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Torsten: A Pharmacokinetic/Pharmacodynamic Model Library for Stan

User's Guide (Torsten Version 0.91.0, Stan version 2.33.1)

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#### Institutions

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### Individuals

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# Introduction

Stan is an open source probabilistic programing language designed primarily to do Bayesian data analysis [1]. It provides an expressive syntax for statistic modeling and contains an efficient variant of No U-Turn Sampler(NUTS), an adaptative Hamiltonian Monte Carlo algorithm that was proven more efficient than commonly used Monte Carlo Markov Chains (MCMC) samplers for complex high dimensional problems [2, 3].

Torsten is a collection of Stan functions to facilitate analysis of pharmacometric data. Given an event schedule and an ODE system, it calculates amounts in each compartment. The current version includes <sup>1</sup>:

- Specific linear compartment models:
  - One compartment model with first order absorption.
  - Two compartment model with elimination from and first order absorption into central compartment
- General linear compartment model described by a system of first-order <u>linear</u> Ordinary Differential Equations (ODEs).
- General compartment model described by a system of first order ODEs.
- Coupled model with PK forcing function described by a linear one or two compartment model and PD components solved by numerical ODE integration.

The models and data format are based on NONMEM ® <sup>2</sup>/NMTRAN/PREDPP conventions including:

- recursive calculation of model predictions, which permits piecewise constant covariate values.
- bolus or constant rate inputs into any compartment,
- single dose and multiple dose events,
- steady state dosing events,
- NMTRAN-compartible data items such as TIME, EVID, CMT, AMT, RATE, ADDL, II, and SS.

All real variable arguments in Torsten functions can be passed as Stan parameters.

# 1.1. Implementation summary

• Current Torsten v0.91.0 is based on Stan v2.33.1.

<sup>&</sup>lt;sup>1</sup>WARNING: The current version of Torsten is a *prototype*. It is being released for review and comment, and to support limited research applications. It has not been rigorously tested and should not be used for critical applications without further testing or cross-checking by comparison with other methods. We encourage interested users to try Torsten out and are happy to assist. Please report issues, bugs, and feature requests on our GitHub page.

<sup>&</sup>lt;sup>2</sup>NONMEM® is licensed and distributed by ICON Development Solutions.

- All functions are programmed in C++ and are compatible with the Stan math automatic differentiation library [4]
- One and two compartment models are based on analytical solutions of governing ODEs.
- General linear compartment models are based on semi-analytical solutions using the built-in matrix exponential function
- General compartment models are solved numerically using built-in ODE integrators in Stan. The tuning parameters of the solver are adjustable. The steady state solution is calculated using a numerical algebraic solver.
- Coupled model that has PK forcing function solved analytically and PD ODE components solved numerically.

# 1.2. Development plans

Our current plans for future development of Torsten include the following:

- Build a system to easily share packages of Stan functions (written in C++ or in the Stan language)
- Optimize Matrix exponential functions
  - Function for the action of Matrix Exponential on a vector
  - Hand-coded gradients
  - Special algorithm for matrices with special properties
- Develop new method for large-scale hierarchical models with costly ODE solving.
- Fix issue that arises when computing the adjoint of the lag time parameter (in a dosing compartment) evaluated at  $t_{\text{lag}} = 0$ .
- Extend formal tests
  - More unit tests and better CD/CI support.
  - Comparison with simulations from the R package mrgsolve and the software NONMEM®
  - Recruit non-developer users to conduct beta testing

# Changelog

- Version 0.91.1 <2024-12-01 Sun>
  - Changed
    - \* More model examples fixes.
    - \* Remove ODE group integrator functions.
    - \* Improve documentation.
- Version 0.91.0 < 2023-12-31 Sun>
  - Changed
    - \* Update model examples according to new array syntax.
    - \* Update to Stan version 2.33.1.
    - \* Removed experimental cross-chain feature.
    - \* Deprecation of old Torsten functions not using 'pmx\_' prefix.
- Version 0.90.0 <2022-05-18 Wed>
  - Changed
    - \* Update model examples according to new array syntax.
    - \* Update to Stan version 2.29.2.
- Version 0.89 < 2021-06-15 Tue>
  - Changed
    - \* New backend for ODE events solvers.
    - \* Use vector instead of array as ODE function state & return type.
    - \* Simplified ODE integrator naming, e.g. pmx ode bdf[ctrl].
    - \* Update to Stan version 2.27.0.
- Version 0.88 < 2020-12-18 Fri>
  - Added
    - \* Bioavailability, lag time, ODE real & integer data are optional in PMX function signatures.
    - \* Support all EVID options from NM-TRAN and mrgsolve.
    - \* Support steady-state infusion through multiple interdose intervals.
  - Changed
    - \* More efficient memory management of COVDES implementation.
    - \* Update of MPI framework to adapt multilevel paralleism.
    - \* Update to Stan version 2.25.0.
    - \* Use cmdstanr as R interface.
    - \* Stop supporting rstan as R interface.
- Version 0.87 < 2019-07-26 Fri>
  - Added
    - \* MPI dynamic load balance for Torsten's population ODE integrators
      - · pmx\_integrate\_ode\_group\_adams
      - pmx\_integrate\_ode\_group\_bdf

pmx\_integrate\_ode\_group\_rk45

To invoke dynamic load balance instead of default static balance for MPI, issue TORSTEN\_MPI=2 in make/local.

- \* Support RATE as parameter in pmx\_solve\_rk45/bdf/adams functions.
- Changed
  - \* Some fixes on steady-state solvers
  - \* Update to rstan version 2.19.2.
- Version 0.86 < 2019-05-15 Wed>
  - Added
    - \* Torsten's ODE integrator functions
      - · pmx\_integrate\_ode\_adams
      - pmx\_integrate\_ode\_bdf
      - pmx\_integrate\_ode\_rk45

and their counterparts to solve a population/group of subjects governed by an ODE

- · pmx\_integrate\_ode\_group\_adams
- pmx\_integrate\_ode\_group\_bdf
- · pmx\_integrate\_ode\_group\_rk45
- \* Torsten's population PMX solver functions for general ODE models
  - pmx\_solve\_group\_adams
  - · pmx\_solve\_group\_bdf
  - pmx\_solve\_group\_rk45
- \* Support time step ts as parameter in pmx\_integrate\_ode\_xxx solvers.
- Changed
  - \* Renaming Torsten functions in previous releases, the old-new name mapping is
    - $\cdot$  PKModelOneCpt  $\rightarrow$  pmx\_solve\_onecpt
    - $\cdot$  PKModelTwoCpt  $\rightarrow$  pmx\_solve\_onecpt
    - $\cdot$  linOdeModel  $\rightarrow$  pmx\_solve\_linode
    - $\cdot$  generalOdeModel\_adams  $\rightarrow$  pmx\_solve\_adams
    - $\cdot$  generalOdeModel\_bdf  $\rightarrow$  pmx\_solve\_bdf
    - · generalOdeModel\_rk45  $\rightarrow$  pmx\_solve\_rk45
    - $\cdot$  mixOde1CptModel\_bdf  $\rightarrow$  pmx\_solve\_onecpt\_bdf
    - $\cdot$  mixOde1CptModel\_rk45  $\rightarrow$  pmx\_solve\_onecpt\_rk45
    - $\cdot$  mixOde2CptModel\_bdf  $\rightarrow$  pmx\_solve\_twocpt\_bdf
    - · mixOde2CptModel\_rk45  $\rightarrow$  pmx\_solve\_twocpt\_rk45

Note that the new version of the above functions return the *transpose* of the matrix returned by the old versions, in order to improve memory efficiency. The old version are retained but will be deprecated in the future.

- \* Update to Stan version 2.19.1.
- Version 0.85 < 2018-12-04 Tue>
  - Added
    - \* Dosing rate as parameter
  - Changed
  - Update to Stan version 2.18.0.

- $\bullet$  Version 0.84 <2018-02-24 Sat>
  - Added
    - \* Piecewise linear interpolation function.
    - \* Univariate integral functions.
  - Changed
    - \* Update to Stan version 2.17.1.
    - \* Minor revisions to User Manual.
    - \* Bugfixes.
- Version 0.83 < 2017-08-02 Wed>
  - Added
    - \* Work with TorstenHeaders
    - \* Each chain has a different initial estimate
  - Changed
    - \* User manual
    - \* Fix misspecification in ODE system for TwoCpt example.
    - \* Other bugfixes
- Version 0.82 < 2017-01-29 Sun>
  - Added
    - \* Allow parameter arguments to be passed as 1D or 2D arrays
    - \* More unit tests
    - \* Unit tests check automatic differentiation against finite differentiation.
  - Changed
    - \* Split the parameter argument into three arguments: pMatrix (parameters for the ODEs note: for linOdeModel, pMatrix is replaced by the constant rate matrix K), biovar (parameters for the biovariability), and tlag (parameters for the lag time).
    - \* bugfixes
- Version 0.81 <2016-09-27 Tue>
  - Added
    - \* linCptModel (linear compartmental model) function
- Version 0.80a < 2016-09-21 Wed>
  - Added
    - \* check\_finite statements in pred\_1 and pred\_2 to reject metropolis proposal if initial conditions are not finite

# Installation

Currently Torsten is based on a forked version of Stan and hosted on GitHub

• https://github.com/metrumresearchgroup/Torsten

The latest v0.91.0 is compatible with Stan v2.33.1. Torsten can be accessed from command line for cmdstan interface and cmdstanr (https://mc-stan.org/cmdstanr/) for R interface. It requires a modern C++11 compiler as well as a Make utility. See [5] for details of installation and required toolchain. In particular, we recommend the following versions of C++ compilers:

- Linux: g++>=7.5 or clang >=8.0,
- macOS: the XCode version of clang,
- Windows: g++8.1 (available with RTools 4.0).

On windows, the Make utility mingw32-make can be installed as part of RTools.

#### 3.1. Command line interface

The command line interface cmdstan is available along with Torsten and can be found at Torsten/cmdstan.

After installation, one can use the following command to build a Torsten model\_name in model\_path

```
cd Torsten/cmdstan
make model_path/model_name # replace "make" with "mingw32-make" on

→ Windows platform
```

### 3.2. R interface

After installing cmdstanr from https://mc-stan.org/cmdstanr/, use the following command to set path

```
cmdstanr::set_cmdstan_path("Torsten/cmdstan")
```

Then one can follow https://mc-stan.org/cmdstanr/articles/cmdstanr.html to compile and run Torsten models.

# 3.3. MPI support

Torsten's MPI support is of a different flavour than reduce\_sum found in Stan. To be able to utilize MPI parallelisation, one first needs to ensure an MPI library such as

- https://www.mpich.org/downloads/
- https://www.open-mpi.org/software/ompi/

3.4. TESTING

is available. Torsen's implementation is tested on both MPICH and OpenMPI. To use MPI-supported population/group solvers, add/edit make/local

```
# path to MPI headers
CXXFLAGS += -isystem /usr/local/include
# if you are using Metrum's metworx platform, add MPICH3's
# headers with
# CXXFLAGS += -isystem /usr/local/mpich3/include
```

Note that currently TORSTEN\_MPI and STAN\_MPI flags conflict on processes management and cannot be used in a same Stan model, and MPI support is only available through cmdstan interface.

# 3.4. Testing

Models in example-models directory are for tutorial and demonstration. The following shows how to build and run the two-compartment model using cmdstanr, and use bayesplot to examine posterior density of CL.

# Using Torsten

The reader should have a basic understanding of how Stan works. In this section we go through the different functions Torsten adds to Stan. The code for the examples can be found at Torsten's example models.

# 4.1. Events specification

Torsten's functions are prefixed with pmx\_. For some of their arguments we adopt NM-TRAN format for events specification(Table 4.1).

TABLE 4.1. NM-TRAN compatible event specification arguments. All arrays should have the same length corresponding to the number of events.

Argument Name	Definition	Stan data type
time	event time	real[]
amt	dosing amount	real[]
rate	infusion rate	real[]
ii	interdose interval	real[]
evid	event ID	int[]
cmt	event compartment	int[]
addl	additionial identical doses	int[]
SS	steady-state dosing flag	int[]

All the real[] arguments above are allowed to be parameters in a Stan model. In addition, Torsten functions support optional arguments and overloaded signatures. Optional arguments are indicated by surrounding square bracket []. Table below shows three commonly used PMX model arguments that support overloading. In the rest of this document we assume this convention unless indicated otherwise.

