V	alpha						2
ka	vc	alpha					2
vc	alpha						2
ka	v1	alpha					2
v1	alpha						2
ka	a	alpha	b	beta	c	gamma	3
a	alpha	b	beta	c	gamma		3
ka	a	alpha	b	beta			2
a	alpha	b	beta				2
ka	a	alpha					1
a	alpha						1

Table 12.3: alpha Based linCmt() parameterizations

Once the linCmt() sleuthing is complete, the 1, 2 or 3 compartment model solution is used as the value of linCmt().

The compartments where you can dose in a linear solved system are depot and central when there is an linear absorption constant in the model ka. Without any additional ODEs, these compartments are numbered depot=1 and central=2.

When the absorption constant ka is missing, you may only dose to the central compartment. Without any additional ODEs the compartment number is central=1.

These compartments take the same sort of events that a ODE model can take, and are discussed in the RxODE events vignette.

```
mod <- RxODE({
    ke <- 0.5
```

```
V <- 1
   ipre <- linCmt();</pre>
})
mod
#> RxODE 1.0.5 model named rx_d41c572184c0c5d136339f94b81154e8 model (ready).
#> x$stateExtra: central
#> x$params: ke, V
#> x$lhs: ipre
This then acts as an ODE model; You specify a dose to the depot compartment and
then solve the system:
et <- et(amt=10,time=0,cmt=depot) %>%
   et(seq(0,24,length.out=50))
cmt1 <- rxSolve(mod,et,params=c(ke=0.5))</pre>
cmt1
#> _____ Solved RxODE object _____
#> -- Parameters (x$params): -----
#> ke V
#> 0.5 1.0
#> -- Initial Conditions (x$inits): -----
#> named numeric(0)
#> -- First part of data (object): -----
\#> \# A tibble: 50 x 2
     time ipre
#>
    <dbl> <dbl>
#> 1 0
        10
#> 2 0.490 7.83
#> 3 0.980 6.13
#> 4 1.47 4.80
#> 5 1.96 3.75
#> 6 2.45 2.94
#> # ... with 44 more rows
```

12.3 Mixing Solved Systems and ODEs

In addition to pure ODEs, you may mix solved systems and ODEs. The prior 2-compartment indirect response model can be simplified with a linCmt() function:

```
library(RxODE)
## Setup example model
mod1 <-RxODE({</pre>
   C2 = centr/V2;
   C3 = peri/V3;
    d/dt(depot) =-KA*depot;
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) =
                                     Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
});
## Seup parameters and initial conditions
theta <-
    c(KA=2.94E-01, CL=1.86E+01, V2=4.02E+01, # central
      Q=1.05E+01, V3=2.97E+02, # peripheral
      Kin=1, Kout=1, EC50=200)
                                            # effects
inits <- c(eff=1);</pre>
## Setup dosing event information
ev <- eventTable(amount.units="mg", time.units="hours") %>%
    add.dosing(dose=10000, nbr.doses=10, dosing.interval=12) %>%
    add.dosing(dose=20000, nbr.doses=5, start.time=120,dosing.interval=24) %>%
    add.sampling(0:240);
## Setup a mixed solved/ode system:
mod2 <- RxODE({</pre>
    ## the order of variables do not matter, the type of compartmental
    ## model is determined by the parameters specified.
    C2 = linCmt(KA, CL, V2, Q, V3);
    eff(0) = 1 ## This specifies that the effect compartment starts at 1.
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})
```

This allows the indirect response model above to assign the 2-compartment model to the C2 variable and the used in the indirect response model.

When mixing the solved systems and the ODEs, the solved system's compartment is always the last compartment. This is because the solved system technically isn't a compartment to be solved. Adding the dosing compartment to the end will not interfere with the actual ODE to be solved.

Therefore,in the two-compartment indirect response model, the effect compartment is compartment #1 while the PK dosing compartment for the depot is compartment #2.

This compartment model requires a new event table since the compartment number changed:

```
ev <- eventTable(amount.units='mg', time.units='hours') %>%
    add.dosing(dose=10000, nbr.doses=10, dosing.interval=12,dosing.to=2) %>%
    add.dosing(dose=20000, nbr.doses=5, start.time=120,dosing.interval=24,dosing.to=2) %>%
    add.sampling(0:240);
```

This can be solved with the following command:

```
x <- mod2 %>% solve(theta, ev)
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): -----
   CL V2 Q V3
                          KA
                               Kin Kout EC50
#> 18.600 40.200 10.500 297.000 0.294 1.000 1.000 200.000
#> -- Initial Conditions ($inits): ------
#> eff
#> -- First part of data (object): -----
#> # A tibble: 241 x 3
    time C2 eff
#>
    [h] <dbl> <dbl>
#> 1
    0 249. 1
#> 2
     1 121. 1.35
#> 3
    2 60.3 1.38
#> 4
    3 31.0 1.28
#> 5
   4 17.0 1.18
#> # ... with 235 more rows
```

Note this solving did not require specifying the effect compartment initial condition to be 1. Rather, this is already pre-specified by eff(0)=1.

This can be solved for different initial conditions easily:

```
#> 18.600 40.200 10.500 297.000 0.294 1.000 1.000 200.000
#> -- Initial Conditions ($inits): -----
#> eff
#> -- First part of data (object): -----
#> # A tibble: 241 x 3
   time
          C2
     [h] <dbl> <dbl>
#>
#> 1
     0 249.
#> 2
      1 121. 1.93
#> 3
      2 60.3 1.67
#> 4
      3 31.0 1.41
#> 5
       4 17.0 1.23
#> 6
    5 10.2 1.13
#> # ... with 235 more rows
```

The RxODE detective also does not require you to specify the variables in the linCmt() function if they are already defined in the block. Therefore, the following function will also work to solve the same system.

```
mod3 <- RxODE({</pre>
   KA=2.94E-01;
   CL=1.86E+01;
   V2=4.02E+01;
   Q=1.05E+01;
   V3=2.97E+02;
   Kin=1;
   Kout=1;
   EC50=200:
   ## The linCmt() picks up the variables from above
   C2 = linCmt();
   eff(0) = 1 ## This specifies that the effect compartment starts at 1.
   d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})
x <- mod3 %>% solve(ev)
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): ------
                       Q
                                             Kout EC50
      ΚA
         CL V2
                              V3
                                      Kin
#> 0.294 18.600 40.200 10.500 297.000 1.000 1.000 200.000
#> -- Initial Conditions ($inits): ------
#> eff
```

```
#>
#> -- First part of data (object): -----
#> # A tibble: 241 x 3
    time C2 eff
     [h] <dbl> <dbl>
#>
#> 1
       0 249. 1
#> 2
      1 121. 1.35
      2 60.3 1.38
#> 3
      3 31.0 1.28
#> 4
#> 5
      4 17.0 1.18
    5 10.2 1.11
#> # ... with 235 more rows
```

Note that you do not specify the parameters when solving the system since they are built into the model, but you can override the parameters:

```
x <- mod3 %>% solve(c(KA=10),ev)
print(x)
#> _____ Solved RxODE object _____
#> -- Parameters ($params): ------
    KA CL V2 Q V3 Kin Kout EC50
#> 10.0 18.6 40.2 10.5 297.0 1.0 1.0 200.0
#> -- Initial Conditions ($inits): ------
#> eff
#>
#> -- First part of data (object): -----
#> # A tibble: 241 x 3
    time C2 eff
    [h] <dbl> <dbl>
#>
#> 1
     0 249. 1
#> 2
     1 121.
            1.35
     2 60.3 1.38
#> 3
#> 4
     3 31.0 1.28
#> 5
     4 17.0 1.18
     5 10.2 1.11
#> # ... with 235 more rows
#> ______
```

12.4 Weight based dosing

This is an example model for weight based dosing of daptomycin. Daptomycin is a cyclic lipopeptide antibiotic from fermented *Streptomyces roseosporus*.

There are 3 stages for weight-based dosing simulations: - Create RxODE model - Simulate Covariates - Create event table with weight-based dosing (merged back to covariates)

12.4.1 Creating a 2-compartment model in RxODE

```
library(RxODE)

## Note the time covariate is not included in the simulation
m1 <- RxODE({
    CL ~ (1-0.2*SEX)*(0.807+0.00514*(CRCL-91.2))*exp(eta.cl)
    V1 ~ 4.8*exp(eta.v1)
    Q ~ (3.46+0.0593*(WT-75.1))*exp(eta.q);
    V2 ~ 1.93*(3.13+0.0458*(WT-75.1))*exp(eta.v2)
    A1 ~ centr;
    A2 ~ peri;
    d/dt(centr) ~ - A1*(CL/V1 + Q/V1) + A2*Q/V2;
    d/dt(peri) ~ A1*Q/V1 - A2*Q/V2;
    DV = centr / V1 * (1 + prop.err)
})</pre>
```

12.4.2 Simulating Covariates

This simulation correlates age, sex, and weight. Since we will be using weight based dosing, this needs to be simulated first

```
set.seed(42)
library(dplyr)
nsub=30
### Simulate Weight based on age and gender
AGE<-round(runif(nsub,min=18,max=70))
SEX<-round(runif(nsub,min=0,max=1))
HTm<-round(rnorm(nsub,176.3,0.17*sqrt(4482)),digits=1)
HTf<-round(rnorm(nsub,162.2,0.16*sqrt(4857)),digits=1)
WTm<-round(exp(3.28+1.92*log(HTm/100))*exp(rnorm(nsub,0,0.14)),digits=1)
WTf<-round(exp(3.49+1.45*log(HTf/100))*exp(rnorm(nsub,0,0.17)),digits=1)
WT<-ifelse(SEX==1,WTf,WTm)
CRCL<-round(runif(nsub,30,140))
## id is in lower case to match the event table
cov.df <- tibble(id=seq_along(AGE), AGE=AGE, SEX=SEX, WT=WT, CRCL=CRCL)
print(cov.df)</pre>
```

```
#> # A tibble: 30 x 5
#>
             AGE
                   SEX
                          WT CRCL
        id
      <int> <dbl> <dbl> <dbl> <dbl>
#>
                     1 49.4
#>
   1
         1
              66
                                83
                     1 52.5
              67
                                79
#>
   2
         2
#>
   3
         3
              33
                     0 97.9
                                37
#>
  4
         4
              61
                     1 63.8
                                66
  5
#>
         5
              51
                     0 71.8
                               127
  6
         6
                     1 69.6
#>
              45
                               132
#>
  7
         7
              56
                     0 61
                                73
#> 8
         8
              25
                     0 57.7
                                47
#> 9
         9
              52
                     1 58.7
                                65
#> 10
        10
              55
                     1 73.1
                                64
#> # ... with 20 more rows
```

