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 minilanguage 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 5) related to Leibnitz notation.
- The event data, which can be:
 - a NONMEM or deSolve compatible data frame (Chapter ??), or
 - created with et() or EventTable() for easy simulation of events(Chapter ??)
 - 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 uncertanty can be used to simulate while solving.

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

This book was assembled on Fri Dec $11\ 23:15:40\ 2020$ with RxODE version 1.0.0.0 automatically by github actions.

Related R packages

2.1 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 RxODE maintainers has had positive interactions with all of the ODE-solving pharmacometric projects listed.

2.2 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.

2.2.1 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.

2.2.2 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")
```

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:

3.0.1 Windows

In windows you may simply use install rto install rtools:

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

Alternatively you can download and install rtools directly.

3.0.2 Mac OSX

To get the most speed you need OpenMP enabled and compile RxODE against that binary. Here is some discussion about this:

https://mac.r-project.org/openmp/

3.0.3 Linux

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

3.1 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

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)
library(units)

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>
```

4.1 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
```

4.2 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 %>%

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

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

4.3 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

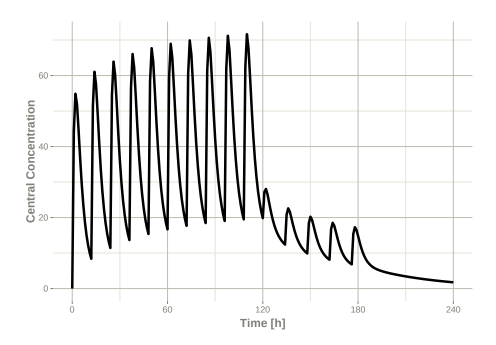
You can also solve this and create a RxODE data frame:

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

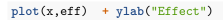
```
#> _____ Solved RxODE object _____
#> -- Parameters (x$params): -----
                       CL Q
#>
     V2
          V3 KA
                                  Kin
                                       Kout
#> 40.200 297.000  0.294  18.600  10.500  1.000
                                      1.000 200.000
#> -- Initial Conditions (x$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
#> 1
                     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
     3 51.9 4.46 4140. 2087. 1324. 1.23
#> 4
     4 44.5 5.98 3085. 1789. 1776. 1.23
#> 5
                2299. 1467. 2132. 1.21
     5 36.5 7.18
#> # ... with 235 more rows
#> ______
```

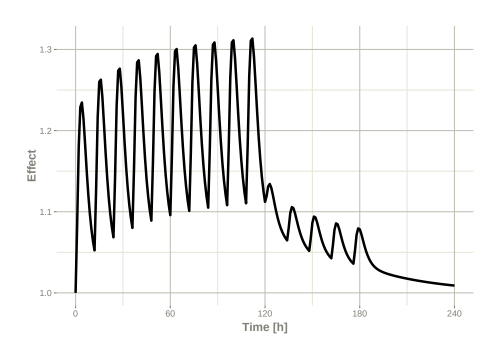
This returns a modified data frame. You can see the compartment values in the plot below:

```
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.

5.1 Example

5.2 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 vignette.
- 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, etc.

5.2.1 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.

5.3 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.

5.4 cmt() changing compartment numbers for states

The compartment order can be changed with the <code>cmt()</code> syntax in the model. To understand what the <code>cmt()</code> can do you need to understand how <code>RxODE</code> numbers the compartments.

Below is an example of how RxODE numbers compartments

5.4.1 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*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;
    # 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 +
```

VLU = (0.76 * WT/100)/1.051

#>

```
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 compart-
ment number(s)
summary(pbpk)
#> RxODE 1.0.0-0 model named rx 2e57f6731796bc01c8971d9f443861b6 model (ready).
#> DLL: /home/matt/.cache/R/RxODE/rx_2e57f6731796bc01c8971d9f443861b6__.rxd/rx_2e57f6731796bc01c8
#> NULL
#>
#> Calculated Variables:
                                "CLint" "KbBO"
#> [1] "KbBR"
                        "KbAD"
                                                         "CO"
                                                                 "QHT"
                "KbMU"
                                                 "KbRB"
                                                                         "QBR"
#> [10] "QMU"
                "QAD"
                        "QSK"
                                "QSP"
                                        "QPA"
                                                 "QLI"
                                                         "QST"
                                                                 "QGU"
                                                                         "QHA"
#> [19] "QBO"
                "QKI"
                                        "VLU"
                                                 "VHT"
                                                         "VBR"
                                                                 "VMU"
                                                                         "VAD"
                        "QRB"
                                "QLU"
#> [28] "VSK"
                "VSP"
                        "VPA"
                                "VLI"
                                        "VST"
                                                 "VGU"
                                                         "VBO"
                                                                 "VKI"
                                                                         "VAB"
                "VRB"
                                                        "KbSP"
#> [37] "VVB"
                                "KbLU"
                                        "KbHT"
                                                 "KbSK"
                                                                 "KbPA" "KbLI"
                        "fub"
#> [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
```

```
#>
       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)
       d/dt(Arterial Blood) = QLU * (Lungs/KbLU/VLU - Arterial Blood/VAB)
#>
#>
       d/dt(Venous_Blood) = QHT * Heart/KbHT/VHT + QBR * Brain/KbBR/VBR +
```

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.

5.4.2 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 1.0.0-0 model named rx_dfc43b162c4ff916310e6ca1def028ac model (ready).
#> x$state: Venous_Blood, Skin, Lungs, Heart, Brain, Muscles, Adipose, Spleen, Pancreas, Liver, S
#> x$params: 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.

5.4.3 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 1.0.0-0 model named rx_de6db2c40ddb8e5bf666a1a942c0c10a 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.
```