TABLE 4.2. PMX model parameter overloadings. One can use 1d array real 1 to indicate constants of all events, or 2d array real 1, 1 so that the *i*th row of the array describes the model arguments for time interval  $(t_{i-1}, t_i)$ , and the number of the rows equals to the size of time.

Argument Name	Definition	Stan data type	Optional
theta	model parameters	real[] or real[ ,	] N
biovar	bioavailability fraction	real[] or real[ ,	Y (default to 1.0)
tlag	lag time	<pre>real[] or real[ ,</pre>	Y (default to 0.0)

# 4.2. One Compartment Model

**4.2.1. Description.** Function pmx\_solve\_onecpt solves a one-compartment PK model (Figure 4.1). The model obtains the mass  $(y_1, y_2)$  in each compartment by solving the ordinary differential equations (ODEs)

$$y_1' = -k_a y_1, (4.1)$$

$$y_2' = k_a y_1 - \left(\frac{CL}{V_2} + \frac{Q}{V_2}\right) y_2. \tag{4.2}$$

The plasma concentrations of parent drug in the central compartment can then be calculated as  $c = y_2/V_2$ .

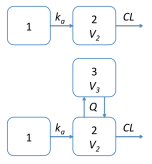


FIGURE 4.1. One and two compartment models with first order absorption implemented in Torsten.

#### 4.2.2. Usage.

- **4.2.3.** Arguments. See Tables in Section 4.1.
- **4.2.4.** Return value. An ncmt-by-nt matrix, where nt is the number of time steps and ncmt=2 is the number of compartments.

#### 4.2.5. Note.

- ODE Parameters theta should consist of CL,  $V_2$ ,  $k_a$ , in that order.
- biovar and tlag are optional, so that the following are allowed:

```
pmx_solve_onecpt(..., theta);
pmx_solve_onecpt(..., theta, biovar);
pmx_solve_onecpt(..., theta, biovar, tlag);
```

• Setting  $k_a = 0$  eliminates the first-order absorption.

## 4.3. Two Compartment Model

**4.3.1. Description.** Function pmx\_solve\_twocpt solves a two-compartment PK model (Figure 4.1). The model obtains the mass  $(y_1, y_2, y_3)$  in each compartment by solving the ODEs

$$y_1' = -k_a y_1 (4.3)$$

$$y_2' = k_a y_1 - \left(\frac{CL}{V_2} + \frac{Q}{V_2}\right) y_2 + \frac{Q}{V_3} y_3 \tag{4.4}$$

$$y_3' = \frac{Q}{V_2} y_2 - \frac{Q}{V_3} y_3 \tag{4.5}$$

The plasma concentrations of parent drug in the central compartment can then be calculated as  $c = y_2/V_2$ .

# 4.3.2. Usage.

- **4.3.3. Arguments.** See Tables in Section 4.1.
- **4.3.4. Return value.** An ncmt-by-nt matrix, where nt is the number of time steps and ncmt=3 is the number of compartments.

#### 4.3.5. Note.

- ODE Parameters theta consists of CL, Q,  $V_2$ ,  $V_3$ ,  $k_a$ .
- biovar and tlag are optional, so that the following are allowed:

```
pmx_solve_twocpt(..., theta);
pmx_solve_twocpt(..., theta, biovar);
pmx_solve_twocpt(..., theta, biovar, tlag);
```

• Setting  $k_a = 0$  eliminates the first-order absorption.

#### 4.4. General Linear ODE Model Function

**4.4.1. Description.** Function  $pmx\_solve\_linode$  solves a (piecewise) linear ODEs model with coefficients in form of matrix K

$$y'(t) = Ky(t) \tag{4.6}$$

For example, in a two-compartment model with first order absorption, K is

$$K = \begin{bmatrix} -k_a & 0 & 0\\ k_a & -(k_{10} + k_{12}) & k_{21}\\ 0 & k_{12} & -k_{21} \end{bmatrix}$$

$$(4.7)$$

where  $k_{10} = CL/V_2$ ,  $k_{12} = Q/V_2$ , and  $k_{21} = Q/V_3$ .

## 4.4.2. Usage.

## 4.4.3. Arguments.

- K System parameters. K can be either
  - a matrix for constant parameters in all events, or
  - an array of matrices matrix K[nt] so that the *i*th entry of the array describes the model parameters for time interval  $(t_{i-1}, t_i)$ , and the number of the rows equals to the number of event time nt.
- See Tables in Section 4.1 for the rest of arguments.
- **4.4.4.** Return value. An n-by-nt matrix, where nt is the number of time steps and n is the number of rows(columns) of square matrix K.

#### 4.5. General ODE Model Function

**4.5.1. Description.** Function pmx\_solve\_adams, pmx\_solve\_bdf, and pmx\_solve\_rk45 solve a first-order ODE system specified by user-specified right-hand-side function ODE\_rhs f

$$y'(t) = f(t, y(t))$$

In the case where the rate vector r is non-zero, this equation becomes:

$$y'(t) = f(t, y(t)) + r$$

# 4.5.2. Usage.

Here [adams || rk45 || bdf] indicates the function name can use any of the three suffixes. See below.

# 4.5.3. Arguments.

• ODE\_rhs ODE right-hand-side f. It should be defined in functions block and has the following format

Here t is time, y the unknowns of ODE, param the parameters, dat\\_r the real data, dat\\_i the integer data. param, dat\\_r, and dat\\_i are from the entry of theta, x\_r, and x\_i corresponding to t, respectively. f should not include dosing rates in its definition, as Torsten automatically update f when the corresponding event indicates infusion dosage.

• nCmt The number of compartments, equivalently, the dimension of the ODE system.

- x\_r 2d array real data to be passed to ODE RHS. If specified, its 1st dimension should have the same size as time.
- x\_i 2d array integer data to be passed to ODE RHS. If specified, its 1st dimension should have the same size as time.
- rel\_tol The relative tolerance for numerical integration, default to 1.0E-6.
- abs\_tol The absolute tolerance for numerical integration, default to 1.0E-6.
- max\_step The maximum number of steps in numerical integration, default to 10<sup>6</sup>.
- as\_rel\_tol The relative tolerance for algebra solver for steady state solution, default to 1.0E-6.
- as\_abs\_tol The absolute tolerance for algebra solver for steady state solution, default to 1.0E-6.
- as\_max\_step The maximum number of interations in algebra solver for steady state solution, default to 10<sup>2</sup>.
- See Tables in Section 4.1 for the rest of arguments.

# **4.5.4.** Return value. An nCmt-by-nt matrix, where nt is the size of time.

### 4.5.5. Note.

- See Section 4.8 for different types of integrator and general guidance.
- See Section 4.8 for comments on accuracy and tolerance.
- The default values of atol, rtol, and max\_step are based on a limited amount of PKPD test problems and should not be considered as universally applicable. We strongly recommend user to set these values according to physical intuition and numerical tests. See also Section 4.8.
- With optional arguments indicated by square bracket, the following calls are allowed:

```
pmx_solve_[adams || rk45 || bdf](..., theta);
pmx_solve_[adams || rk45 || bdf](..., theta, rel_tol, abs_tol, max_step);
pmx_solve_[adams || rk45 || bdf](..., theta, rel_tol, abs_tol, max_step,
\hookrightarrow as rel tol, as abs tol, as max step);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, rel_tol, abs_tol,
\hookrightarrow max step);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, rel_tol, abs_tol,

→ max_step, as_rel_tol, as_abs_tol, as_max_step);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, tlag);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, tlag, rel_tol,

    abs_tol, max_step);

pmx_solve_[adams | rk45 | bdf](..., theta, biovar, tlag, rel_tol,
→ abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);
pmx_solve_[adams | | rk45 | | bdf](..., theta, biovar, tlag, x_r);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r, rel_tol,
\hookrightarrow abs_tol, max_step);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r, rel_tol,
→ abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);
pmx_solve_{[adams || rk45 || bdf](..., theta, biovar, tlag, x_r, x_i);
```

```
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r, x_i, \hookrightarrow rel_tol, abs_tol, max_step);
pmx_solve_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r, x_i, \hookrightarrow rel_tol, abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);
```

# 4.6. Coupled ODE Model Function

**4.6.1.** Description. When the ODE system consists of two subsystems in form of

$$y'_1 = f_1(t, y_1),$$
  
 $y'_2 = f_2(t, y_1, y_2),$ 

with  $y_1$ ,  $y_2$ ,  $f_1$ , and  $f_2$  being vector-valued functions, and  $y_1'$  independent of  $y_2$ , the solution can be accelerated if  $y_1$  admits an analytical solution which can be introduced into the ODE for  $y_2$  for numerical integration. This structure arises in PK/PD models, where  $y_1$  describes a forcing PK function and  $y_2$  the PD effects. In the example of a Friberg-Karlsson semi-mechanistic model(see below), we observe an average speedup of  $\sim 47 \pm 18\%$  when using the mix solver in lieu of the numerical integrator. In the context, currently the couple solver supports one- & two-compartment for PK model, and rk45 & bdf integrator for nonlinear PD model.

# 4.6.2. Usage.

# 4.6.3. Arguments.

• reduced\_ODE\_rhs The system numerically solve ( $y_2$  in the above discussion, also called the *reduced system* and node the number of equations in the <u>reduced</u> system. The function that defines a reduced system has an almost identical signature to that used for a full system, but takes one additional argument:  $y_1$ , the PK states, i.e. solution to the PK ODEs.

```
vector reduced_ODE_rhs(real t, vector y2, vector y1, real[] theta, \hookrightarrow real[] x_r, int[] x_i)
```

- nCmt The number of compartments. Equivalently, the dimension of the ODE system
- rel\_tol The relative tolerance for numerical integration, default to 1.0E-6.
- abs\_tol The absolute tolerance for numerical integration, default to 1.0E-6.
- max\_step The maximum number of steps in numerical integration, default to 10<sup>6</sup>.
- See Tables in Section 4.1 for the rest of arguments.

**4.6.4.** Return value. An (nPk + nOde) × nt matrix, where nt is the size of time, and nPk equals to 2 in pmx\_solve\_onecpt\_ functions and 3 in pmx\_solve\_twocpt\_ functions.

# 4.7. General ODE-based Population Model Function

**4.7.1. Description.** All the preivous functions solves for a single sunject. Torsten also provides population modeling counterparts for ODE solutions. The functions solve for a population that share an ODE model but with subject-level parameters and event specifications and have similar signatures to single-subject functions, except that now events arguments time, amt, rate, ii, evid, cmt, addl, ss specifies the entire population, one subject after another.

## 4.7.2. Usage.

Here [adams || rk45 || bdf] indicates the function name can be of any of the three suffixes. See Section 4.8.

## 4.7.3. Arguments.

- ODE rhs Same as in Section 4.8.
- time, amt, rate, ii, evid, cmt, addl, ss 2d-array arguments that describe data record for the entire population (see also Tables in Section 4.1). They must have same size in the first dimension. Take evid for example. Let N be the population size, then evid[1,] to evid[n1,] specifies events ID for subject 1, evid[n1 + 1,] to evid[n1 + n2,] for subject 2, etc. With  $n_i$  being the number of events for subject i,  $i = 1, 2, \ldots, N$ , the size of evid's first dimension is  $\sum_i n_i$ .
- len The length of data for each subject within the above events arrays. The size of len equals to population size N.
- nCmt The number of compartments. Equivalently, the dimension of the ODE system.
- x\_r 2d array real data to be passed to ODE RHS. If specified, its 1st dimension should have the same size as time.
- x\_i 2d array integer data to be passed to ODE RHS. If specified, its 1st dimension should have the same size as time.
- rel\_tol The relative tolerance for numerical integration, default to 1.0E-6.
- abs\_tol The absolute tolerance for numerical integration, default to 1.0E-6.
- max\_step The maximum number of steps in numerical integration, default to 10<sup>6</sup>.
- as\_rel\_tol The relative tolerance for algebra solver for steady state solution, default to 1.0E-6.
- as\_abs\_tol The absolute tolerance for algebra solver for steady state solution, default to 1.0E-6.
- as\_max\_step The maximum number of interations in algebra solver for steady state solution, default to 10<sup>2</sup>.