12.4.3 Creating weight based event table

```
s<-c(0,0.25,0.5,0.75,1,1.5,seq(2,24,by=1))
s <- lapply(s, function(x){.x <- 0.1 * x; c(x - .x, x + .x)})

e <- et() %>%
    ## Specify the id and weight based dosing from covariate data.frame
    ## This requires RxODE XXX
    et(id=cov.df$id, amt=6*cov.df$WT, rate=6 * cov.df$WT) %>%
    ## Sampling is added for each ID
    et(s) %>%
    as.data.frame %>%
    ## Merge the event table with the covarite information
    merge(cov.df, by="id") %>%
    as_tibble
```

```
#> # A tibble: 900 x 12
#>
              low time
                                                            AGE
                                                                         WT CRCL
        id
                         high cmt
                                          amt rate evid
                                                                  SEX
                        <dbl> <chr>
      <int> <dbl> <dbl>
                                         <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
#>
#>
         1 0
                  0
                         0
                               (obs)
                                          NA
                                                             66
                                                                       49.4
                                                                               83
   1
                                                NA
                                                        0
                                                                    1
                                                                       49.4
#>
   2
         1 NA
                  0
                        NA
                               (default)
                                         296.
                                               296.
                                                             66
                                                                    1
                                                                               83
                                                        1
         1 0.225 0.246 0.275 (obs)
#>
  3
                                          NA
                                                NA
                                                        0
                                                             66
                                                                    1
                                                                       49.4
                                                                               83
         1 0.45 0.516 0.55 (obs)
                                          NA
                                                NA
                                                        0
                                                             66
                                                                    1
                                                                       49.4
                                                                               83
                                                                    1 49.4
#> 5
         1 0.675 0.729 0.825 (obs)
                                          NA
                                                NA
                                                        0
                                                             66
                                                                               83
                                                                               83
#> 6
         1 0.9 0.921 1.1
                               (obs)
                                          NA
                                                        0
                                                             66
                                                                    1 49.4
                                                NA
```

```
#> 7
        1 1.35 1.42
                      1.65 (obs)
                                       NA
                                             NA
                                                    0
                                                        66
                                                              1 49.4
                                                                         83
        1 1.8
                 1.82
                       2.2
                             (obs)
                                       NA
                                             NA
                                                    0
                                                        66
                                                              1 49.4
                                                                         83
#> 8
#> 9
                                             NA
                                                    0
                                                        66
                                                             1 49.4
                                                                         83
        1 2.7
                 2.97
                       3.3
                             (obs)
                                       NA
#> 10
                                                        66
                                                             1 49.4
                                                                         83
        1 3.6
                 3.87
                       4.4
                             (obs)
                                       NA
                                             NA
#> # ... with 890 more rows
```

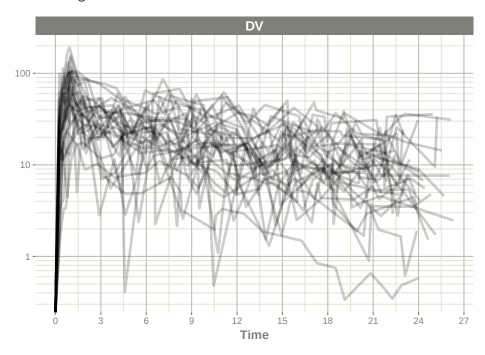
12.4.4 Solving Daptomycin simulation

```
______ Solved RxODE object ______
#> -- Parameters ($params): -----
#> # A tibble: 30 x 5
  id
         eta.cl eta.v1 eta.q eta.v2
#>
    <fct>
         <dbl> <dbl> <dbl>
                           <dbl>
#> 1 1
         #> 2 2
        0.0341 0.406 -0.139 -0.481
#> 3 3
        -0.447 0.0952 -0.185 -0.249
#> 4 4
        -0.988 0.248 -0.131 -0.449
#> 5 5
        0.144 -1.14
                     0.106
                          0.360
        -0.689
#> 6 6
              0.407 -0.193 -0.200
#> 7 7
        -0.426 -0.706 -0.190 -0.234
#> 8 8
        -0.212 0.728
                    0.335 0.0665
#> 9 9
         0.0884 -0.934 0.337 0.154
#> 10 10
        -0.557 1.29
                     0.0163 -0.140
#> # ... with 20 more rows
#> -- Initial Conditions ($inits): -------
#> centr peri
    0
#> -- First part of data (object): -------
#> # A tibble: 900 x 9
     id evid cmt
                  amt time
                          DV SEX
                                    WT CRCL
#> <int> <int> <int> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
#> 1 1 1 296.0 0
                               1 49.4 83
```

```
1
                   NA
                        NA O
                                            1 49.4
                                                       83
        1
                   NA
                        NA
                            0.246 2.32
                                            1
                                               49.4
                                                       83
              0
                   NA
                                                       83
        1
                            0.516 19.6
                                            1 49.4
                   NA
                            0.729 23.2
                                                       83
        1
                        NA
                                            1 49.4
              0
                            0.921 21.1
                                               49.4
                                                       83
#> 6
        1
                   NA
                        NA
#> # ... with 894 more rows
```

```
plot(data, log="y")
```

- #> Warning in self\$trans\$transform(x): NaNs produced
- #> Warning: Transformation introduced infinite values in continuous y-axis



12.4.5 Daptomycin Reference

This weight-based simulation is adapted from the Daptomycin article below:

Dvorchik B, Arbeit RD, Chung J, Liu S, Knebel W, Kastrissios H. Population pharmacokinetics of daptomycin. Antimicrob Agents Che mother 2004; 48: 2799-2807. doi:(10.1128/AAC.48.8.2799-2807.2004)[https://dx.doi.org/10.1128%2FAAC.48.8. 2799-2807.2004]

This simulation example was made available from the work of Sherwin Sy with modifications by Matthew Fidler

12.5 Inter-occasion and other nesting examples

More than one level of nesting is possible in RxODE; In this example we will be using the following uncertainties and sources of variability:

Level	Variable	Matrix specified	Integrated Matrix
Model uncertainty	NA	thetaMat	thetaMat
Investigator	inv.Cl,inv.Ka	omega	theta
Subject	eta.Cl,eta.Ka	omega	omega
Eye	eye.Cl,eye.Ka	omega	omega
Occasion	iov.Cl,occ.Ka	omega	omega
Unexplained Concentration	prop.sd	sigma	sigma
Unexplained Effect	add.sd	sigma	sigma

12.5.1 Event table

This event table contains nesting variables:

- · inv: investigator id
- id: subject id
- eye: eye id (left or right)
- occ: occasion

```
library(RxODE)
library(dplyr)

et(amountUnits="mg", timeUnits="hours") %>%
    et(amt=10000, addl=9,ii=12,cmt="depot") %>%
    et(time=120, amt=2000, addl=4, ii=14, cmt="depot") %>%
    et(seq(0, 240, by=4)) %>% # Assumes sampling when there is no dosing information et(seq(0, 240, by=4) + 0.1) %>% ## adds 0.1 for separate eye et(id=1:20) %>%
    ## Add an occasion per dose
    mutate(occ=cumsum(!is.na(amt))) %>%
    mutate(occ=ifelse(occ == 0, 1, occ)) %>%
    mutate(occ=2- occ %% 2) %>%
    mutate(eye=ifelse(round(time) == time, 1, 2)) %>%
    mutate(inv=ifelse(id < 10, 1, 2)) %>% as_tibble ->
ev
```

12.5.2 RxODE model

This creates the RxODE model with multi-level nesting. Note the variables inv.Cl, inv.Ka, eta.Cl etc; You only need one variable for each level of nesting.

```
mod <- RxODE({
    ## Clearance with individuals
    eff(0) = 1
    C2 = centr/V2*(1+prop.sd);
    C3 = peri/V3;
    CL = TCl*exp(eta.Cl + eye.Cl + iov.Cl + inv.Cl)
    KA = TKA * exp(eta.Ka + eye.Ka + iov.Cl + inv.Ka)
    d/dt(depot) =-KA*depot;
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) = Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
    ef0 = eff + add.sd
})</pre>
```

12.5.3 Uncertainty in Model parameters

```
#>
             TKA
                        TCl
                                   V2
                                              Q
                                                        V3
#> TKA 0.084901793 0.008491535 -0.028211905 -0.083460075 0.058187918
#> TCl
     -0.028211905 -0.047388281 0.094485793 -0.001142105 -0.031424916
#> Q
      -0.083460075 -0.049711532 -0.001142105 0.271685090 -0.042012559
#> V3
       #> Kin -0.046169545 -0.016205743 -0.014705754 0.094478654 0.001534101
#> Kout -0.023946567 -0.040201822 0.008377087 0.062087105 -0.053276590
#> EC50 -0.017733886 -0.011145761 -0.004038295 0.025496213 0.006360558
                                 EC50
             Kin
                       Kout.
#> TKA -0.046169545 -0.023946567 -0.017733886
#> TCl -0.016205743 -0.040201822 -0.011145761
```

12.5.4 Nesting Variability

To specify multiple levels of nesting, you can specify it as a nested lotri matrix; When using this approach you use the condition operator | to specify what variable nesting occurs on; For the Bayesian simulation we need to specify how much information we have for each parameter; For RxODE this is the nu parameter.

In this case: - id, nu=100 or the model came from 100 subjects - eye, nu=200 or the model came from 200 eyes - occ, nu=200 or the model came from 200 occasions - inv, nu=10 or the model came from 10 investigators

To specify this in lotri you can use | var(nu=X), or:

```
#> $id
#>
          eta.Cl eta.Ka
#> eta.Cl
             0.1
                    0.0
#> eta.Ka
             0.0
                    0.1
#>
#> $eye
#>
          eye.Cl eye.Ka
            0.05
                   0.00
#> eye.Cl
#> eye.Ka
            0.00
                   0.05
#>
#> $occ
#>
          iov.Cl iov.Ka
#> iov.Cl 0.01
                   0.00
#> iov.Ka 0.00
                   0.01
#>
```

```
#> $inv
#> inv.Cl inv.Ka
#> inv.Cl 0.02 0.00
#> inv.Ka 0.00 0.02
#>
#> Properties: nu
```