**4.7.4. Return value.** An nCmt-by-nt matrix, where nt is the total size of events  $\sum_{i} n_{i}$ .

#### 4.7.5. Note.

- Similar to single-subject solvers, three numerical integrator are provided:
  - pmx\_solve\_group\_adams: Adams-Moulton method,
  - pmx\_solve\_group\_bdf: Backward-differentiation formular,
  - pmx\_solve\_group\_rk45: Runge-Kutta 4/5 method.
- With optional arguments indicated by square bracket, the following calls are allowed:

```
pmx_solve_group_[adams || rk45 || bdf](..., theta);
pmx_solve_group_[adams || rk45 || bdf](..., theta, rel_tol, abs_tol,
\hookrightarrow max_step);
pmx_solve_group_[adams || rk45 || bdf](..., theta, rel_tol, abs_tol,

→ max_step, as_rel_tol, as_abs_tol, as_max_step);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, rel_tol,

    abs_tol, max_step);

pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, rel_tol,
→ abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, rel_tol,

    abs_tol, max_step);

pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, rel_tol,

    abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);

pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r,

    rel_tol, abs_tol, max_step);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r,

    rel_tol, abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r,
\hookrightarrow x_i);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r,

    x_i, rel_tol, abs_tol, max_step);
pmx_solve_group_[adams || rk45 || bdf](..., theta, biovar, tlag, x_r,

→ x_i, rel_tol, abs_tol, max_step, as_rel_tol, as_abs_tol, as_max_step);
```

• The group solvers support parallelisation through Message Passing Interface (MPI). One can access this feature through cmdstan or cmdstanr interface.

Here n denotes number of MPI processes, so that N ODE systems (each specified by a same RHS function and subject-dependent events) are distributed to and solved by n processes evenly. Note that to access this feature user must have MPI installed, and some MPI installation may require set additional compiler arguments, such as CXXLFAGS and LDFLAGS.

## 4.8. ODE integrator function

**4.8.1. Description.** Torsten provides its own implementation of ODE solvers that solves

$$y'(t) = f(t, y(t)), \quad y(t_0) = y_0$$

for y. These solvers are customized for Torsten applications and different from those found in Stan. The general ODE PMX solvers in previous sections are internally powered by these functions.

# 4.8.2. Usage.

#### 4.8.3. Arguments.

 $\bullet$  ODE\_rhs Function that specifies the right-hand-side f. It should be defined in functions block and has the following format

```
vector = f(real t, vector y, ...) {}
```

Here t is time, y the unknowns of ODE, ... the arbitrary number number (and type) of parameters.

- y0 Initial condition  $y_0$ .
- t0 Initial time  $t_0$ .
- ts Output time when solution is seeked.
- ... Parameters to be passed to ODE\_rhs function.
- rtol Relative tolerance, default to 1.e-6(rk45) and 1.e-8(adams and bdf).
- atol Absolute tolerance, default to 1.e-6(rk45) and 1.e-8(adams and bdf).
- max\_step Maximum number of steps allowed between neighboring time in ts, default to 100000.

**4.8.4.** Return value. A n-array of nd-vectors, where n is the size of ts and nd the dimension of the system.

#### 4.8.5. Note.

- Four numerical integrator are provided:
  - pmx\_ode\_adams: Adams-Moulton method,
  - pmx\_ode\_bdf: Backward-differentiation formular,
  - pmx\_ode\_rk45: Runge-Kutta 4/5 method.
  - pmx\_ode\_ckrk: Cash-Karp 4/5 method.

When not equipped with further understanding of the ODE system, as a rule of thumb we suggest user try rk45 integrator first, bdf integrator when the system is suspected to be stiff, and adams and ckrk when a non-stiff system needs to be solved with higher accuracy/smaller tolerance, or long integration time.

• All four integrators support adaptive stepping. To achieve that, at step i estimated error  $e_i$  is calculated and compared with given tolerance so that

$$e_i < \|\text{rtol} \times \tilde{y} + \text{atol}\|$$
 (4.8)

Here  $\tilde{y}$  is the numerical solution of y at current step and  $\|\cdot\|$  indicates certain norm. When the above check fails, the solver attempts to reduce step size and retry. The default values of atol, rtol, and max\_step are based on Stan's ODE functions and should not be considered as optimal. User should make problem-dependent decision on rtol and atol, according to estimated scale of the unknowns, so that the error would not affect inference on statistical variance of quantities that enter the Stan model. In particular, when an unknown can be neglected below certain threshold without affecting the rest of the dynamic system, setting atol greater than that threshold will avoid spurious and error-prone computation. See [6] and 1.4 of [7] for details.

### 4.9. ODE group integrator Function

**4.9.1. Description.** All the preivous functions solves for a single ODE system. Torsten also provides group modeling counterparts for ODE integrators. The functions solve for a group of ODE systems that share an ODE RHS but with different parameters. They have similar signatures to single-ODE integration functions.

## 4.9.2. Usage.

```
matrix pmx_integrate_ode_group_[adams || rk45 || bdf](ODE_system, real[, \hookrightarrow ] y0, real t0, int[] len, real[] ts, real[, ] theta, real[, ] x_r, \hookrightarrow int[, ] x_i, [ real rtol, real atol, int max_step ]);
```

Here [adams || rk45 || bdf] indicates the function name can be of any of the three suffixes. See Section 4.8.

- ODE\_rhs Function that specifies the right-hand-side f. See Section 4.8.
- y0 Initial condition  $y_0$  for each subsystem in the group. The first dimension equals to the size of the group.
- t0 Initial time  $t_0$ .

- len A vector that contains the number of output time points for each subsystem. The length of the vector equals to the size of the group.
- ts Output time when solution is seeked, consisting of ts of each subsystem concatenated.
- theta 2d-array parameters to be passed to ODE\_rhs function. Each row corresponds to one subsystem.
- x\_r 2d-array real data to be passed to ODE\_rhs function. Each row corresponds to one subsystem.
- x\_i 2d-array integer data to be passed to ODE\_rhs function. Each row corresponds to one subsystem.
- rtol Relative tolerance.
- at ol Absolute tolerance.
- max\_step Maximum number of steps allowed between neighboring time in ts.

**4.9.3. Return value.** An n-by-nd matrix, where n is the size of ts and nd the dimension of the system.

#### 4.9.4. Note.

• With optional arguments indicated by square bracket, the following calls are allowed:

• The group integrators support parallelesation through Message Passing Interface (MPI). One can access this feature through cmdstan or cmdstanr interface.

```
library("cmdstanr")
cmdstan_make_local(cpp_options = list("TORSTEN_MPI" = "1",
    "TBB_CXX_TYPE"="gcc")) # "gcc" should be replaced by user's C
    compiler
rebuild_cmdstan()
mod <- cmdstan_model(path-to-model-file, quiet=FALSE,
    force_recompile=TRUE)
f <- mod$sample_mpi(data = ..., chains = 1, mpi_args = list("n" =
    number_of_processes), refresh = 200)</pre>
```

Here n denotes number of MPI processes, so that N ODE systems are distributed to and solved by n processes evenly. Note that to access this feature user must have MPI installed, and some MPI installation may require set additional compiler arguments, such as CXXLFAGS and LDFLAGS.

## 4.10. Univariate integral

```
real univariate_integral_rk45(f, t0, t1, theta, x_r, x_i)
```

```
real univariate_integral_bdf(f, t0, t1, theta, x_r, x_i)
```

Based on the ODE solver capability in Stan, Torsten provides functions calculating the integral of a univariate function. The integrand function f must follow the signature

```
real f(real t, real[] theta, real[] x_r, int[] x_i) {
   /* ... */
}
```

## 4.11. Piecewise linear interpolation

```
real linear_interpolation(real xout, real[] x, real[] y)
```

```
real[] linear_interpolation(real[] xout, real[] x, real[] y)
```

Torsten also provides function  $linear_interpolation$  for piecewise linear interpolation over a set of x, y pairs. It returns the values of a piecewise linear function at specified values xout of the first function argument. The function is specified in terms of a set of x, y pairs. Specifically,  $linear_interpolation$  implements the following function

$$y_{\text{out}} = \begin{cases} y_1, & x_{\text{out}} < x_1 \\ y_i + \frac{y_{i+1} - y_i}{x_{i+1} - x_i} (x_{\text{out}} - x_i), & x_{\text{out}} \in [x_i, x_{i+1}) \\ y_n, & x_{\text{out}} \ge x_n \end{cases}$$

- The x values must be in increasing order, i.e.  $x_i < x_{i+1}$ .
- All three arguments may be data or parameters.

# **Examples**

All the PMX models in this chapter can be found in Torsten/example-models directory:

- Two-compartment model
- Two-compartment model by linear ODEs
- two-compartment model by numerical ODEs
- Joint PK/PD model
- Two-compartment popPK model
- Group of ODEs
- Effective compartment model
- PopPKPD model

## 5.1. Two-compartment model for single patient

We model drug absorption in a single patient and simulate plasma drug concentrations:

- Multiple Doses: 1250 mg, every 12 hours, for a total of 15 doses
- PK measured at 0.083, 0.167, 0.25, 0.5, 0.75, 1, 1.5, 2, 4, 6, 8, 10 and 12 hours after 1st, 2nd, and 15th dose. In addition, the PK is measured every 12 hours throughout the trial.

With the plasma concentration  $\hat{c}$  using 4.3, we simulate c according to:

$$\log(c) \sim N \left( \log(\widehat{c}), \sigma^2 \right)$$

$$(CL, Q, V_2, V_3, ka) = \left( 5 \text{ L/h}, 8 \text{ L/h}, 20 \text{ L}, 70 \text{ L}, 1.2 \text{ h}^{-1} \right)$$

$$\sigma^2 = 0.01$$

The data are generated using the R package mrgsolve [8].

Code below shows how Torsten function pmx\_solve\_twocpt can be used to fit the above model.

```
data{
  int<lower = 1> nt;  // number of events
  int<lower = 1> nObs;  // number of observation
  int<lower = 1> iObs[nObs];  // index of observation

// NONMEM data
  int<lower = 1> cmt[nt];
  int evid[nt];
  int addl[nt];
  int ss[nt];
  real amt[nt];
  real time[nt];
```

```
real rate[nt];
  real ii[nt];
  vector<lower = 0>[nObs] cObs; // observed concentration (Dependent
  → Variable)
}
transformed data{
 vector[nObs] logCObs = log(cObs);
  int nTheta = 5; // number of ODE parameters in Two Compartment Model
  int nCmt = 3; // number of compartments in model
}
parameters{
 real<lower = 0 > CL;
 real<lower = 0 > Q;
 real<lower = 0 > V1;
  real<lower = 0> V2;
 real<lower = 0> ka;
 real<lower = 0> sigma;
}
transformed parameters{
 real theta[nTheta]; // ODE parameters
 row_vector<lower = 0>[nt] cHat;
  vector<lower = 0>[nObs] cHatObs;
 matrix<lower = 0>[nCmt, nt] x;
  theta[1] = CL;
  theta[2] = Q;
  theta[3] = V1;
  theta[4] = V2;
  theta[5] = ka;
 x = pmx_solve_twocpt(time, amt, rate, ii, evid, cmt, addl, ss, theta);
  cHat = x[2, :] ./ V1; // we're interested in the amount in the second

→ compartment

  cHatObs = cHat'[iObs]; // predictions for observed data recors
}
model{
 // informative prior
 CL \sim lognormal(log(10), 0.25);
  Q \sim lognormal(log(15), 0.5);
 V1 \sim lognormal(log(35), 0.25);
  V2 \sim lognormal(log(105), 0.5);
 ka ~ lognormal(log(2.5), 1);
 sigma \sim cauchy(0, 1);
 logCObs ~ normal(log(cHatObs), sigma);
```