12.5.5 Unexplained variability

The last piece of variability to specify is the unexplained variability

12.5.6 Solving the problem

```
s <- rxSolve(mod, theta, ev,
          thetaMat=tMat, omega=omega,
          sigma=sigma, sigmaDf=400,
          nStud=400)
#> unhandled error message: EE:[lsoda] 70000 steps taken before reaching tout
#> @(lsoda.c:748
#> Warning: some ID(s) could not solve the ODEs correctly; These values are
#> replaced with 'NA'
print(s)
#> ______ Solved RxODE object _____
#> -- Parameters ($params): -----
#> # A tibble: 8,000 x 24
    sim.id id    `inv.Cl(inv==1)` `inv.Cl(inv==2)` `inv.Ka(inv==1)`
#>
#>
     <int> <fct>
                                     <dbl>
                       <dbl>
                                                   <dbl>
#> 1
      1 1
                       0.0186
                                    0.116
                                                   0.188
#> 2
       1 2
                      0.0186
                                    0.116
                                                  0.188
       1 3
#> 3
                      0.0186
                                    0.116
                                                  0.188
#> 4
       1 4
                      0.0186
                                    0.116
                                                   0.188
#> 5
       15
                      0.0186
                                    0.116
                                                  0.188
       1 6
#> 6
                     0.0186
                                    0.116
                                                  0.188
#> 7 1 7
                     0.0186
                                    0.116
                                                   0.188
```

```
#>
  8
        1 8
                      0.0186
                                    0.116
                                                 0.188
        1 9
                      0.0186
#> 9
                                    0.116
                                                 0.188
#> 10
        1 10
                      0.0186
                                    0.116
                                                 0.188
#> # ... with 7,990 more rows, and 19 more variables: inv.Ka(inv==2) <dbl>,
     eye.Cl(eye==1) <dbl>, eye.Cl(eye==2) <dbl>, eye.Ka(eye==1) <dbl>,
#> #
     eye.Ka(eye==2) <dbl>, iov.Cl(occ==1) <dbl>, iov.Cl(occ==2) <dbl>,
#> #
     iov.Ka(occ==1) <dbl>, iov.Ka(occ==2) <dbl>, V2 <dbl>, V3 <dbl>, TC1 <dbl>,
#> #
     eta.Cl <dbl>, TKA <dbl>, eta.Ka <dbl>, Q <dbl>, Kin <dbl>, Kout <dbl>,
#> #
     EC50 <dbl>
#> -- Initial Conditions ($inits): -------
#> depot centr peri
                 eff
#>
     Λ
          0
              0
#>
#> Simulation with uncertainty in:
#> * parameters (s$thetaMat for changes)
#> * omega matrix (s$omegaList)
#>
#> -- First part of data (object): ------
#> # A tibble: 976,000 x 18
#>
           id time inv.Cl inv.Ka eye.Cl eye.Ka iov.Cl iov.Ka
                                                      C2
                                                            C3
   sim.id
#>
    <int> <int>
               [h] <dbl> <dbl> <dbl> <dbl> <dbl>
                                               <dbl> <dbl>
                                                          <dbl>
#> 1
       1
               0
#> 2
               1
                                                         0.0281
#> 3
               1
            1
                                                         9.90
#> 4
       1
            1
               4.1 0.0186   0.188   -0.296   0.0588   0.0247   -0.0530   57.9   10.1
#> 5
               1
            1
               #> # ... with 975,994 more rows, and 7 more variables: CL <dbl>, KA <dbl>,
   efO <dbl>, depot <dbl>, centr <dbl>, peri <dbl>, eff <dbl>
```

There are multiple investigators in a study; Each investigator has a number of individuals enrolled at their site. RxODE automatically determines the number of investigators and then will simulate an effect for each investigator. With the output, inv.Cl(inv==1) will be the inv.Cl for investigator l, inv.Cl(inv==2) will be the inv.Cl for investigator 2, etc.

inv.Cl(inv==1), inv.Cl(inv==2), etc will be simulated for each study and then combined to form the between investigator variability. In equation form these represent the following:

```
inv.Cl = (inv == 1) * `inv.Cl(inv==1)` + (inv == 2) * `inv.Cl(inv==2)`
```

If you look at the simulated parameters you can see inv.Cl(inv==1) and inv.Cl(inv==2) are in the s\$params; They are the same for each study:

```
print(head(s$params))
     sim.id id inv.Cl(inv==1) inv.Cl(inv==2) inv.Ka(inv==1) inv.Ka(inv==2)
          1
            1
                   0.01864161
                                   0.1159198
                                                  0.1878234
                                                                 -0.292727
#> 2
          1 2
                   0.01864161
                                                  0.1878234
                                                                 -0.292727
                                   0.1159198
#> 3
          1 3
                   0.01864161
                                   0.1159198
                                                  0.1878234
                                                                 -0.292727
          1 4
#> 4
                   0.01864161
                                   0.1159198
                                                  0.1878234
                                                                 -0.292727
#> 5
          1 5
                   0.01864161
                                   0.1159198
                                                  0.1878234
                                                                 -0.292727
#> 6
          1 6
                   0.01864161
                                   0.1159198
                                                  0.1878234
                                                                 -0.292727
     eye.Cl(eye==1) eye.Cl(eye==2) eye.Ka(eye==1) eye.Ka(eye==2) iov.Cl(occ==1)
#>
       -0.10410790
                       -0.29632211
                                       0.07855205
                                                      0.05884539
                                                                     0.02466606
#>
#> 2
       -0.06820792
                        0.06585538
                                       0.34603340
                                                      0.20141355
                                                                    -0.06976537
#> 3
       -0.04885836
                        0.13135196
                                      -0.13256387
                                                      0.21645151
                                                                    -0.03121109
#> 4
        0.20667975
                       -0.08775327
                                      -0.01404241
                                                     -0.04239568
                                                                    -0.08207797
#> 5
         0.04877033
                       -0.22890756
                                      -0.21685969
                                                      0.04846680
                                                                    -0.01393029
                                      -0.16960164
#> 6
         0.19830134
                       -0.33204702
                                                      0.06823678
                                                                    -0.17462695
     iov.Cl(occ==2) iov.Ka(occ==1) iov.Ka(occ==2)
                                                        V2
                                                                  VЗ
                    -0.052971164
                                    -0.106088075 39.57653 297.3736 18.81115
#> 1
       -0.11264526
#> 2
        0.03970507
                      -0.073742566
                                      0.090882718 39.57653 297.3736 18.81115
                                    0.067412442 39.57653 297.3736 18.81115
#> 3
       -0.23892944
                     -0.136470596
                     -0.061910605
                                    -0.072601879 39.57653 297.3736 18.81115
#> 4
       -0.02134625
#> 5
        0.05580236
                      0.099876044
                                    -0.094708943 39.57653 297.3736 18.81115
#> 6
        -0.03152016
                      -0.002074008
                                    -0.004758332 39.57653 297.3736 18.81115
#>
          eta.Cl
                       TKA
                                eta.Ka
                                                     Kin
                                                              Kont.
#> 1 -0.02740884 0.4375889 0.07529548 10.97588 1.179938 0.9161172 200.2625
#> 2 -0.11896272 0.4375889 0.07490355 10.97588 1.179938 0.9161172 200.2625
#> 3 -0.61026874 0.4375889 -0.15964154 10.97588 1.179938 0.9161172 200.2625
#> 4 -0.17447915 0.4375889 -0.19377239 10.97588 1.179938 0.9161172 200.2625
#> 5  0.26213020  0.4375889  -0.38954283  10.97588  1.179938  0.9161172  200.2625
#> 6 -0.22932331 0.4375889 -0.49123723 10.97588 1.179938 0.9161172 200.2625
print(head(s$params %>% filter(sim.id == 2)))
     sim.id id inv.Cl(inv==1) inv.Cl(inv==2) inv.Ka(inv==1) inv.Ka(inv==2)
#> 1
                                                                 0.0902204
          2 1
                  -0.01105301
                                  -0.1209402
                                                  0.3370577
#> 2
          2 2
                  -0.01105301
                                  -0.1209402
                                                  0.3370577
                                                                 0.0902204
#> 3
          2 3
                  -0.01105301
                                  -0.1209402
                                                                 0.0902204
                                                  0.3370577
          2 4
                                  -0.1209402
#> 4
                  -0.01105301
                                                  0.3370577
                                                                 0.0902204
#> 5
          2 5
                  -0.01105301
                                  -0.1209402
                                                  0.3370577
                                                                 0.0902204
#> 6
          2 6
                  -0.01105301
                                  -0.1209402
                                                  0.3370577
                                                                  0.0902204
#>
     eye.Cl(eye==1) eye.Cl(eye==2) eye.Ka(eye==1) eye.Ka(eye==2) iov.Cl(occ==1)
#> 1
       -0.01262553
                     -0.08161227
                                    -0.238499594
                                                      0.17178813
                                                                     0.08330981
#> 2
       -0.06778157
                        0.29410669
                                   -0.003700213
                                                     -0.03805489
                                                                    -0.12869095
#> 3
     0.06059738
                      -0.16831575 -0.085582067
                                                    0.22970053
                                                                     0.05711749
```

```
#> 4
        -0.13086494
                        0.02748735
                                     -0.056454551
                                                     -0.23331112
                                                                     0.07216869
#> 5
        -0.23416424
                       -0.13568099
                                     -0.436719663
                                                     -0.03106162
                                                                     -0.13191139
#> 6
         0.24092815
                        0.66166495
                                     -0.345840539
                                                      0.13552870
                                                                      0.03987511
#>
    iov.Cl(occ==2) iov.Ka(occ==1) iov.Ka(occ==2)
                                                        ٧2
                                                                  ٧3
                                                                          TCl
#> 1
       -0.148592318
                        0.10100830
                                      0.050123275 40.32538 296.5826 18.65152
#> 2
       -0.002200205
                       -0.04045931 -0.077835601 40.32538 296.5826 18.65152
#> 3
                        0.02903611
                                      0.076384740 40.32538 296.5826 18.65152
      -0.185116411
#> 4
        0.101902663
                        0.04680555
                                      0.054662894 40.32538 296.5826 18.65152
#> 5
                                      0.100946619 40.32538 296.5826 18.65152
        0.103696663
                        0.02589958
#> 6
        0.023244426
                       -0.03067335
                                     -0.009347601 40.32538 296.5826 18.65152
         eta.Cl
                       TKA
                                eta.Ka
                                                     Kin
                                                             Kout
#> 1 -0.2518665 -0.2160777  0.33092617 11.02462 1.122102 1.022177 199.9981
#> 2 0.3495104 -0.2160777 -0.35774607 11.02462 1.122102 1.022177 199.9981
#> 3 -0.3101379 -0.2160777 -0.09014428 11.02462 1.122102 1.022177 199.9981
#> 4 -0.1665144 -0.2160777 -0.11974060 11.02462 1.122102 1.022177 199.9981
#> 5  0.3184297 -0.2160777 -0.06982612 11.02462 1.122102 1.022177 199.9981
#> 6 -0.1216137 -0.2160777 0.30275205 11.02462 1.122102 1.022177 199.9981
```

For between eye variability and between occasion variability each individual simulates a number of variables that become the between eye and between occasion variability; In the case of the eye:

```
eye.C1 = (eye == 1) * `eye.C1(eye==1)` + (eye == 2) * `eye.C1(eye==2)`
```

So when you look the simulation each of these variables (ie eye.Cl(eye==1), eye.Cl(eye==2), etc) they change for each individual and when combined make the between eye variability or the between occasion variability that can be seen in some pharamcometric models.