Four MCMC chains of 2000 iterations (1000 warmup iterations and 1000 sampling iterations) are simulated. 1000 samples per chain were used for the subsequent analyses. The MCMC history plots(Figure 5.1) suggest that the 4 chains have converged to common distributions for all of the key model parameters. The fit to the plasma concentration data (Figure 5.3) are in close agreement with the data, which is not surprising since the fitted model is identical to the one used to simulate the data. Similarly the parameter posterior density can be examined in Figure 5.2 and shows consistency with the values used for simulation. Another way to summarize the posterior is through cmdstanr's summary method.

```
## fit is a CmdStanMCMC object returned by sampling. See cmdstanr
  reference.
> pars = c("CL", "Q", "V1", "V2", "ka", "sigma")
> fit$summary(pars)
# A tibble: 6 x 10
  variable
            mean median
                               sd
                                     mad
                                               q5
                                                      q95
                                                          rhat ess bulk
  <dbl>
                   <dbl>
                           <dbl>
                                   <dbl>
                                            <dbl>
                                                    <dbl> <dbl>
                                                                    <dbl>
  <chr>
  \hookrightarrow <dbl>
                    4.83
                                           4.68
1 CL
            4.82
                          0.0910 0.0870
                                                    4.97
                                                           1.00
                                                                    1439.

→ 1067.

2 Q
            7.56
                    7.55
                           0.588
                                  0.586
                                           6.61
                                                    8.56
                                                           1.00
                                                                    1256.
\hookrightarrow 1235.
3 V1
                   21.1
                           2.50
                                  2.45
                                          17.1
                                                   25.3
                                                           1.00
                                                                    1057.
           21.1
\hookrightarrow 1177.
4 V2
           76.1
                   76.1
                           5.33
                                  4.93
                                          67.5
                                                   84.9
                                                           1.01
                                                                    1585.
5 ka
            1.23
                    1.23 0.175
                                  0.174
                                           0.958
                                                    1.52
                                                           1.00
                                                                    1070.
\hookrightarrow 1122.
            0.109 0.108 0.0117 0.0111
                                          0.0911 0.130
                                                                    1414.
6 sigma
                                                           1.01
  905.
```

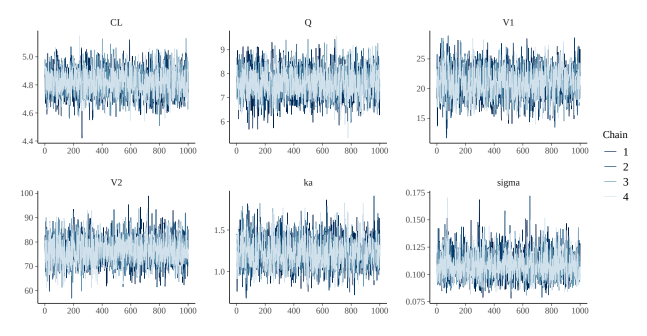


FIGURE 5.1. MCMC history plots for the parameters of a two compartment model with first order absorption (each color corresponds to a different chain)

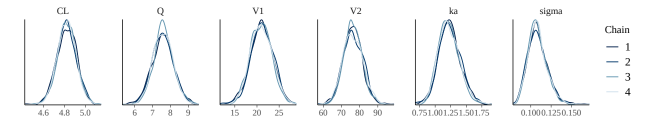


FIGURE 5.2. Posterior marginal densities of the Model Parameters of a two compartment model with first order absorption (each color corresponds to a different chain)

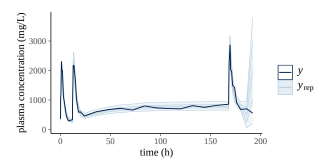


FIGURE 5.3. Predicted  $(y_{\text{rep}})$  and observed (y) plasma drug concentrations of a two compartment model with first order absorption.  $y_{\text{rep}}$  is shown with posterior median, 50%, 90% credible intervals.

## 5.2. Two-compartment model as a linear ODE model for single patient

Using pmx\_solve\_linode, the following example fits a two-compartment model with first order absorption. We omit data and model block as they are identical to 5.1 Example.

```
transformed data{
 row_vector[nObs] logCObs = log(cObs);
  int nCmt = 3;
 real biovar[nCmt];
  real tlag[nCmt];
 for (i in 1:nCmt) {
   biovar[i] = 1;
    tlag[i] = 0;
 }
}
parameters{
 real < lower = 0 > CL;
 real<lower = 0 > Q;
 real<lower = 0> V1;
 real<lower = 0 > V2;
 real<lower = 0> ka;
 real<lower = 0> sigma;
transformed parameters{
 matrix[3, 3] K;
 real k10 = CL / V1;
 real k12 = Q / V1;
 real k21 = Q / V2;
 row_vector<lower = 0>[nt] cHat;
 row_vector<lower = 0>[nObs] cHatObs;
 matrix<lower = 0>[3, nt] x;
 K = rep_matrix(0, 3, 3);
  K[1, 1] = -ka;
  K[2, 1] = ka;
  K[2, 2] = -(k10 + k12);
  K[2, 3] = k21;
  K[3, 2] = k12;
  K[3, 3] = -k21;
  x = pmx_solve_linode(time, amt, rate, ii, evid, cmt, addl, ss, K,
  cHat = row(x, 2) ./ V1;
  for(i in 1:nObs){
    cHatObs[i] = cHat[iObs[i]]; // predictions for observed data records
  }
}
```

# 5.3. Two-compartment model solved by numerical integrator for single patient

Using pmx\_solve\_rk45, the following example fits a two-compartment model with first order absorption. User-defined function ode\_rhs describes the RHS of the ODEs.

```
functions{
 vector ode_rhs(real t, vector x, real[] parms, real[] x_r, int[] x_i){
   real CL = parms[1];
   real Q = parms[2];
   real V1 = parms[3];
   real V2 = parms[4];
   real ka = parms[5];
   real k10 = CL / V1;
   real k12 = Q / V1;
   real k21 = Q / V2;
   vector[3] y;
   y[1] = -ka*x[1];
    y[2] = ka*x[1] - (k10 + k12)*x[2] + k21*x[3];
   y[3] = k12*x[2] - k21*x[3];
   return y;
 }
}
```

We omit data and model block as they are identical to 5.1 Example.

```
transformed data {
  row_vector[nObs] logCObs = log(cObs);
  int nTheta = 5;  // number of parameters
  int nCmt = 3;  // number of compartments
}
parameters {
 real<lower = 0 > CL;
 real<lower = 0 > Q;
  real<lower = 0> V1;
 real<lower = 0 > V2;
 real<lower = 0> ka;
  real<lower = 0> sigma;
}
transformed parameters {
 real theta[nTheta];
 row_vector<lower = 0>[nt] cHat;
 row_vector<lower = 0>[nObs] cHatObs;
 matrix < lower = 0 > [3, nt] x;
  theta[1] = CL;
  theta[2] = Q;
```

```
theta[3] = V1;
  theta[4] = V2;
  theta[5] = ka;
  x = pmx_solve_rk45(ode_rhs, 3, time, amt, rate, ii, evid, cmt, addl,
    ss, theta, 1e-5, 1e-8, 1e5);
  cHat = x[2, ] ./ V1;
  for(i in 1:nObs){
    cHatObs[i] = cHat[iObs[i]]; // predictions for observed data records
}
model {
```

### 5.4. Joint PK-PD model

Neutropenia is observed in patients receiving an ME-2 drug. Our goal is to model the relation between neutrophil counts and drug exposure. As shown in Figure 5.4, the Friberg-Karlsson Semi-Mechanistic model [9] couples a PK model with a PD effect to describe a delayed feedback mechanism that keeps the absolute neutrophil count (ANC) at the baseline in a circulatory compartment (Circ), and the drug's effect in reducing the proliferation rate (prol). The delay between prol and Circ is modeled using n transit comparements with mean transit time MTT =  $(n+1)/k_{\rm tr}$ , with  $k_{\rm tr}$  the transit rate constant. In the current example, we use the 4.3 for PK model, and set n = 3.

$$\log(ANC) \sim N(\log(y_{\rm circ}), \sigma_{\rm ANC}^2),$$
 (5.1)

$$y_{\text{circ}} = f_{\text{FK}}(\text{MTT}, \text{Circ}_0, \alpha, \gamma, c),$$
 (5.2)

where c is the drug concentration calculated from the PK model, and function  $f_{\rm FK}$  represents solving the following nonlinear ODE for  $y_{\rm circ}$ 

$$\frac{dy_{\text{prol}}}{dt} = k_{\text{prol}}y_{\text{prol}}(1 - E_{\text{drug}}) \left(\frac{\text{Circ}_0}{y_{\text{circ}}}\right)^{\gamma} - k_{\text{tr}}y_{\text{prol}},\tag{5.3}$$

$$\frac{dy_{\text{trans1}}}{dt} = k_{\text{tr}}y_{\text{prol}} - k_{\text{tr}}y_{\text{trans1}},\tag{5.4}$$

$$\frac{dy_{\text{trans1}}}{dt} = k_{\text{tr}}y_{\text{prol}} - k_{\text{tr}}y_{\text{trans1}},$$

$$\frac{dy_{\text{trans2}}}{dt} = k_{\text{tr}}y_{\text{trans1}} - k_{\text{tr}}y_{\text{trans2}},$$

$$\frac{dy_{\text{trans3}}}{dt} = k_{\text{tr}}y_{\text{trans2}} - k_{\text{tr}}y_{\text{trans3}},$$

$$\frac{dy_{\text{trans3}}}{dt} = k_{\text{tr}}y_{\text{trans2}} - k_{\text{tr}}y_{\text{trans3}},$$
(5.6)

$$\frac{dy_{\text{trans3}}}{dt} = k_{\text{tr}}y_{\text{trans2}} - k_{\text{tr}}y_{\text{trans3}},\tag{5.6}$$

$$\frac{dy_{\rm circ}}{dt} = k_{\rm tr} y_{\rm trans3} - k_{\rm tr} y_{\rm circ},\tag{5.7}$$

We use  $E_{\text{drug}} = \alpha c$  to model the linear effect of drug concentration in central compartment, with  $c = y_{\text{cent}}/V_{\text{cent}}$  based on PK solutions.

Since the ODEs specifying the Two Compartment Model (Equation (4.3)) do not depend on the PD ODEs (Equation (5.3)) and can be solved analytically using Torsten's pmx\_sol, ve twocpt function we can specify solve the system using a coupled solver function. We do not expect our system to be stiff and use the Runge-Kutta 4th/5th order integrator.

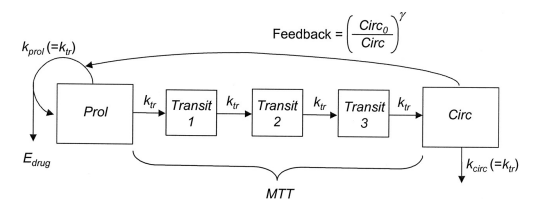


FIGURE 5.4. Friberg-Karlsson semi-mechanistic Model.