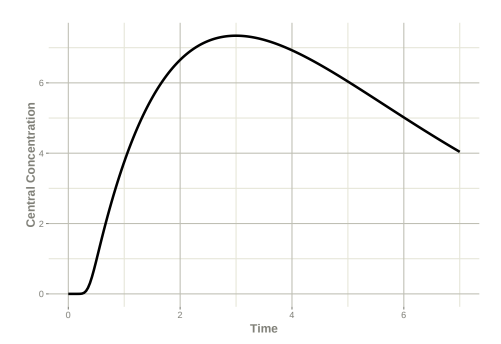
12.6 Transit compartment models

Savic 2008 first introduced the idea of transit compartments being a mechanistic explanation of a a lag-time type phenomena. RxODE has special handling of these models:

You can specify this in a similar manner as the original paper:

```
library(RxODE)
mod <- RxODE({
    ## Table 3 from Savic 2007
    cl = 17.2 # (L/hr)
    vc = 45.1 # L
    ka = 0.38 # 1/hr</pre>
```

```
mtt = 0.37 # hr
bio=1
n = 20.1
k = cl/vc
ktr = (n+1)/mtt
## note that lgammafn is the same as lgamma in R.
d/dt(depot) = exp(log(bio*podo)+log(ktr)+n*log(ktr*t)-ktr*t-lgammafn(n+1))-ka*depot
d/dt(cen) = ka*depot-k*cen
})
et <- eventTable();
et$add.sampling(seq(0, 7, length.out=200));
et$add.dosing(20, start.time=0);
transit <- rxSolve(mod, et, transit_abs=TRUE)
plot(transit, cen, ylab="Central Concentration")</pre>
```

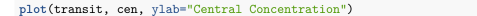


Another option is to specify the transit compartment function transit syntax. This specifies the parameters transit(number of transit compartments, mean transit time, bioavailability). The bioavailability term is optional.

Using the transit code also automatically turns on the transit_abs option. Therefore, the same model can be specified by:

```
mod <- RxODE({</pre>
    ## Table 3 from Savic 2007
    cl = 17.2 \# (L/hr)
    vc = 45.1 \# L
    ka = 0.38 \# 1/hr
    mtt = 0.37 \# hr
    bio=1
    n = 20.1
    k = c1/vc
    ktr = (n+1)/mtt
    d/dt(depot) = transit(n,mtt,bio)-ka*depot
    d/dt(cen) = ka*depot-k*cen
})
et <- eventTable();</pre>
et$add.sampling(seq(0, 7, length.out=200));
et$add.dosing(20, start.time=0);
transit <- rxSolve(mod, et)</pre>
```

#> Warning: assumed transit compartment model since 'podo' is in the model





Chapter 13

Advanced & Miscellaneous Topics

This covers advanced or miscellaneous topics in RxODE

13.1 Covariates in RxODE

13.1.1 Individual Covariates

If there is an individual covariate you wish to solve for you may specify it by the iCov dataset:

```
library(RxODE)
library(units)
library(xgxr)

mod3 <- RxODE({
    KA=2.94E-01;
#### Clearance with individuals
    CL=1.86E+01 * (WT / 70) ^ 0.75;
    V2=4.02E+01;
    Q=1.05E+01;
    V3=2.97E+02;
    Kin=1;
    Kout=1;
    EC50=200;
#### The linCmt() picks up the variables from above
    C2 = linCmt();</pre>
```

```
Tz=8
   amp=0.1
   eff(0) = 1 ## This specifies that the effect compartment starts at 1.
   d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})
ev <- et(amount.units="mg", time.units="hours") %>%
   et(amt=10000, cmt=1) %>%
   et(0,48,length.out=100) %>%
   et(id=1:4);
set.seed(10)
#### Now use iCov to simulate a 4-id sample
r1 <- solve(mod3, ev,
### Create individual covariate data-frame
           iCov=data.frame(id=1:4, WT=rnorm(4, 70, 10)),
### in this case it would be useful to keep the WT in the output dataset
          keep="WT")
print(r1)
#> ______ Solved RxODE object _____
#> -- Parameters ($params): ------
#> # A tibble: 4 x 11
#> id
                              Q V3 Kin Kout EC50
            KA
                  WT
                       V2
                                                        Tz
                                                             amp
#> <fct> <dbl> <
                                                      8 0.1
#> 1 1
        0.294 70.2 40.2 10.5 297 1 1 200

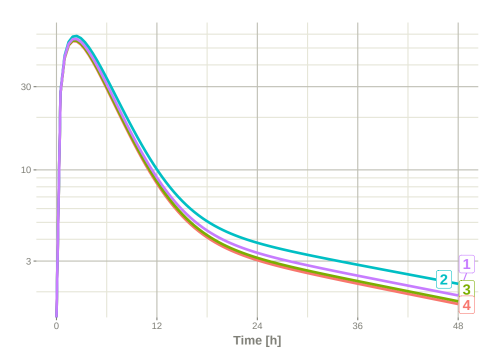
      0.294
      68.2
      40.2
      10.5
      297
      1
      1
      200

      0.294
      56.3
      40.2
      10.5
      297
      1
      1
      200

        0.294 68.2 40.2 10.5 297
                                                        8 0.1
#> 2 2
#> 3 3
                                                        8 0.1
         0.294 64.0 40.2 10.5 297 1 1 200 8 0.1
#> -- Initial Conditions ($inits): ------
#> eff
#> 1
#> -- First part of data (object): ------
#> # A tibble: 400 x 6
#>
       id
              time
                     CL
                          C2 eff WT
               [h] <dbl> <dbl> <dbl> <dbl>
#> <int>
#> 1
      1 0.0000000 18.6 0 1
        1 0.4848485 18.6 27.8 1.03 70.2
#> 2
#> 3
        1 0.9696970 18.6 43.7 1.08 70.2
#> 4
      1 1.4545455 18.6 51.7 1.13 70.2
#> 5
      1 1.9393939 18.6 54.7 1.18 70.2
#> 6
       1 2.4242424 18.6 54.5 1.21 70.2
#> # ... with 394 more rows
```

```
plot(r1, C2, log="y")
```

#> Warning: Transformation introduced infinite values in continuous y-axis



13.1.2 Time Varying Covariates

Covariates are easy to specify in RxODE, you can specify them as a variable. Time-varying covariates, like clock time in a circadian rhythm model, can also be used. Extending the indirect response model already discussed, we have:

```
library(RxODE)
library(units)

mod3 <- RxODE({
    KA=2.94E-01;
    CL=1.86E+01;
    V2=4.02E+01;
    Q=1.05E+01;
    V3=2.97E+02;
    KinO=1;
    Kout=1;</pre>
```

```
EC50=200;
#### The linCmt() picks up the variables from above
   C2 = linCmt();
   Tz=8
   amp=0.1
    eff(0) = 1 ## This specifies that the effect compartment starts at 1.
#### Kin changes based on time of day (like cortosol)
   Kin = Kin0 + amp *cos(2*pi*(ctime-Tz)/24)
   d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})
ev <- eventTable(amount.units="mg", time.units="hours") %>%
    add.dosing(dose=10000, nbr.doses=1, dosing.to=1) %>%
    add.sampling(seq(0,48,length.out=100));
#### Create data frame of 8 am dosing for the first dose This is done
#### with base R but it can be done with dplyr or data.table
ev$ctime <- (ev$time+set_units(8,hr)) %% 24
```

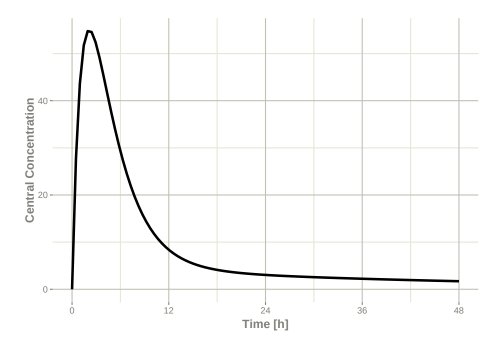
Now there is a covariate present in the event dataset, the system can be solved by combining the dataset and the model:

```
r1 <- solve(mod3, ev, covs_interpolation="linear")</pre>
print(r1)
#> ______ Solved RxODE object ______
#> -- Parameters ($params): -------
    KA CL V2 Q V3 KinO
                                                    Kout
#> 0.294000 18.600000 40.200000 10.500000 297.000000 1.000000 1.000000
#> EC50 Tz amp
                         pi
#> 200.000000 8.000000 0.100000 3.141593
#> -- Initial Conditions ($inits): ------
#> eff
#> -- First part of data (object): ------
#> # A tibble: 100 x 4
     time C2 Kin eff
       [h] <dbl> <dbl> <dbl>
#> 1 0.0000000 0 1.1 1
#> 2 0.4848485 27.8 1.10 1.07
#> 3 0.9696970 43.7 1.10 1.15
#> 4 1.4545455 51.8 1.09 1.22
#> 5 1.9393939 54.8 1.09 1.27
#> 6 2.4242424 54.6 1.08 1.30
```

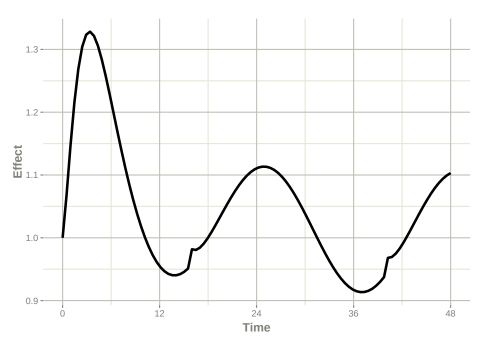
```
#> # ... with 94 more rows
#> ______
```

When solving ODE equations, the solver may sample times outside of the data. When this happens, this ODE solver can use linear interpolation between the covariate values. It is equivalent to R's approxfun with method="linear".

```
plot(r1,C2, ylab="Central Concentration")
```



```
plot(r1,eff) + ylab("Effect") + xlab("Time");
```



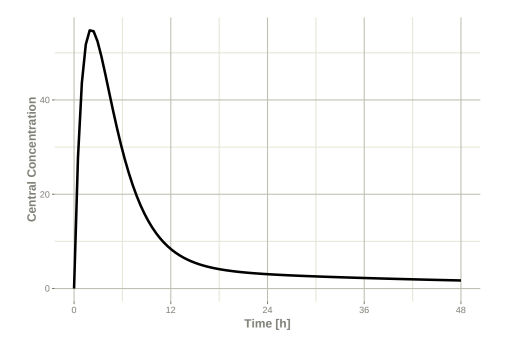
Note that the linear approximation in this case leads to some kinks in the solved system at 24-hours where the covariate has a linear interpolation between near 24 and near 0. While linear seems reasonable, cases like clock time make other interpolation methods more attractive.