The model fitting is based on simulated data

(MTT, Circ<sub>0</sub>, 
$$\alpha$$
,  $\gamma$ ,  $k_{tr}$ ) = (125, 5.0, 3 × 10<sup>-4</sup>, 0.17)  
 $\sigma_{ANC}^2 = 0.001$ .

```
functions{
 vector FK_ODE(real t, vector y, vector y_pk, real[] theta, real[]
  → rdummy, int[] idummy) {
   /* PK variables */
   real VC = theta[3];
   /* PD variable */
   real mtt = theta[6];
   real circ0 = theta[7];
   real alpha
               = theta[8];
   real gamma = theta[9];
   real ktr
                = 4.0 / mtt;
                = y[1] + circ0;
   real prol
   real transit1 = y[2] + circ0;
   real transit2 = y[3] + circ0;
   real transit3 = y[4] + circ0;
               = fmax(machine_precision(), y[5] + circ0);
   real circ
   real conc
                = y_pk[2] / VC;
   real EDrug = alpha * conc;
   vector[5] dydt;
   dydt[1] = ktr * prol * ((1 - EDrug) * ((circ0 / circ)^gamma) - 1);
   dydt[2] = ktr * (prol - transit1);
   dydt[3] = ktr * (transit1 - transit2);
   dydt[4] = ktr * (transit2 - transit3);
   dydt[5] = ktr * (transit3 - circ);
   return dydt;
 }
}
```

```
data{
 int<lower = 1> nt;
  int<lower = 1> nObsPK;
 int<lower = 1> nObsPD;
 int<lower = 1> iObsPK[nObsPK];
  int<lower = 1> iObsPD[nObsPD];
 real<lower = 0> amt[nt];
 int<lower = 1> cmt[nt];
 int<lower = 0> evid[nt];
  real<lower = 0> time[nt];
 real<lower = 0> ii[nt];
 int<lower = 0> addl[nt];
 int<lower = 0> ss[nt];
  real rate[nt];
 vector<lower = 0>[nObsPK] cObs;
 vector<lower = 0>[nObsPD] neutObs;
  real<lower = 0> circ0Prior;
 real<lower = 0> circ0PriorCV;
 real<lower = 0> mttPrior;
 real<lower = 0> mttPriorCV;
 real<lower = 0> gammaPrior;
 real<lower = 0> gammaPriorCV;
 real<lower = 0> alphaPrior;
  real<lower = 0> alphaPriorCV;
}
transformed data{
 int nOde = 5;
 vector[nObsPK] logCObs;
 vector[nObsPD] logNeutObs;
 int nTheta = 9; // number of parameters
 int nIIV = 7; // parameters with IIV
 int n = 8;
                                    /* ODE dimension */
 real rtol = 1e-8;
 real atol = 1e-8;;
 int max_step = 100000;
 logCObs = log(cObs);
 logNeutObs = log(neutObs);
}
parameters{
 real<lower = 0 > CL;
 real<lower = 0 > Q;
 real<lower = 0 > VC;
 real<lower = 0 > VP;
 real<lower = 0> ka;
  real<lower = 0> mtt;
```

```
real<lower = 0> circ0;
  real<lower = 0> alpha;
  real<lower = 0> gamma;
  real<lower = 0> sigma;
 real<lower = 0> sigmaNeut;
 // IIV parameters
 cholesky_factor_corr[nIIV] L;
 vector<lower = 0>[nIIV] omega;
}
transformed parameters{
 row_vector[nt] cHat;
 vector<lower = 0>[nObsPK] cHatObs;
 row_vector[nt] neutHat;
 vector<lower = 0>[nObsPD] neutHatObs;
 real<lower = 0> theta[nTheta];
 matrix[nOde + 3, nt] x;
  real biovar[nTheta] = rep_array(1.0, nTheta);
 real tlag[nTheta] = rep_array(0.0, nTheta);
  theta[1] = CL;
  theta[2] = Q;
  theta[3] = VC;
  theta[4] = VP;
  theta[5] = ka;
  theta[6] = mtt;
  theta[7] = circ0;
  theta[8] = alpha;
  theta[9] = gamma;
 x = pmx_solve_twocpt_rk45(FK_ODE, nOde, time, amt, rate, ii, evid, cmt,

→ addl, ss, theta, biovar, tlag, rtol, atol, max_step);
 cHat = x[2, ] / VC;
 neutHat = x[8, ] + circ0;
 for(i in 1:nObsPK) cHatObs[i] = cHat[iObsPK[i]];
 for(i in 1:nObsPD) neutHatObs[i] = neutHat[iObsPD[i]];
}
model {
 // Priors
  CL ~ normal(0, 20);
      ~ normal(0, 20);
  VC ~ normal(0, 100);
      ~ normal(0, 1000);
 VP
     \sim normal(0, 5);
  ka
 sigma \sim cauchy(0, 1);
           ~ lognormal(log(mttPrior), mttPriorCV);
  mtt
  circ0 ~ lognormal(log(circ0Prior), circ0PriorCV);
          ~ lognormal(log(alphaPrior), alphaPriorCV);
```

```
gamma ~ lognormal(log(gammaPrior), gammaPriorCV);
sigmaNeut ~ cauchy(0, 1);

// Parameters for Matt's trick
L ~ lkj_corr_cholesky(1);
omega ~ cauchy(0, 1);

// observed data likelihood
logCObs ~ normal(log(cObs), sigma);
logNeutObs ~ normal(log(neutObs), sigmaNeut);
}
```

# 5.5. Two-compartment population model

Using pmx\_solve\_group\_bdf, the following example fits a two-compartment population model.

```
functions {
  // define ODE system for two compartmnt model
  real[] twoCptModelODE(real t,
                         real[] x,
                         real[] parms,
                         real[] rate, // in this example, rate is treated
                         \hookrightarrow as data
                         int[] dummy) {
    // Parameters
    real CL = parms[1];
    real Q = parms[2];
    real V1 = parms[3];
    real V2 = parms[4];
    real ka = parms[5];
    // Re-parametrization
    real k10 = CL / V1;
    real k12 = Q / V1;
    real k21 = Q / V2;
    // Return object (derivative)
    real y[3]; // 1 element per compartment of
                // the model
    // PK component of the ODE system
    y[1] = -ka * x[1];
    y[2] = ka*x[1] - (k10 + k12)*x[2] + k21*x[3];
    y[3] = k12*x[2] - k21*x[3];
    return y;
  }
}
data{
```

```
int<lower = 1> np;  /* population size */
  int<lower = 1> nt; // number of events
  int<lower = 1> nObs; // number of observations
  int<lower = 1> iObs[nObs]; // index of observation
 // NONMEM data
  int<lower = 1> cmt[np * nt];
 int evid[np * nt];
 int addl[np * nt];
 int ss[np * nt];
 real amt[np * nt];
 real time[np * nt];
 real rate[np * nt];
 real ii[np * nt];
 real<lower = 0> cObs[np*nObs]; // observed concentration (dependent
  → variable)
transformed data {
 real logCObs[np*nObs];
 int<lower = 1> len[np];
 int<lower = 1> len_theta[np];
 int<lower = 1> len_biovar[np];
 int<lower = 1> len_tlag[np];
 int nTheta = 5;  // number of parameters
 int nCmt = 3; // number of compartments
 real biovar[np * nt, nCmt];
 real tlag[np * nt, nCmt];
 logCObs = log(cObs);
 for (id in 1:np) {
   for (j in 1:nt) {
     for (i in 1:nCmt) {
       biovar[(id - 1) * nt + j, i] = 1;
       tlag[(id - 1) * nt + j, i] = 0;
    }
   }
   len[id] = nt;
   len_theta[id] = nt;
   len biovar[id] = nt;
   len_tlag[id] = nt;
 }
}
parameters{
real<lower = 0 > CL[np];
real<lower = 0> Q[np];
 real<lower = 0> V1[np];
 real<lower = 0 > V2[np];
  real<lower = 0 > ka[np];
```

```
real<lower = 0> sigma[np];
}
transformed parameters{
 real theta[np * nt, nTheta];
 vector<lower = 0>[nt] cHat[np];
  real<lower = 0> cHatObs[np*nObs];
  matrix[3, nt * np] x;
  for (id in 1:np) {
    for (it in 1:nt) {
      theta[(id - 1) * nt + it, 1] = CL[id];
      theta[(id - 1) * nt + it, 2] = Q[id];
      theta[(id - 1) * nt + it, 3] = V1[id];
      theta[(id - 1) * nt + it, 4] = V2[id];
      theta[(id - 1) * nt + it, 5] = ka[id];
  }
  x = pmx\_solve\_group\_bdf(twoCptModelODE, 3, len,
                           time, amt, rate, ii, evid, cmt, addl, ss,
                           theta, biovar, tlag);
  for (id in 1:np) {
    for (j in 1:nt) {
      cHat[id][j] = x[2, (id - 1) * nt + j] ./ V1[id];
  }
  for (id in 1:np) {
    for(i in 1:nObs){
     cHatObs[(id - 1)*nObs + i] = cHat[id][iObs[i]]; // predictions for
      → observed data records
  }
}
model{
 // informative prior
  for(id in 1:np){
    CL[id] \sim lognormal(log(10), 0.25);
    Q[id] \sim lognormal(log(15), 0.5);
    V1[id] \sim lognormal(log(35), 0.25);
    V2[id] \sim lognormal(log(105), 0.5);
    ka[id] \sim lognormal(log(2.5), 1);
    sigma[id] \sim cauchy(0, 1);
    for(i in 1:nObs){
     \log CObs[(id - 1)*nObs + i] \sim normal(log(cHatObs[(id - 1)*nObs + i]))
      \hookrightarrow i]), sigma[id]);
  }
}
```

When the above model is compiled with MPI support (see Section MPI support), one can run it using within-chain parallelization:

Here nproc indicates the number of parallel processes participating ODE solution. For example, with np=8 for a population of 8, nproc=4 indicates solving 8 subjects' ODEs in parallel, with each process solving 2 subjects.

## 5.6. Lotka-Volterra group model

Using pmx\_integrate\_ode\_group\_rk45, the following example fits a Lotka-Volterra group model, based on Stan's case study.

```
functions {
  real[] dz dt(real t,
                           // time
              real[] theta, // parameters
              real[] x_r, // unused data
              int[] x_i) {
   real u = z[1];
   real v = z[2];
   real alpha = theta[1];
   real beta = theta[2];
   real gamma = theta[3];
   real delta = theta[4];
   real du_dt = (alpha - beta * v) * u;
   real dv_dt = (-gamma + delta * u) * v;
   return { du_dt, dv_dt };
  }
}
data {
 int<lower = 0> N_subj; // number of subjects
 int < lower = 0 > N;
                           // number of measurement times
  real ts_0[N];
                           // measurement times > 0
 real y0_0[2];  // initial measured populations
  real<lower = 0> y_0[N, 2];  // measured populations
}
transformed data {
 int len[N_subj] = rep_array(N, N_subj);
 real y0[N_subj, 2] = rep_array(y0_0, N_subj);
  real y[N_subj, N, 2] = rep_array(y_0, N_subj);
 real ts[N_subj * N];
 for (i in 1:N_subj) {
    ts[((i-1)*N + 1) : (i*N)] = ts_0;
}
```

```
parameters {
  real<lower = 0> theta[N_subj, 4]; // { alpha, beta, gamma, delta }
  real<lower = 0> z_init[N_subj, 2]; // initial population
  real<lower = 0> sigma[N subj, 2]; // measurement errors
}
transformed parameters {
 matrix[2, N_subj * N] z;
  z = pmx_integrate_ode_group_rk45(dz_dt, z_init, 0, len, ts, theta,

→ rep_array(rep_array(0.0, 0), N_subj), rep_array(rep_array(0,
  \hookrightarrow 0), N_subj));
}
model {
 for (isub in 1:N_subj) {
    theta[isub, \{1, 3\}] ~ normal(1, 0.5);
    theta[isub, \{2, 4\}] ~ normal(0.05, 0.05);
    sigma[isub] ~ lognormal(-1, 1);
    z init[isub] ~ lognormal(10, 1);
    for (k in 1:2) {
      y0[isub, k] ~ lognormal(log(z_init[isub, k]), sigma[isub, k]);
     y[isub, , k] \sim lognormal(log(z[k, ((isub-1)*N + 1):(isub*N)]),
      \hookrightarrow sigma[isub, k]);
    }
  }
```

#### 5.7. Univariate integral of a quadratic function

integral of a quadratic function. This example shows how to use univariate\_integral\_rk45 to calculate the integral of a quadratic function.

```
functions {
  real fun_ord2(real t, real[] theta, real[] x_r, int[] x_i) {
    real a = 2.3;
    real b = 2.0;
   real c = 1.5;
    real res;
    res = a + b * t + c * t * t;
    return res;
  }
}
data {
 real t0;
 real t1;
 real dtheta[2];
  real x_r[0];
 int x_i[0];
transformed data {
  real univar_integral;
  univar_integral = univariate_integral_rk45(func, t0, t1, dtheta,
                          x r, x i);
/* ... */
```

# 5.8. Linear intepolation

This example illustrates how to use linear\_intepolationi to fit a piecewise linear function to a data set consisting of (x, y) pairs.

```
data{
 int nObs;
 real x0bs[n0bs];
 real yObs[nObs];
 int nx;
 int nPred;
 real xPred[nPred];
transformed data{
 real xmin = min(xObs);
 real xmax = max(xObs);
}
parameters{
 real y[nx];
 real<lower = 0> sigma;
  simplex[nx - 1] xSimplex;
}
transformed parameters{
 real yHat[nObs];
 real x[nx];
 x[1] = xmin;
 x[nx] = xmax;
  for (i in 2: (nx-1))
   x[i] = x[i-1] + xSimplex[i-1] * (xmax - xmin);
 yHat = linear_interpolation(xObs, x, y);
 xSimplex ~ dirichlet(rep_vector(1, nx - 1));
  y \sim normal(0, 25);
 yObs ~ normal(yHat, sigma);
generated quantities{
 real yHatPred[nPred];
 real yPred[nPred];
 yHatPred = linear_interpolation(xPred, x, y);
 for(i in 1:nPred)
    yPred[i] = normal_rng(yHatPred[i], sigma);
```

## 5.9. Effect Compartment Population Model

Here we expand the 5.1 to a population model fitted to the combined data from phase I and phase II as tudies. The parameters exhibit inter-individual variations (IIV), due to both random effects and to the patients' body weight, treated as a covariate and denoted bw.

## 5.9.1. Population Model for Plasma Drug Concentration c.

$$\begin{aligned} \log\left(c_{ij}\right) &\sim N\left(\log\left(\widehat{c}_{ij}\right), \sigma^{2}\right), \\ \widehat{c}_{ij} &= f_{2cpt}\left(t_{ij}, D_{j}, \tau_{j}, CL_{j}, Q_{j}, V_{1j}, V_{2j}, k_{aj}\right), \\ \log\left(CL_{j}, Q_{j}, V_{ssj}, k_{aj}\right) &\sim N\left(\log\left(\widehat{CL}\left(\frac{bw_{j}}{70}\right)^{0.75}, \widehat{Q}\left(\frac{bw_{j}}{70}\right)^{0.75}, \widehat{V}_{ss}\left(\frac{bw_{j}}{70}\right), \widehat{k}_{a}\right), \Omega\right), \\ V_{1j} &= f_{V_{1}}V_{ssj}, \\ V_{2j} &= (1 - f_{V_{1}})\,V_{ssj}, \\ \left(\widehat{CL}, \widehat{Q}, \widehat{V}_{ss}, \widehat{k}_{a}, f_{V_{1}}\right) &= \left(10\,\,\mathrm{L/h}, 15\,\,\mathrm{L/h}, 140\,\,\mathrm{L}, 2\,\,\mathrm{h^{-1}}, 0.25\right), \\ \Omega &= \begin{pmatrix} 0.25^{2} & 0 & 0 & 0 \\ 0 & 0.25^{2} & 0 & 0 \\ 0 & 0 & 0.25^{2} & 0 \\ 0 & 0 & 0 & 0.25^{2} \end{pmatrix}, \\ \sigma &= 0.1 \end{aligned}$$

Furthermore we add a fourth compartment in which we measure a PD effect (Figure 5.5).

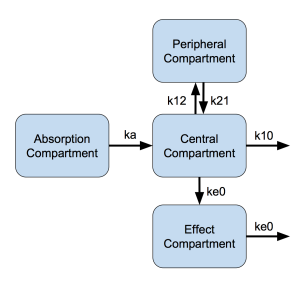


FIGURE 5.5. Effect Compartment Model

## 5.9.2. Effect Compartment Model for PD response R.

$$R_{ij} \sim N\left(\widehat{R}_{ij}, \sigma_R^2\right),$$

$$\widehat{R}_{ij} = \frac{E_{max}c_{eij}}{EC_{50j} + c_{eij}},$$

$$c'_{e \cdot j} = k_{e0j}\left(c_{\cdot j} - c_{e \cdot j}\right),$$

$$\log\left(EC_{50j}, k_{e0j}\right) \sim N\left(\log\left(\widehat{EC}_{50}, \widehat{k}_{e0}\right), \Omega_R\right),$$

$$\left(E_{max}, \widehat{EC}_{50}, \widehat{k}_{e0}\right) = (100, 100.7, 1),$$

$$\Omega_R = \begin{pmatrix} 0.2^2 & 0\\ 0 & 0.25^2 \end{pmatrix}, \quad \sigma_R = 10.$$

The PK and the PD data are simulated using the following treatment.

- Phase I study
  - Single dose and multiple doses
  - Parallel dose escalation design
  - 25 subjects per dose
  - Single doses: 5, 10, 20, and 40 mg
  - PK: plasma concentration of parent drug (c)
  - PD response: Emax function of effect compartment concentration (R)
  - PK and PD measured at 0.125, 0.25, 0.5, 0.75, 1, 2, 3, 4, 6, 8, 12, 18, and 24 hours
- Phase IIa trial in patients
  - 100 subjects
  - Multiple doses: 20 mg
  - sparse PK and PD data (3-6 samples per patient)

The model is simultaneously fitted to the PK and the PD data. For this effect compartment model, we construct a constant rate matrix and use pmx\_solve\_linode. Correct use of Torsten requires the user pass the entire event history (observation and dosing events) for an individual to the function. Thus the Stan model shows the call to pmx\_solve\_linode within a loop over the individual subjects rather than over the individual observations. Note that the correlation matrix  $\rho$  does not explicitly appear in the model, but it is used to construct  $\Omega$ , which parametrizes the PK IIV.

```
data{
  int<lower = 1> nSubjects;
  int<lower = 1> nt;
  int<lower = 1> nObs;
  int<lower = 1> iObs[nObs];
  real<lower = 0> amt[nt];
  int<lower = 1> cmt[nt];
  int<lower = 0> evid[nt];
  int<lower = 0> evid[nt];
  int<lower = 1> start[nSubjects];
  int<lower = 1> end[nSubjects];
  real<lower = 0> time[nt];
  vector<lower = 0>[nObs] cObs;
```

```
vector[nObs] respObs;
  real<lower = 0> weight[nSubjects];
  real<lower = 0> rate[nt];
  real<lower = 0> ii[nt];
 int<lower = 0> addl[nt];
  int<lower = 0> ss[nt];
}
transformed data{
 vector[nObs] logCObs = log(cObs);
  int<lower = 1> nRandom = 5;
 int nCmt = 4;
 real biovar[nCmt] = rep_array(1.0, nCmt);
 real tlag[nCmt] = rep_array(0.0, nCmt);
parameters{
 real<lower = 0> CLHat;
  real<lower = 0> QHat;
 real<lower = 0> V1Hat;
 real<lower = 0> V2Hat;
  // real<lower = 0> kaHat;
  real<lower = (CLHat / V1Hat + QHat / V1Hat + QHat / V2Hat +
                sqrt((CLHat / V1Hat + QHat / V1Hat + QHat / V2Hat)^2 -
                     4 * CLHat / V1Hat * QHat / V2Hat)) / 2> kaHat; // ka
                      \hookrightarrow > lambda 1
 real<lower = 0> ke0Hat;
 real<lower = 0> EC50Hat;
  vector<lower = 0>[nRandom] omega;
 corr_matrix[nRandom] rho;
 real<lower = 0> omegaKe0;
 real<lower = 0> omegaEC50;
  real<lower = 0> sigma;
 real<lower = 0> sigmaResp;
 // reparameterization
 vector[nRandom] logtheta_raw[nSubjects];
 real logKe0_raw[nSubjects];
 real logEC50_raw[nSubjects];
}
transformed parameters{
 vector<lower = 0>[nRandom] thetaHat;
 cov_matrix[nRandom] Omega;
 real<lower = 0> CL[nSubjects];
 real<lower = 0> Q[nSubjects];
 real<lower = 0> V1[nSubjects];
  real<lower = 0> V2[nSubjects];
 real<lower = 0> ka[nSubjects];
 real<lower = 0> ke0[nSubjects];
 real<lower = 0> EC50[nSubjects];
 matrix[nCmt, nCmt] K;
  real k10;
```

```
real k12;
real k21;
row_vector<lower = 0>[nt] cHat;
row_vector<lower = 0>[nObs] cHatObs;
row_vector<lower = 0>[nt] respHat;
row_vector<lower = 0>[nObs] respHatObs;
row_vector<lower = 0>[nt] ceHat;
matrix[nCmt, nt] x;
matrix[nRandom, nRandom] L;
vector[nRandom] logtheta[nSubjects];
real logKe0[nSubjects];
real logEC50[nSubjects];
thetaHat[1] = CLHat;
thetaHat[2] = QHat;
thetaHat[3] = V1Hat;
thetaHat[4] = V2Hat;
thetaHat[5] = kaHat;
Omega = quad_form_diag(rho, omega); // diag_matrix(omega) * rho *

→ diag matrix(omega)

L = cholesky_decompose(Omega);
for(j in 1:nSubjects){
  logtheta[j] = log(thetaHat) + L * logtheta_raw[j];
  logKe0[j] = log(ke0Hat) + logKe0_raw[j] * omegaKe0;
  logEC50[j] = log(EC50Hat) + logEC50_raw[j] * omegaEC50;
  CL[j] = exp(logtheta[j, 1]) * (weight[j] / 70)^0.75;
  Q[j] = \exp(\log t_{j}, 2]) * (weight[j] / 70)^0.75;
  V1[j] = \exp(\log theta[j, 3]) * weight[j] / 70;
  V2[j] = exp(logtheta[j, 4]) * weight[j] / 70;
  ka[j] = exp(logtheta[j, 5]);
  ke0[j] = exp(logKe0[j]);
  EC50[j] = exp(logEC50[j]);
  k10 = CL[j] / V1[j];
  k12 = Q[\dot{1}] / V1[\dot{1}];
  k21 = Q[j] / V2[j];
  K = rep_matrix(0, nCmt, nCmt);
  K[1, 1] = -ka[j];
  K[2, 1] = ka[j];
  K[2, 2] = -(k10 + k12);
  K[2, 3] = k21;
  K[3, 2] = k12;
  K[3, 3] = -k21;
  K[4, 2] = ke0[j];
  K[4, 4] = -ke0[j];
```

```
x[, start[j]:end[j]] = pmx_solve_linode(time[start[j]:end[j]],
        amt[start[j]:end[j]],
                                              rate[start[j]:end[j]],
                                              \rightarrow ii[start[j]:end[j]],
                                              evid[start[j]:end[j]],

    cmt[start[j]:end[j]],

                                              addl[start[j]:end[j]],
                                                ss[start[j]:end[j]], K,
                                                 biovar, tlag);
    cHat[start[j]:end[j]] = 1000 * x[2, start[j]:end[j]] ./ V1[j];
    ceHat[start[j]:end[j]] = 1000 * x[4, start[j]:end[j]] ./ V1[j];
    respHat[start[j]:end[j]] = 100 * ceHat[start[j]:end[j]] ./
       (EC50[j] + ceHat[start[j]:end[j]]);
  }
  cHatObs = cHat[iObs];
  respHatObs = respHat[iObs];
model{
  // Prior
  CLHat \sim \log(10), 0.2);
  QHat \sim \log(15), 0.2);
  V1Hat \sim \log(30), 0.2);
  V2Hat \sim \log(100), 0.2);
  kaHat ~ lognormal(log(5), 0.25);
  keOHat \sim lognormal(log(10), 0.25);
  EC50Hat \sim lognormal(log(1.0), 0.2);
  omega \sim normal(0, 0.2);
```

**5.9.3.** Results. We use the same diagnosis tools as for the previous examples. Table 5.1 summarises the statistics and diagnostics of the parameters. In particular, rhat for all parameters being close to 1.0 indicates convergence of the 4 chains. Figure 5.6 shows the posterior density of the parameters.

Posterior prediction check (PPC) in Figure 5.7 - 5.11 show that the fits to the plasma concentration are in close agreement with the data, notably for the sparse data case (phase IIa study). The fits to the PD data (Figure 5.12 - 5.16) look reasonable considering data being more noisy so the model produces larger credible intervals. Both the summary table and PPC plots show that the estimated values of the parameters are consistent with the values used to simulate the data.

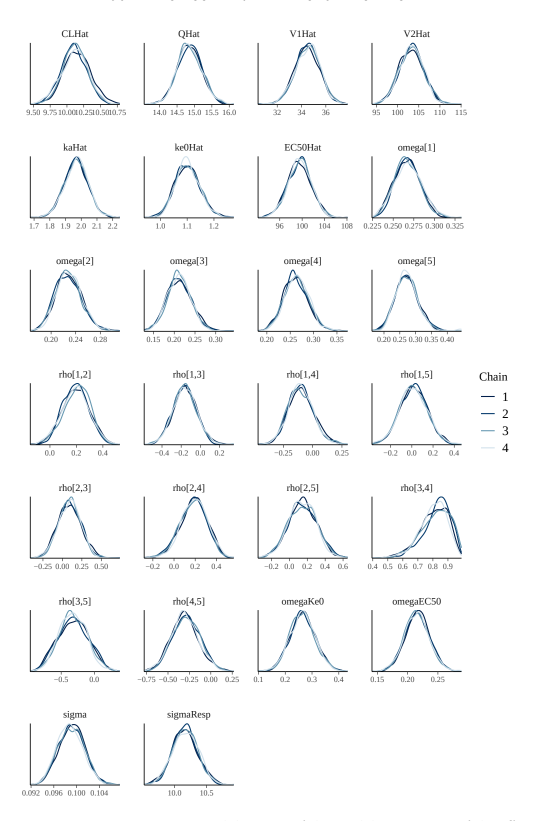


FIGURE 5.6. Posterior marginal densities of the model parameters of the effect compartment model.