In RxODE the default covariate interpolation is be the last observation carried forward (locf), or constant approximation. This is equivalent to R's approxfun with method="constant".

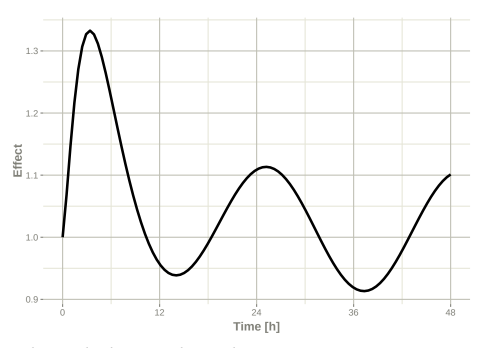
```
r1 <- solve(mod3, ev,covs_interpolation="constant")
print(r1)
              ______ Solved RxODE object _____
#> _____
#> -- Parameters ($params): -----
                          V2 Q
        KA CL
#>
   0.294000 18.600000 40.200000 10.500000 297.000000 1.000000 1.000000
#>
       EC50
                Tz
                         amp
                                  pi
#> 200.000000 8.000000
                    0.100000 3.141593
#> -- Initial Conditions ($inits): -----
#> eff
#> -- First part of data (object): ------
#> # A tibble: 100 x 4
        time C2 Kin eff
#>
#>
        [h] <dbl> <dbl> <dbl>
#> 1 0.0000000 0 1.1 1
#> 2 0.4848485 27.8 1.10 1.07
```

which gives the following plots:

```
plot(r1,C2, ylab="Central Concentration", xlab="Time")
```



```
plot(r1,eff, ylab="Effect", xlab="Time")
```



In this case, the plots seem to be smoother.

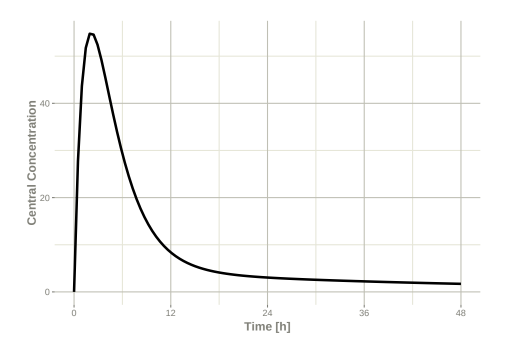
You can also use NONMEM's preferred interpolation style of next observation carried backward (NOCB):

```
r1 <- solve(mod3, ev,covs_interpolation="nocb")
print(r1)
#> ______ Solved RxODE object _____
#> -- Parameters ($params): ------
             CL V2
                            Q
      KA
                                          V3
                                                  KinO
                                                           Kout
   0.294000 18.600000 40.200000 10.500000 297.000000 1.000000 1.0000000
#>
                         amp pi
      EC50
#>
           Tz
#> 200.000000 8.000000 0.100000 3.141593
#> -- Initial Conditions ($inits): ------
#> eff
#> -- First part of data (object): -----
#> # A tibble: 100 x 4
      time C2 Kin eff
        [h] < db l> < db l>
#>
#> 1 0.0000000 0 1.1 1
#> 2 0.4848485 27.8 1.10 1.07
#> 3 0.9696970 43.7 1.10 1.15
#> 4 1.4545455 51.8 1.09 1.21
#> 5 1.9393939 54.8 1.09 1.27
#> 6 2.4242424 54.6 1.08 1.30
```

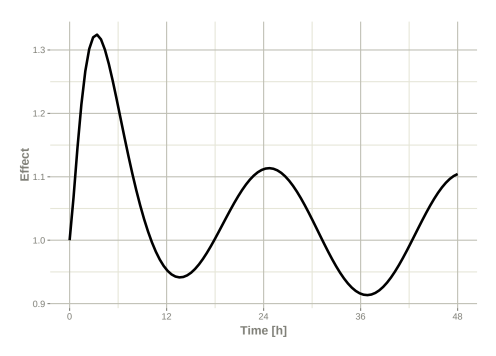
```
#> # ... with 94 more rows
#> ______
```

which gives the following plots:

```
plot(r1,C2, ylab="Central Concentration", xlab="Time")
```



```
plot(r1,eff, ylab="Effect", xlab="Time")
```



13.2 Shiny and RxODE

13.2.1 Facilities for generating R shiny applications

An example of creating an R shiny application to interactively explore responses of various complex dosing regimens is available at http://qsp.engr.uga.edu:3838/RxODE/RegimenSimulator. Shiny applications like this one may be programmatically created with the experimental function genShinyApp.template().

The above application includes widgets for varying the dose, dosing regimen, dose cycle, and number of cycles.

```
genShinyApp.template(appDir = "shinyExample", verbose=TRUE)
library(shiny)
runApp("shinyExample")
```

Click here to go to the Shiny App

13.2.2 Exploring parameter fits graphically using shiny

An RxODE object can be explored with rxShiny(obj). rxShiny() will also allow you to try new models to see how they behave.

13.3 Using RxODE with a pipeline

13.3.1 Setting up the RxODE model for the pipeline

In this example we will show how to use RxODE in a simple pipeline.

We can start with a model that can be used for the different simulation workflows that RxODE can handle:

```
library(RxODE)
Ribba2012 <- RxODE({</pre>
   k = 100
    tkde = 0.24
    eta.tkde = 0
    kde ~ tkde*exp(eta.tkde)
    tkpq = 0.0295
    eta.kpq = 0
    kpq ~ tkpq * exp(eta.kpq)
    tkqpp = 0.0031
    eta.kqpp = 0
    kqpp ~ tkqpp * exp(eta.kqpp)
    tlambdap = 0.121
    eta.lambdap = 0
    lambdap ~ tlambdap*exp(eta.lambdap)
    tgamma = 0.729
    eta.gamma = 0
    gamma ~ tgamma*exp(eta.gamma)
    tdeltagp = 0.00867
    eta.deltaqp = 0
    deltaqp ~ tdeltaqp*exp(eta.deltaqp)
    prop.err <- 0
    pstar <- (pt+q+qp)*(1+prop.err)</pre>
    d/dt(c) = -kde * c
    d/dt(pt) = lambdap * pt *(1-pstar/k) + kqpp*qp -
        kpq*pt - gamma*c*kde*pt
    d/dt(q) = kpq*pt -gamma*c*kde*q
    d/dt(qp) = gamma*c*kde*q - kqpp*qp - deltaqp*qp
```

```
#### initial conditions

tpt0 = 7.13
  eta.pt0 = 0
  pt0 ~ tpt0*exp(eta.pt0)

tq0 = 41.2
  eta.q0 = 0
  q0 ~ tq0*exp(eta.q0)
  pt(0) = pt0
  q(0) = q0
}
```

This is a tumor growth model described in Ribba 2012. In this case, we compiled the model into an R object Ribba2012, though in an RxODE simulation pipeline, you do not *have* to assign the compiled model to any object, though I think it makes sense.

13.3.2 Simulating one event table

Simulating a single event table is quite simple:

pstar (total tumor tissue)

- You pipe the RxODE simulation object into an event table object by et().
- When the events are completely specified, you simply solve the ODE system with rxSolve().
- In this case you can pipe the output to plot() to conveniently view the results.
- Note for the plot we are only selecting the selecting following:

```
pt (Proliferative Tissue),q (quiescent tissue)qp (DNA-Damaged quiescent tissue) and
```

```
Ribba2012 %>% # Use RxODE

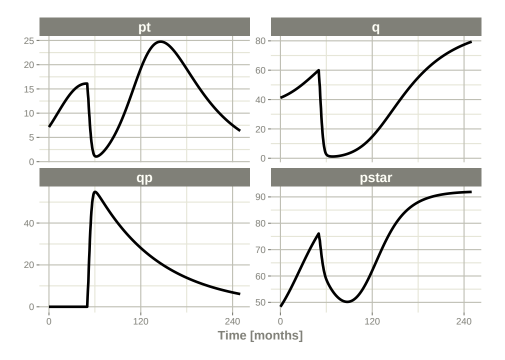
et(time.units="months") %>% # Pipe to a new event table

et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months

et(0, 250, by=0.5) %>% # Add some sampling times (not required)

rxSolve() %>% # Solve the simulation

plot(pt, q, qp, pstar) # Plot it, plotting the variables of interest
```



13.3.3 Simulating multiple subjects from a single event table

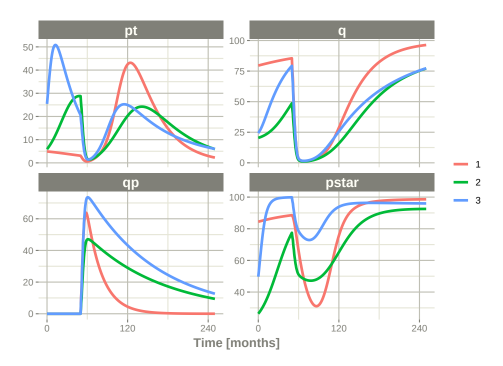
13.3.3.1 Simulating with between subject variability

The next sort of simulation that may be useful is simulating multiple patients with the same treatments. In this case, we will use the omega matrix specified by the paper:

```
omega
#>
            eta.pt0
                    eta.q0 eta.lambdap
                                   eta.kqp
                                          eta.qpp eta.deltaqp
#> eta.pt0
          0.0000000
#> eta.q0
          0.0000000 0.2558818
                           0.0000000 0.0000000 0.0000000
                                                  0.0000000
#> eta.lambdap 0.0000000 0.0000000 0.4176571 0.0000000 0.0000000
                                                  0.0000000
#> eta.kqp 0.0000000 0.0000000
                          0.0000000 0.4559047 0.0000000
                                                  0.0000000
#> eta.qpp
         0.0000000 0.0000000 0.0000000 0.0000000 0.6631518
                                                  0.0000000
0.8426442
0.0000000
#>
            eta.kde
#> eta.pt0
          0.0000000
#> eta.q0
         0.0000000
#> eta.lambdap 0.0000000
#> eta.kqp 0.0000000
#> eta.qpp
          0.0000000
#> eta.deltaqp 0.0000000
#> eta.kde 0.3987761
```

With this information, it is easy to simulate 3 subjects from the model-based parameters:

```
set.seed(1089)
Ribba2012 %>% # Use RxODE
et(time.units="months") %>% # Pipe to a new event table
et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months
et(0, 250, by=0.5) %>% # Add some sampling times (not required)
rxSolve(nSub=3, omega=omega) %>% # Solve the simulation
plot(pt, q, qp, pstar) # Plot it, plotting the variables of interest
```



Note there are two different things that were added to this simulation: - nSub to specify how many subjects are in the model - omega to specify the between subject variability.

13.3.3.2 Simulation with unexplained variability

You can even add unexplained variability quite easily:

```
Ribba2012 %>% # Use RxODE

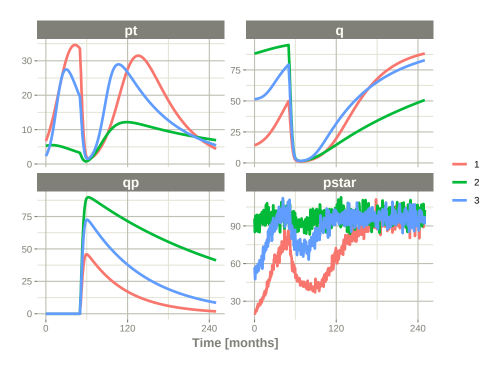
et(time.units="months") %>% # Pipe to a new event table

et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months

et(0, 250, by=0.5) %>% # Add some sampling times (not required)

rxSolve(nSub=3, omega=omega, sigma=lotri(prop.err ~ 0.05^2)) %>% # Solve the simulation

plot(pt, q, qp, pstar) # Plot it, plotting the variables of interest
```



In this case we only added the sigma matrix to have unexplained variability on the pstar or total tumor tissue.

You can even simulate with uncertainty in the theta omega and ${\tt sigma}$ values if you wish.

13.3.3.3 Simulation with uncertainty in all the parameters (by matrices)

If we assume these parameters came from 95 subjects with 8 observations apiece, the degrees of freedom for the omega matrix would be 95, and the degrees of freedom of the sigma matrix would be 95*8=760 because 95 items informed the omega matrix, and 760 items informed the sigma matrix.

```
Ribba2012 %>% # Use RxODE

et(time.units="months") %>% # Pipe to a new event table

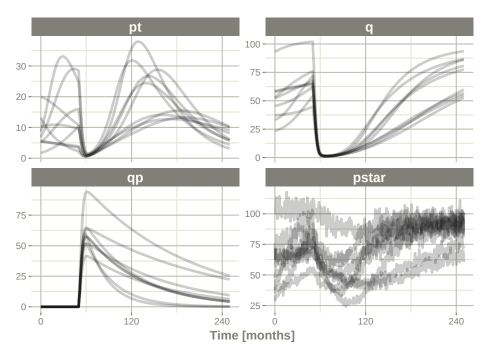
et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months

et(0, 250, by=0.5) %>% # Add some sampling times (not required)

rxSolve(nSub=3, nStud=3, omega=omega, sigma=lotri(prop.err ~ 0.05^2),

dfSub=760, dfObs=95) %>% # Solve the simulation

plot(pt, q, qp, pstar) # Plot it, plotting the variables of interest
```



Often in simulations we have a full covariance matrix for the fixed effect parameters. In this case, we do not have the matrix, but it could be specified by thetaMat.

While we do not have a full covariance matrix, we can have information about the diagonal elements of the covariance matrix from the model paper. These can be converted as follows:

```
rseVar <- function(est, rse){</pre>
    return(est*rse/100)^2
}
thetaMat <- lotri(tpt0 ~ rseVar(7.13,25),</pre>
                  tq0 \sim rseVar(41.2,7),
                  tlambdap ~ rseVar(0.121, 16),
                  tkqpp ~ rseVar(0.0031, 35),
                  tdeltaqp ~ rseVar(0.00867, 21),
                  tgamma ~ rseVar(0.729, 37),
                  tkde ~ rseVar(0.24, 33)
                  );
thetaMat
#>
                     tq0 tlambdap
                                      tkqpp tdeltaqp tgamma
              tpt0
#> tpt0
            1.7825 0.000 0.00000 0.000000 0.0000000 0.00000 0.0000
            0.0000 2.884 0.00000 0.000000 0.0000000 0.00000 0.0000
#> tq0
#> tlambdap 0.0000 0.000 0.01936 0.000000 0.0000000 0.00000 0.0000
```

Now we have a thetaMat to represent the uncertainty in the theta matrix, as well as the other pieces in the simulation. Typically you can put this information into your simulation with the thetaMat matrix.

With such large variability in theta it is easy to sample a negative rate constant, which does not make sense. For example:

```
Ribba2012 %>% # Use RxODE
et(time.units="months") %>% # Pipe to a new event table
et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months
et(0, 250, by=0.5) %>% # Add some sampling times (not required)
rxSolve(nSub=2, nStud=2, omega=omega, sigma=lotri(prop.err ~ 0.05^2),
thetaMat=thetaMat,
dfSub=760, dfObs=95) %>% # Solve the simulation
plot(pt, q, qp, pstar) # Plot it, plotting the variables of interest

#> unhandled error message: EE:[lsoda] 70000 steps taken before reaching tout
#> @(lsoda.c:750
#> Warning message:
#> In rxSolve_(object, .ctl, .nms, .xtra, params, events, inits, setupOnly = .setupOnly
#> Some ID(s) could not solve the ODEs correctly; These values are replaced with NA.
```

To correct these problems you simply need to use a truncated multivariate normal and specify the reasonable ranges for the parameters. For theta this is specified by thetaLower and thetaUpper. Similar parameters are there for the other matrices: omegaLower, omegaUpper, sigmaLower and sigmaUpper. These may be named vectors, one numeric value, or a numeric vector matching the number of parameters specified in the thetaMat matrix.

In this case the simulation simply has to be modified to have thetaLower=0 to make sure all rates are positive:

```
Ribba2012 %>% # Use RxODE

et(time.units="months") %>% # Pipe to a new event table

et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months

et(0, 250, by=0.5) %>% # Add some sampling times (not required)

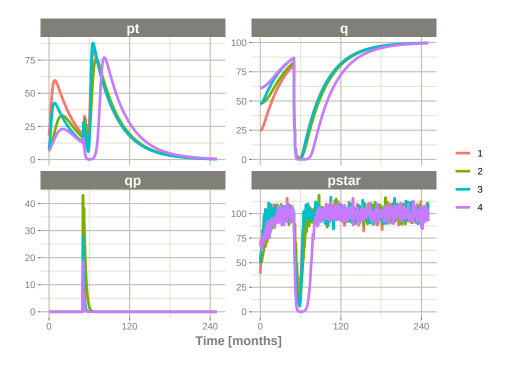
rxSolve(nSub=2, nStud=2, omega=omega, sigma=lotri(prop.err ~ 0.05^2),

thetaMat=thetaMat,

thetaLower=0, # Make sure the rates are reasonable

dfSub=760, dfObs=95) %>% # Solve the simulation

plot(pt, q, qp, pstar) # Plot it, plotting the variables of interest
```



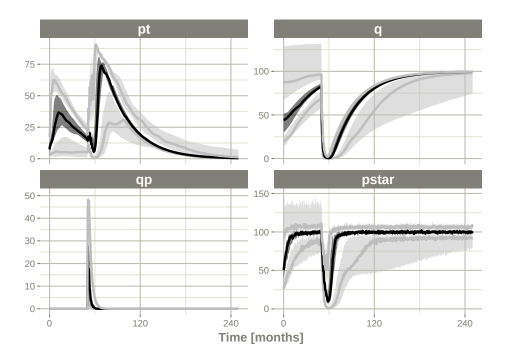
13.3.4 Summarizing the simulation output

While it is easy to use dplyr and data.table to perform your own summary of simulations, RxODE also provides this ability by the confint function.

```
#### This takes a little more time; Most of the time is the summary
#### time.

sim0 <- Ribba2012 %>% # Use RxODE
    et(time.units="months") %>% # Pipe to a new event table
    et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months
    et(0, 250, by=0.5) %>% # Add some sampling times (not required)
    rxSolve(nSub=10, nStud=10, omega=omega, sigma=lotri(prop.err ~ 0.05^2),
        thetaMat=thetaMat,
        thetaLower=0, # Make sure the rates are reasonable
        dfSub=760, dfObs=95) %>% # Solve the simulation
    confint(c("pt","q","qp","pstar"),level=0.90); # Create Simulation intervals

sim0 %>% plot() # Plot the simulation intervals
```

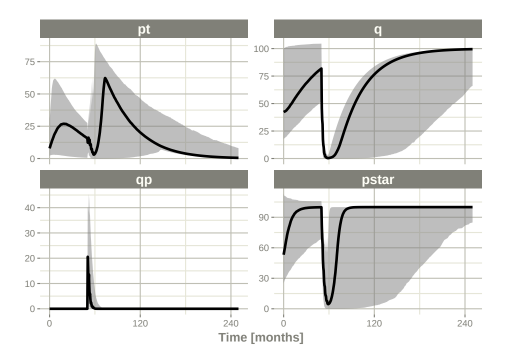


13.3.4.1 Simulating from a data-frame of parameters

While the simulation from matrices can be very useful and a fast way to simulate information, sometimes you may want to simulate more complex scenarios. For instance, there may be some reason to believe that tkde needs to be above tlambdap, therefore these need to be simulated more carefully. You can generate the data frame in whatever way you want. The internal method of simulating the new parameters is exported too.

```
library(dplyr)
pars <- rxInits(Ribba2012);</pre>
pars <- pars[regexpr("(prop|eta)",names(pars)) == -1]</pre>
print(pars)
                                  tkqpp tlambdap
#>
                          tkpq
                                                    tgamma tdeltaqp
                 tkde
                                                                         tpt0
#> 1.00e+02 2.40e-01 2.95e-02 3.10e-03 1.21e-01 7.29e-01 8.67e-03 7.13e+00
        tq0
#> 4.12e+01
#### This is the exported method for simulation of Theta/Omega internally in RxODE
df <- rxSimThetaOmega(params=pars, omega=omega,dfSub=760,</pre>
                       thetaMat=thetaMat, thetaLower=0, nSub=60,nStud=60) %>%
    filter(tkde > tlambdap) %>% as.tbl()
#### You could also simulate more and bind them together to a data frame.
print(df)
```

```
#> # A tibble: 2,340 x 16
         k tkde tkpq tkqpp tlambdap tgamma tdeltaqp tpt0 tq0 eta.pt0 eta.q0
     <dbl> <
                                                                 <dbl> <dbl>
#> 1 100 2.83 0.0295 0.239 0.683 0.861 1.25 7.67 42.0 0.559
                                                                        0.136
#> 2 100 2.83 0.0295 0.239 0.683 0.861
                                              1.25 7.67 42.0 0.0465 -0.581
#> 3 100 2.83 0.0295 0.239 0.683 0.861 1.25 7.67 42.0 -0.188 -0.180 #> 4 100 2.83 0.0295 0.239 0.683 0.861 1.25 7.67 42.0 0.321 0.614
#> 5
      100 2.83 0.0295 0.239 0.683 0.861
                                              1.25 7.67 42.0 0.0656 -0.232
#> 6 100 2.83 0.0295 0.239 0.683 0.861
                                               1.25 7.67 42.0 0.0194 0.517
#> 7 100 2.83 0.0295 0.239 0.683 0.861
                                               1.25 7.67 42.0 -0.218
                                                                       0.260
      100 2.83 0.0295 0.239
                                               1.25 7.67 42.0 -0.258 -0.761
                              0.683 0.861
#> 9 100 2.83 0.0295 0.239 0.683 0.861
                                               1.25 7.67 42.0 -1.28 -1.34
                                             1.25 7.67 42.0 -0.495 0.161
#> # ... with 2,330 more rows, and 5 more variables: eta.lambdap <dbl>,
#> # eta.kqp <dbl>, eta.qpp <dbl>, eta.deltaqp <dbl>, eta.kde <dbl>
#### Quick check to make sure that all the parameters are OK.
all(df$tkde>df$tlambdap)
#> [1] TRUE
sim1 <- Ribba2012 %>% # Use RxODE
   et(time.units="months") %>% # Pipe to a new event table
   et(amt=1, time=50, until=58, ii=1.5) %>% # Add dosing every 1.5 months
   et(0, 250, by=0.5) %>% # Add some sampling times (not required)
   rxSolve(df)
#### Note this information looses information about which ID is in a
#### "study", so it summarizes the confidence intervals by dividing the
#### subjects into sqrt(#subjects) subjects and then summarizes the
#### confidence intervals
sim2 <- sim1 %% confint(c("pt", "q", "qp", "pstar"), level=0.90); # Create Simulation intervals
save(sim2, file = file.path(system.file(package = "RxODE"), "pipeline-sim2.rds"), version = 2)
sim2 %>% plot()
```