TABLE 5.1. Summary of the MCMC simulations of the marginal posterior distributions of the model parameters for the effect compartment model example.

variable	mean	median	$\operatorname{sd}$	$\operatorname{mad}$	q5	q95	rhat	$\operatorname{ess_{bulk}}$	$\operatorname{ess}_{\operatorname{tail}}$
CLHat	10.121	10.120	0.195	0.192	9.797	10.445	1.007	319.942	630.619
$\operatorname{QHat}$	14.858	14.853	0.347	0.344	14.301	15.432	1.000	1106.126	1712.821
V1Hat	34.493	34.516	1.004	0.995	32.814	36.086	1.004	671.777	1563.396
V2Hat	103.269	103.291	2.876	2.878	98.568	108.019	1.002	1689.165	2580.382
kaHat	1.968	1.969	0.076	0.074	1.843	2.087	1.001	1204.531	1747.427
ke0Hat	1.102	1.100	0.046	0.045	1.030	1.180	1.001	4008.337	3167.030
EC50Hat	99.512	99.542	2.124	2.098	95.981	102.987	1.000	2557.436	2773.519
omega[1]	0.268	0.267	0.016	0.016	0.242	0.295	1.008	594.842	978.297
omega[2]	0.229	0.228	0.021	0.021	0.195	0.264	1.002	1245.453	1966.911
omega[3]	0.212	0.211	0.029	0.029	0.165	0.261	1.005	623.820	1692.248
omega[4]	0.263	0.262	0.026	0.026	0.221	0.306	1.002	1396.611	2260.425
omega[5]	0.272	0.271	0.036	0.035	0.217	0.335	1.008	293.132	728.867
rho[1,2]	0.197	0.200	0.100	0.101	0.029	0.360	1.003	1322.261	1955.862
rho[1,3]	-0.161	-0.161	0.122	0.121	-0.361	0.042	1.001	1609.160	2270.515
rho[1,4]	-0.101	-0.105	0.107	0.107	-0.270	0.083	1.001	1685.591	2353.498
rho[1,5]	0.016	0.015	0.128	0.128	-0.192	0.226	1.000	2039.767	2939.988
rho[2,3]	0.091	0.092	0.144	0.148	-0.143	0.328	1.008	718.187	1550.836
rho[2,4]	0.186	0.190	0.125	0.125	-0.025	0.384	1.005	948.704	1819.199
rho[2,5]	0.146	0.145	0.157	0.161	-0.111	0.402	1.003	626.620	1546.157
rho[3,4]	0.815	0.827	0.093	0.094	0.646	0.947	1.010	309.098	736.635
rho[3,5]	-0.318	-0.323	0.219	0.228	-0.678	0.055	1.016	200.806	607.958
rho[4,5]	-0.295	-0.299	0.161	0.162	-0.551	-0.019	1.008	546.998	1151.092
omegaKe0	0.265	0.265	0.047	0.047	0.188	0.346	1.001	1731.276	2049.892
omegaEC50	0.216	0.216	0.020	0.020	0.182	0.249	1.001	1599.567	1844.056
$_{ m sigma}$	0.099	0.099	0.002	0.002	0.095	0.103	1.002	1726.283	2836.027
$\operatorname{sigmaResp}$	10.165	10.166	0.198	0.198	9.844	10.495	1.002	4788.527	2923.203

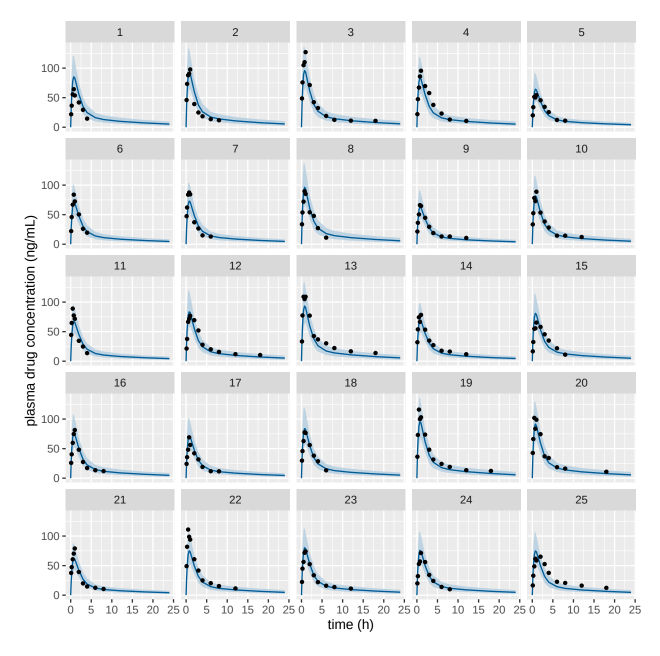


FIGURE 5.7. Predicted (90% credible interval and median) and observed individual plasma drug concentrations in study 1 (5mg dose).

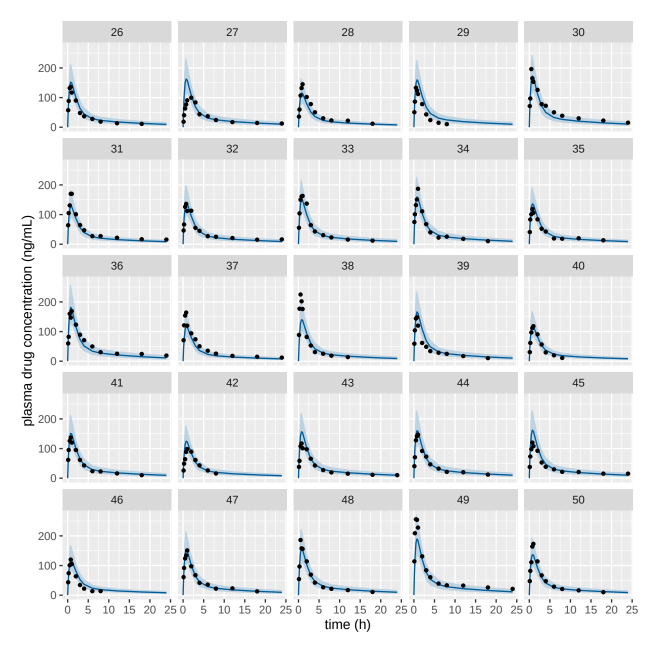


FIGURE 5.8. Predicted (90% credible interval and median) and observed individual plasma drug concentrations in study 1 (10mg dose).

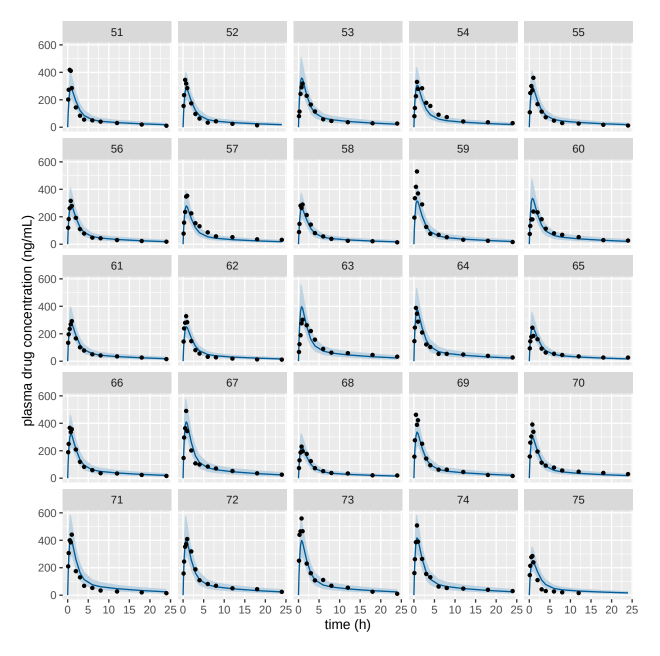


FIGURE 5.9. Predicted (90% credible interval and median) and observed individual plasma drug concentrations in study 1 (20mg dose).

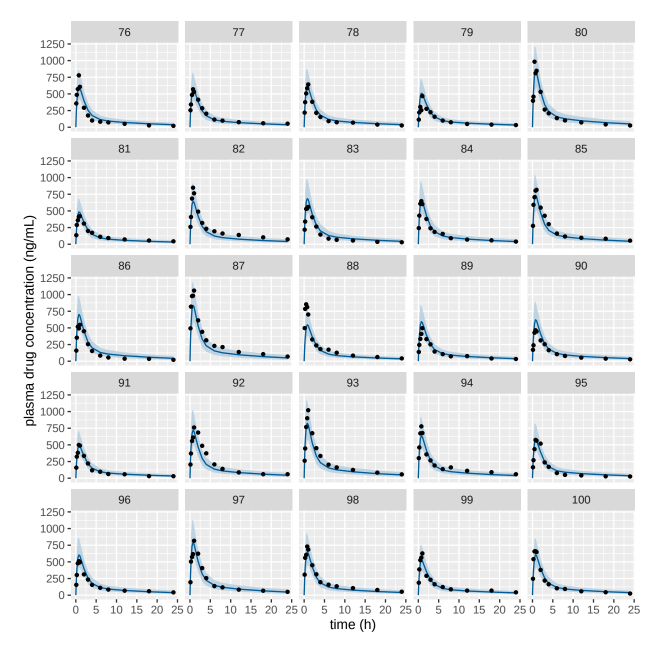


FIGURE 5.10. Predicted (90% credible interval and median) and observed individual plasma drug concentrations in study 1 (40mg dose).

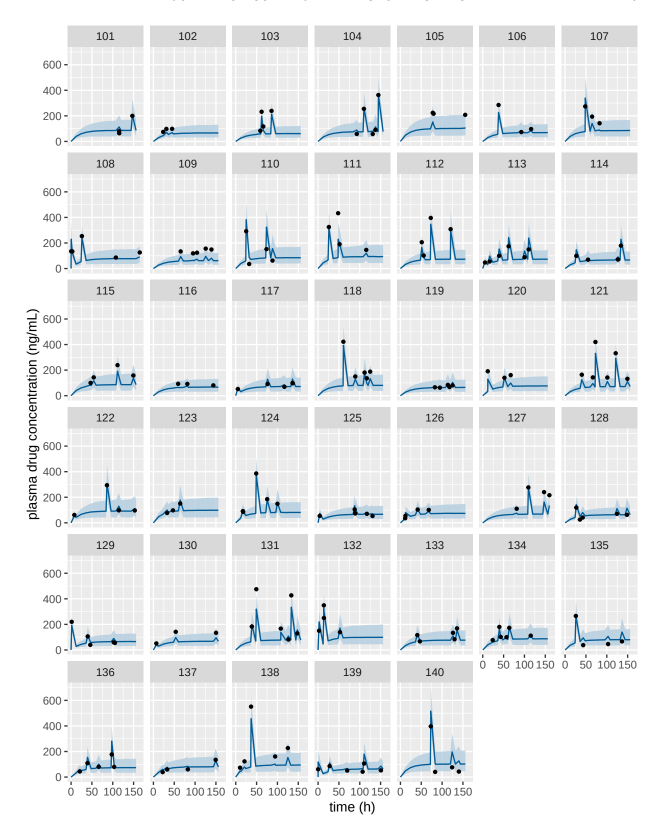


FIGURE 5.11. Predicted (90% credible interval and median) and observed individual plasma drug concentrations in study 2 (first 40 subjects).

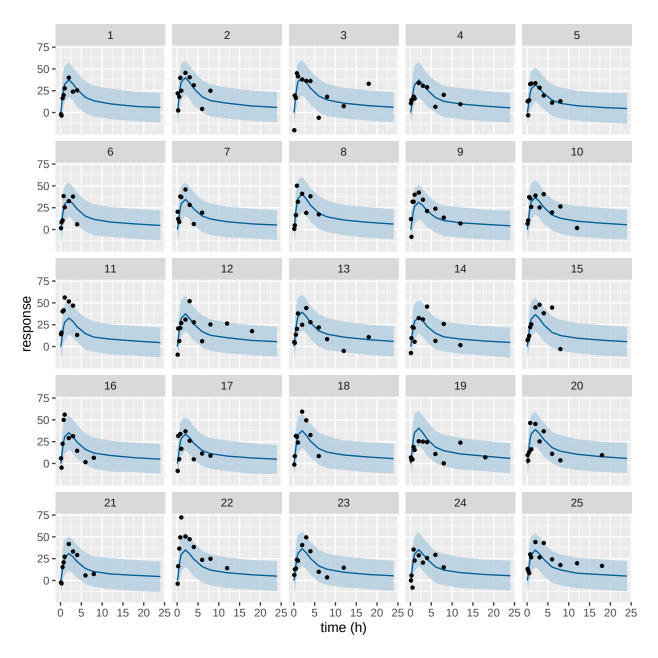


FIGURE 5.12. Predicted (90% credible interval and median) and observed individual PD response in study 1 (5mg dose).