13.4 Speeding up RxODE

13.4.1 Increasing RxODE speed by multi-subject parallel solving

RxODE originally developed as an ODE solver that allowed an ODE solve for a single subject. This flexibility is still supported.

The original code from the RxODE tutorial is below:

```
library(RxODE)

library(microbenchmark)
library(ggplot2)

mod1 <- RxODE({
    C2 = centr/V2;
    C3 = peri/V3;
    d/dt(depot) = -KA*depot;
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) = Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
    eff(0) = 1</pre>
```

13.4.1.1 For Loop

The slowest way to code this is to use a for loop. In this example we will enclose it in a function to compare timing.

```
runFor <- function(){
    res <- NULL
    for (i in 1:nsub) {
        params <- params.all[i,]
        x <- mod1$solve(params, ev, cacheEvent=FALSE)
        ##Store results for effect compartment
        res <- cbind(res, x[, "eff"])
    }
    return(res)
}</pre>
```

13.4.1.2 Running with apply

In general for R, the apply types of functions perform better than a for loop, so the tutorial also suggests this speed enhancement

```
runSapply <- function(){
   res <- apply(params.all, 1, function(theta)
        mod1$run(theta, ev, cacheEvent=FALSE)[, "eff"])
}</pre>
```

13.4.1.3 Run using a single-threaded solve

You can also have RxODE solve all the subject simultaneously without collecting the results in R, using a single threaded solve.

The data output is slightly different here, but still gives the same information:

```
runSingleThread <- function(){
    solve(mod1, params.all, ev, cores=1, cacheEvent=FALSE)[,c("sim.id", "time", "eff")]
}</pre>
```

13.4.1.4 Run a 2 threaded solve

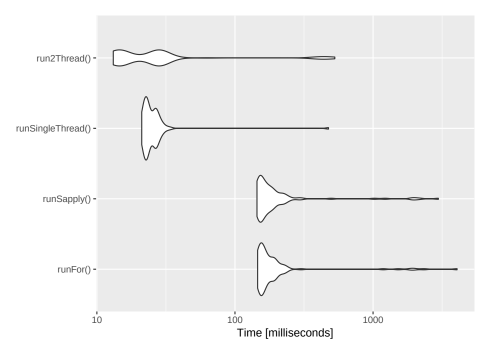
RxODE supports multi-threaded solves, so another option is to have 2 threads (called cores in the solve options, you can see the options in rxControl() or rxSolve()).

```
run2Thread <- function(){
    solve(mod1, params.all, ev, cores=2, cacheEvent=FALSE)[,c("sim.id", "time", "eff")]
}</pre>
```

13.4.1.5 Compare the times between all the methods

Now the moment of truth, the timings:

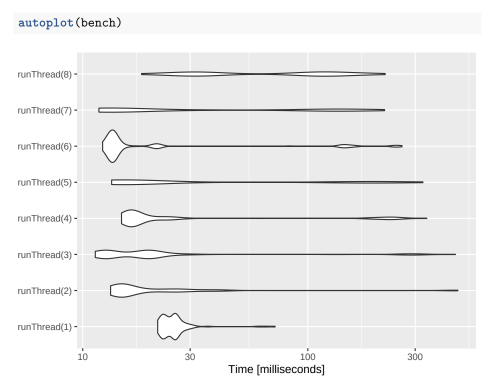
```
bench <- microbenchmark(runFor(), runSapply(), runSingleThread(),run2Thread())</pre>
print(bench)
#> Unit: milliseconds
#>
                                                       median
                           min
                                       lq
                                              mean
                                                                    uq
                                                                              max
#>
             runFor() 145.89505 153.45375 291.60899 164.34831 192.29201 4059.1394
#>
          runSapply() 144.34107 151.90202 279.92188 167.48251 192.32346 2972.0672
#> runSingleThread() 21.12513 22.52377 29.22000 23.66569 26.67664 475.6502
         run2Thread() 13.14746 14.53750 55.53308 27.33067 28.02540 528.5286
#>
#> neval
#>
    100
#>
      100
#>
      100
     100
autoplot(bench)
```



It is clear that the **largest** jump in performance when using the solve method and providing *all* the parameters to RxODE to solve without looping over each subject with either a for or a sapply. The number of cores/threads applied to the solve also plays a role in the solving.

We can explore the number of threads further with the following code:

```
runThread <- function(n){</pre>
    solve(mod1, params.all, ev, cores=n, cacheEvent=FALSE)[,c("sim.id", "time", "eff")]
}
bench <- eval(parse(text=sprintf("microbenchmark(%s)",</pre>
                                     paste(paste0("runThread(", seq(1, 2 * rxCores()),")"),
                                           collapse=","))))
print(bench)
#> Unit: milliseconds
#>
                                               median
                                                                      max neval
                      min
                                lq
                                       mean
                                                             uq
                                                       26.22489 71.53765
#>
   runThread(1) 21.52499 22.84129 26.18048 25.34360
                                                                            100
#> runThread(2) 13.29584 14.71758 32.01578 15.25110
                                                       23.38512 464.10622
                                                                            100
#> runThread(3) 11.37098 12.77958 41.49912 18.90689
                                                       20.27763 452.84269
                                                                            100
#> runThread(4) 14.86808 16.18597 50.41581 16.77116 23.34458 337.75486
                                                                            100
#> runThread(5) 13.43953 14.60734 63.06169
                                             15.11658 134.09386 324.55620
                                                                            100
#> runThread(6) 12.25469 13.35919 36.64898 13.75269 16.53334 262.00588
                                                                            100
#> runThread(7) 11.79991 12.93189 55.71493 13.25709 123.96201 219.25105
                                                                            100
#> runThread(8) 18.25941 32.12943 83.40802 105.28622 124.80418 220.56855
                                                                            100
```



There can be a suite spot in speed vs number or cores. The system type (mac, linux, windows and/or processor), complexity of the ODE solving and the number of subjects may affect this arbitrary number of threads. 4 threads is a good number to use without any prior knowledge because most systems these days have at least 4 threads (or 2 processors with 4 threads).

13.4.2 A real life example

Before some of the parallel solving was implemented, the fastest way to run RxODE was with lapply. This is how Rik Schoemaker created the data-set for nlmixr comparisons, but reduced to run faster automatic building of the pkgdown website.

```
library(RxODE)
library(data.table)
#Define the RxODE model
  ode1 <- "
  d/dt(abs) = -KA*abs;
  d/dt(centr) = KA*abs-(CL/V)*centr;
  C2=centr/V;
"</pre>
```

```
#Create the RxODE simulation object
mod1 <- RxODE(model = ode1)</pre>
#Population parameter values on log-scale
  paramsl \leftarrow c(CL = log(4),
               V = \log(70),
               KA = log(1)
#make 10,000 subjects to sample from:
 nsubg <- 300 # subjects per dose
  doses <-c(10, 30, 60, 120)
 nsub <- nsubg * length(doses)</pre>
#IIV of 30% for each parameter
  omega <- diag(c(0.09, 0.09, 0.09))# IIV covariance matrix
  sigma <- 0.2
#Sample from the multivariate normal
  set.seed(98176247)
 library(MASS)
 mv <-
    mvrnorm(nsub, rep(0, dim(omega)[1]), omega) # Sample from covariance matrix
#Combine population parameters with IIV
  params.all <-
    data.table(
      "ID" = seq(1:nsub),
      "CL" = exp(paramsl['CL'] + mv[, 1]),
      "V" = exp(paramsl['V'] + mv[, 2]),
      "KA" = exp(paramsl['KA'] + mv[, 3])
    )
#set the doses (looping through the 4 doses)
params.all[, AMT := rep(100 * doses,nsubg)]
Startlapply <- Sys.time()</pre>
#Run the simulations using lapply for speed
  s = lapply(1:nsub, function(i) {
#selects the parameters associated with the subject to be simulated
    params <- params.all[i]</pre>
#creates an eventTable with 7 doses every 24 hours
    ev <- eventTable()</pre>
    ev$add.dosing(
      dose = params$AMT,
      nbr.doses = 1,
      dosing.to = 1,
      rate = NULL,
      start.time = 0
```

```
#generates 4 random samples in a 24 hour period
    ev$add.sampling(c(0, sort(round(sample(runif(600, 0, 1440), 4) / 60, 2))))
#runs the RxODE simulation
    x <- as.data.table(mod1$run(params, ev))
#merges the parameters and ID number to the simulation output
    x[, names(params) := params]
})

#runs the entire sequence of 100 subjects and binds the results to
    res = as.data.table(do.call("rbind", s))

Stoplapply <- Sys.time()

print(Stoplapply - Startlapply)
#> Time difference of 4.786675 secs
```

By applying some of the new parallel solving concepts you can simply run the same simulation both with less code and faster:

```
rx <- RxODE({
    CL = log(4)
    V = \log(70)
    KA = log(1)
    CL = exp(CL + eta.CL)
    V = \exp(V + \text{eta.V})
    KA = \exp(KA + \text{eta.}KA)
    d/dt(abs) = -KA*abs;
    d/dt(centr) = KA*abs-(CL/V)*centr;
    C2=centr/V;
})
omega <- lotri(eta.CL ~ 0.09,
                eta.V \sim 0.09,
                eta.KA ~ 0.09)
doses <-c(10, 30, 60, 120)
startParallel <- Sys.time()</pre>
ev <- do.call("rbind",
        lapply(seq_along(doses), function(i){
             et() %>%
                 et(amt=doses[i]) %>% # Add single dose
                 et(0) %>% # Add O observation
#### Generate 4 samples in 24 hour period
```

You can see a striking time difference between the two methods; A few things to keep in mind:

- RxODE use the thread-safe sitmo threefry routines for simulation of eta values. Therefore the results are expected to be different (also the random samples are taken in a different order which would be different)
- This prior simulation was run in R 3.5, which has a different random number generator so the results in this simulation will be different from the actual nlmixr comparison when using the slower simulation.
- This speed comparison used data.table. RxODE uses data.table internally (when available) try to speed up sorting, so this would be different than installations where data.table is not installed. You can force RxODE to use order() when sorting by using forderForceBase(TRUE). In this case there is little difference between the two, though in other examples data.table's presence leads to a speed increase (and less likely it could lead to a slow-down).

13.4.2.1 Want more ways to run multi-subject simulations

The version since the tutorial has even more ways to run multi-subject simulations, including adding variability in sampling and dosing times with et() (see RxODE events for more information), ability to supply both an omega and sigma matrix as well as adding as a thetaMat to R to simulate with uncertainty in the omega, sigma and theta matrices; see RxODE simulation vignette.

13.5 Integrating RxODE models in your package

Using Pre-compiled models in your packages 13.5.1

If you have a package and would like to include pre-compiled RxODE models in your package it is easy to create the package. You simple make the package with the rxPkg() command.

```
library(RxODE);
#### Now Create a model
idr <- RxODE({</pre>
    C2 = centr/V2;
    C3 = peri/V3;
    d/dt(depot) =-KA*depot;
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) =
                                      Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
})
#### You can specify as many models as you want to add
rxPkg(idr, package="myPackage"); ## Add the idr model to your package
```

This will:

- Add the model to your package; You can use the package data as idr once the package loads
- Add the right package requirements to the DESCRIPTION file. You will want to update this to describe the package and modify authors, license etc.
- Create skeleton model documentation files you can add to for your package documentation. In this case it would be the file idr-doc. R in your R directory
- Create a configure and configure.win script that removes and regenerates the src directory based on whatever version of RxODE this is compiled against. This should be modified if you plan to have your own compiled code, though this is not suggested.
- You can write your own R code in your package that interacts with the RxODE object so you can distribute shiny apps and similar things in the package context.

Once this is present you can add more models to your package by rxUse(). Simply compile the RxODE model in your package then add the model with rxUse()

```
rxUse(model)
```

Now both model and idr are in the model library. This will also create model-doc.R in your R directory so you can document this model.

You can then use devtools methods to install/test your model

```
devtools::load_all() # Load all the functions in the package
devtools::document() # Create package documentation
devtools::install() # Install package
devtools::check() # Check the package
devtools::build() # build the package so you can submit it to places like CRAN
```

13.5.2 Using Models in a already present package

To illustrate, lets start with a blank package

```
library(RxODE)
library(usethis)
pkgPath <- file.path(rxTempDir(), "MyRxModel")</pre>
create_package(pkgPath);
use_gpl3_license("Matt")
use_package("RxODE", "LinkingTo")
use_package("RxODE", "Depends") ## library(RxODE) on load; Can use imports instead.
use_roxygen_md()
##use_readme_md()
library(RxODE);
#### Now Create a model
idr <- RxODE({</pre>
   C2 = centr/V2;
    C3 = peri/V3;
    d/dt(depot) =-KA*depot;
    d/dt(centr) = KA*depot - CL*C2 - Q*C2 + Q*C3;
    d/dt(peri) =
                                      Q*C2 - Q*C3;
    d/dt(eff) = Kin - Kout*(1-C2/(EC50+C2))*eff;
});
rxUse(idr); ## Add the idr model to your package
rxUse(); # Update the compiled RxODE sources for all of your packages
```

The rxUse() will: - Create RxODE sources and move them into the package's src/directory. If there is only R source in the package, it will also finish off the directory with an library-init.c which registers all the RxODE models in the package for use in R. - Create stub R documentation for each of the models your are including in your package. You will be able to see the R documentation when loading your package by the standard? interface.

You will still need to: - Export at least one function. If you do not have a function that you wish to export, you can add a re-export of RxODE using roxygen as follows:

##' @importFrom RxODE RxODE
##' @export
RxODE::RxODE

If you want to use Suggests instead of Depends in your package, you way want to export all of RxODE's normal routines

##' @importFrom RxODE RxODE ##' @export RxODE::RxODE ##' @importFrom RxODE et ##' @export RxODE::et ##' @importFrom RxODE etRep ##' @export RxODE::etRep ##' @importFrom RxODE etSeq ##' @export RxODE::etSeq ##' @importFrom RxODE as.et ##' @export RxODE::as.et ##' @importFrom RxODE eventTable ##' @export RxODE::eventTable ##' @importFrom RxODE add.dosing ##' @export RxODE::add.dosing ##' @importFrom RxODE add.sampling ##' @export RxODE::add.sampling ##' @importFrom RxODE rxSolve ##' @export RxODE::rxSolve

```
##' @importFrom RxODE rxControl
##' @export
RxODE::rxControl
##' @importFrom RxODE rxClean
##' @export
RxODE::rxClean
##' @importFrom RxODE rxUse
##' @export
RxODE::rxUse
##' @importFrom RxODE rxShiny
##' @export
RxODE::rxShiny
##' @importFrom RxODE genShinyApp.template
##' @export
RxODE::genShinyApp.template
##' @importFrom RxODE cvPost
##' @export
RxODE::cvPost
### This is actually from `magrittr` but allows less imports
##' @importFrom RxODE %>%
##' @export
RxODE:: `%>%`
```

• You also need to instruct R to load the model library models included in the model's dll. This is done by:

```
### In this case `rxModels` is the package name
##' @useDynLib rxModels, .registration=TRUE
```

If this is a R package with RxODE models and you do not intend to add any other compiled sources (recommended), you can add the following configure scripts

```
#!/bin/sh
### This should be used for both configure and configure.win
echo "unlink('src', recursive=TRUE);RxODE::rxUse()" > build.R
${R_HOME}/bin/Rscript build.R
rm build.R
```

Depending on the check you may need a dummy autoconf script,

```
#### dummy autoconf script
#### It is saved to configure.ac
```

If you want to integrate with other sources in your Rcpp or C/Fortan based packages, you need to include rxModels-compiled.h and: - Add the define macro compiledModelCall to the list of registered .Call functions. - Register C interface to allow model solving by R_initO_rxModels_RxODE_models() (again rxModels would be replaced by your package name).

Once this is complete, you can compile/document by the standard methods:

```
devtools::load_all()
devtools::document()
devtools::install()
```

If you load the package with a new version of RxODE, the models will be recompiled when they are used.

However, if you want the models recompiled for the most recent version of RxODE, you simply need to call rxUse() again in the project directory followed by the standard methods for install/create a package.

```
devtools::load_all()
devtools::document()
devtools::install()
```

Note you do not have to include the RxODE code required to generate the model to regenerate the RxODE c-code in the src directory. As with all RxODE objects, a summary will show one way to recreate the same model.

An example of compiled models package can be found in the rxModels repository.

13.6 Stiff ODEs with Jacobian Specification

13.6.0.1 Stiff ODEs with Jacobian Specification

Occasionally, you may come across a **stiff** differential equation, that is a differential equation that is numerically unstable and small variations in parameters cause different solutions to the ODEs. One way to tackle this is to choose a stiff-solver, or hybrid stiff solver (like the default LSODA). Typically this is enough. However exact Jacobian solutions may increase the stability of the ODE. (Note the Jacobian is the derivative of the ODE specification with respect to each variable). In RXODE

you can specify the Jacobian with the df(state)/dy(variable) = statement. A classic ODE that has stiff properties under various conditions is the Van der Pol differential equations.

In RxODE these can be specified by the following:

```
library(RxODE)
Vtpol2 <- RxODE({</pre>
   d/dt(y) = dy
   d/dt(dy) = mu*(1-y^2)*dy - y
##### Jacobian
   df(y)/dy(dy) = 1
   df(dy)/dy(y) = -2*dy*mu*y - 1
   \frac{df(dy)}{dy}(dy) = mu*(1-y^2)
##### Initial conditions
   y(0) = 2
   dy(0) = 0
##### mu
   mu = 1 ## nonstiff; 10 moderately stiff; 1000 stiff
})
et <- eventTable();</pre>
et$add.sampling(seq(0, 10, length.out=200));
et$add.dosing(20, start.time=0);
s1 <- Vtpol2 %>% solve(et, method="lsoda")
print(s1)
#> ______ Solved RxODE object ______
#> -- Parameters ($params): -----
#> mu
#> 1
#> -- Initial Conditions ($inits): ------
\#> y dy
#> 2 0
#> -- First part of data (object): ------
#> # A tibble: 200 x 3
    time y dy
#> <dbl> <dbl> <dbl>
      22 0
#> 1 0
#> 2 0.0503 22.0 -0.0456
#> 3 0.101 22.0 -0.0456
#> 4 0.151 22.0 -0.0456
#> 5 0.201 22.0 -0.0456
#> 6 0.251 22.0 -0.0456
#> # ... with 194 more rows
```

```
#> _____
```

While this is not stiff at mu=1, mu=1000 is a stiff system

```
s2 <- Vtpol2 %>% solve(c(mu=1000), et)
print(s2)
#> ______ Solved RxODE object _____
#> -- Parameters ($params): ------
#> mu
#> 1000
#> -- Initial Conditions ($inits): ------
#> y dy
#> 2 0
#> -- First part of data (object): ------
#> # A tibble: 200 x 3
\#> time y dy \#> <dbl> <dbl> <math><dbl>
              <db1>
#> 1 0 22 0
#> 2 0.0503 22.0 -0.0000455
#> 3 0.101 22.0 -0.0000455
#> 4 0.151 22.0 -0.0000455
#> 5 0.201 22.0 -0.0000455
#> 6 0.251 22.0 -0.0000455
#> # ... with 194 more rows
#> _____
```

While this is easy enough to do, it is a bit tedious. If you have RxODE setup appropriately, that is you have:

- Python installed in your system
- sympy installed in your system
- SnakeCharmR installed in R

You can use the computer algebra system sympy to calculate the Jacobian automatically.

This is done by the RxODE option calcJac option:

```
Vtpo1 <- RxODE({
    d/dt(y) = dy
    d/dt(dy) = mu*(1-y^2)*dy - y
##### Initial conditions
    y(0) = 2
    dy(0) = 0</pre>
```