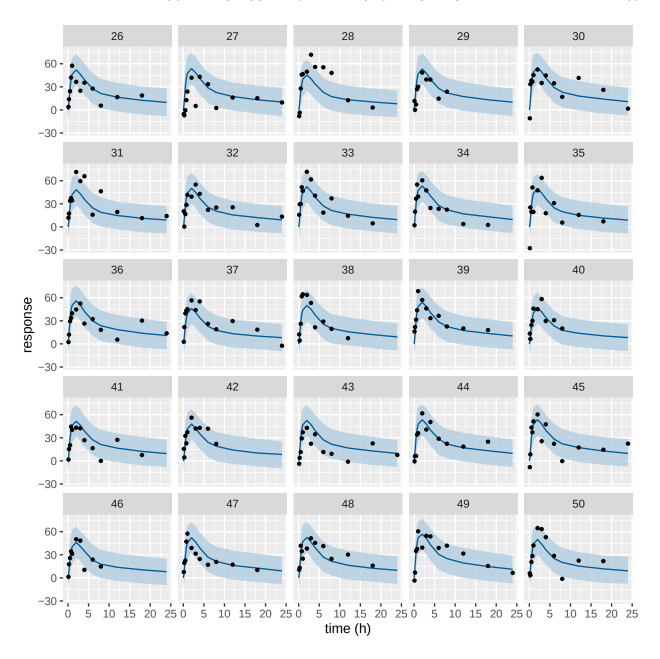


FIGURE 5.13. Predicted (90% credible interval and median) and observed individual PD response in study 1 (10mg dose).

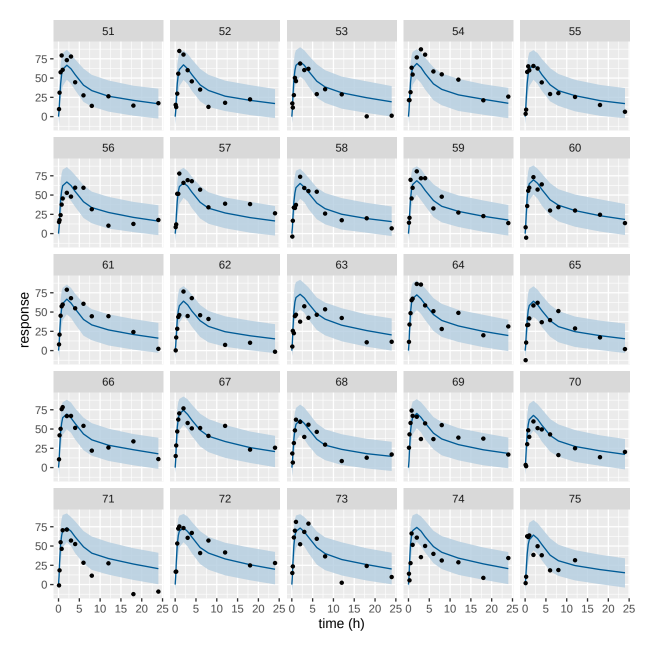


FIGURE 5.14. Predicted (90% credible interval and median) and observed individual PD response in study 1 (20mg dose).

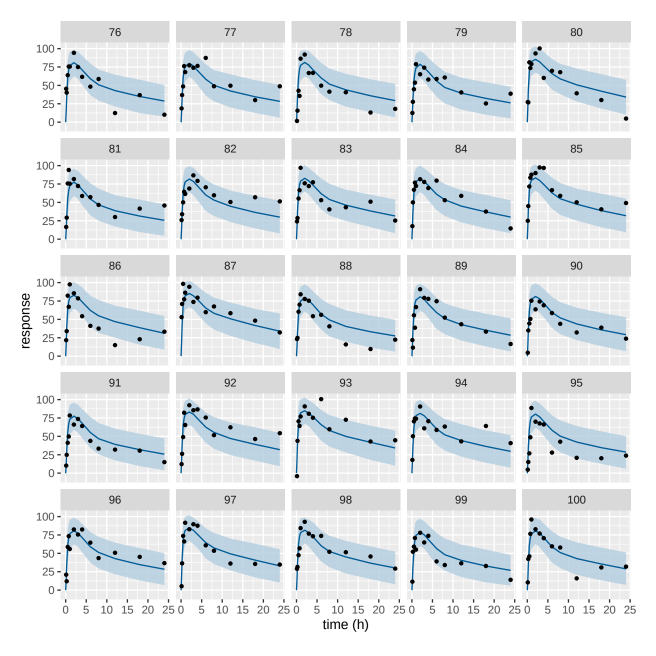


FIGURE 5.15. Predicted (90% credible interval and median) and observed individual PD response in study 1 (40mg dose).

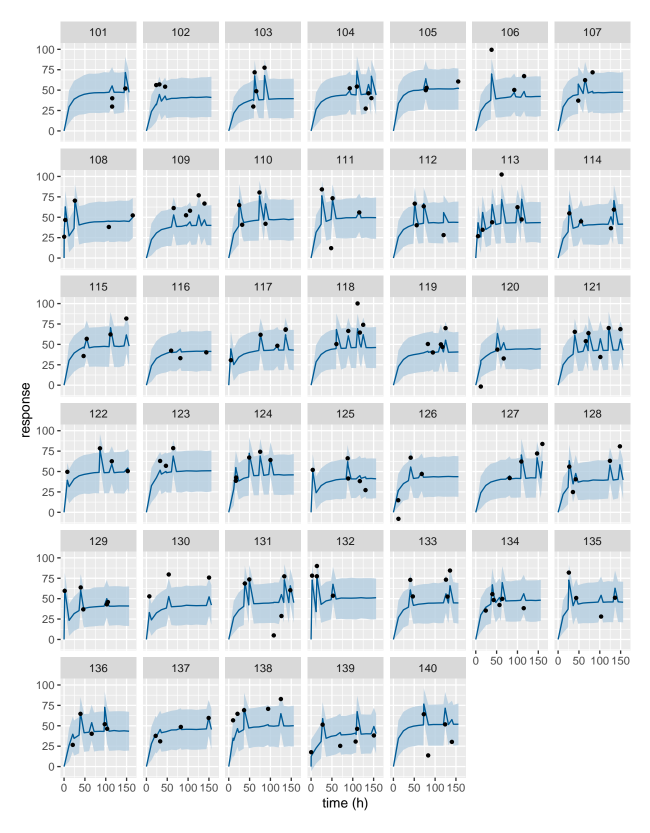


FIGURE 5.16. Predicted (90% credible interval and median) and observed individual PD response in study 2 (first 40 subjects).

# 5.10. Friberg-Karlsson Semi-Mechanistic Population Model

We now return to the example of 5.4 and extend it to a population model. While we recommend using the coupled solver, and this time we solve it using group solver. We leave it as an exercise to the reader to rewrite the model with coupled solver.

# 5.10.1. Friberg-Karlsson Population Model for drug-induced myelosuppression (ANC).

$$\log(ANC_{ij}) \sim N(Circ_{ij}, \sigma_{ANC}^2),$$

$$\log(MTT_j, Circ_{0j}, \alpha_j) \sim N\left(\log\left(\widehat{MTT}, \widehat{Circ_0}, \widehat{\alpha}\right), \Omega_{ANC}\right),$$

$$\left(\widehat{MTT}, \widehat{Circ_0}, \widehat{\alpha}, \gamma\right) = (125, 5, 2, 0.17),$$

$$\Omega_{ANC} = \begin{pmatrix} 0.2^2 & 0 & 0 \\ 0 & 0.35^2 & 0 \\ 0 & 0 & 0.2^2 \end{pmatrix},$$

$$\sigma_{ANC} = 0.1,$$

$$\Omega_{PK} = \begin{pmatrix} 0.25^2 & 0 & a0 & 0 & 0 \\ 0 & 0.4^2 & 0 & 0 & 0 \\ 0 & 0 & 0.25^2 & 0 & 0 \\ 0 & 0 & 0 & 0.4^2 & 0 \\ 0 & 0 & 0 & 0.25^2 \end{pmatrix}$$

The PK and the PD data are simulated using the following treatment.

- Phase IIa trial in patients
  - Multiple doses: 80,000 mg
  - Parallel dose escalation design
  - 15 subjects
  - PK: plasma concentration of parent drug (c)
  - PD response: Neutrophil count (ANC)
  - PK measured at 0.083, 0.167, 0.25, 0.5, 0.75, 1, 2, 3, 4, 6, 8, 12, 18, and 24 hours
  - PD measured once every two days for 28 days.

Once again, we simultaneously fit the model to the PK and the PD data. It pays off to construct informative priors. For instance, we could fit the PK data first, as was done in example 1, and get informative priors on the PK parameters. The PD parameters are drug independent, so we can use information from the neutropenia literature. In this example, we choose to use strongly informative priors on both PK and PD parameters.

The ODE is defined as

```
real CL;
real Q;
real V1;
real V2;
real ka;
real mtt;
real circ0;
real gamma;
real alpha;
real ktr;
vector[8] dxdt;
real conc;
real EDrug;
real transit1;
real transit2;
real transit3;
real circ;
real prol;
CL = parms[1];
Q = parms[2];
V1 = parms[3];
V2 = parms[4];
ka = parms[5];
mtt = parms[6];
circ0 = parms[7];
gamma = parms[8];
alpha = parms[9];
k10 = CL / V1;
k12 = Q / V1;
k21 = Q / V2;
ktr = 4 / mtt;
dxdt[1] = -ka * x[1];
dxdt[2] = ka * x[1] - (k10 + k12) * x[2] + k21 * x[3];
dxdt[3] = k12 * x[2] - k21 * x[3];
conc = x[2]/V1;
EDrug = alpha * conc;
// x[4], x[5], x[6], x[7] and x[8] are differences from circ0.
prol = x[4] + circ0;
transit1 = x[5] + circ0;
transit2 = x[6] + circ0;
transit3 = x[7] + circ0;
circ = fmax(machine\_precision(), x[8] + circ0); // Device for
// initial condition
dxdt[4] = ktr * prol * ((1 - EDrug) * ((circ0 / circ)^gamma) - 1);
dxdt[5] = ktr * (prol - transit1);
dxdt[6] = ktr * (transit1 - transit2);
dxdt[7] = ktr * (transit2 - transit3);
dxdt[8] = ktr * (transit3 - circ);
```

```
return dxdt;
}
}
```

We use the pmx solve group rk45 function to solve the entire population's events.

```
transformed parameters{
 row_vector[nt] cHat;
 vector[nObsPK] cHatObs;
 row_vector[nt] neutHat;
 vector[nObsPD] neutHatObs;
 matrix[8, nt] x;
 real<lower = 0> parms[nSubjects, nTheta]; // The [1] indicates the

→ parameters are constant

 // variables for Matt's trick
 vector<lower = 0>[nIIV] thetaHat;
 matrix<lower = 0>[nSubjects, nIIV] thetaM;
 // Matt's trick to use unit scale
 thetaHat[1] = CLHat;
 thetaHat[2] = QHat;
 thetaHat[3] = V1Hat;
 thetaHat[4] = V2Hat;
 thetaHat[5] = mttHat;
 thetaHat[6] = circOHat;
  thetaHat[7] = alphaHat;
  thetaM = (rep_matrix(thetaHat, nSubjects) .*
             exp(diag_pre_multiply(omega, L * etaStd)))';
  for(i in 1:nSubjects) {
   parms[i, 1] = thetaM[i, 1] * (weight[i] / 70)^0.75; // CL
   parms[i, 2] = thetaM[i, 2] * (weight[i] / 70)^0.75; // Q
   parms[i, 3] = thetaM[i, 3] \star (weight[i] / 70); // V1
   parms[i, 4] = thetaM[i, 4] \star (weight[i] / 70); // V2
   parms[i, 5] = kaHat; // ka
   parms[i, 6] = thetaM[i, 5]; // mtt
   parms[i, 7] = thetaM[i, 6]; // circ0
   parms[i, 8] = gamma;
   parms[i, 9] = thetaM[i, 7]; // alpha
 /* group solver */
 x = pmx_solve_group_rk45(twoCptNeutModelODE, 8, len,
                           time, amt, rate, ii, evid, cmt, addl, ss,
                           parms,
                           1e-6, 1e-6, 500);
 for(i in 1:nSubjects) {
    cHat[start[i]:end[i]] = x[2, start[i]:end[i]] / parms[i, 3]; //
    \hookrightarrow divide by V1
```

This allows us to use within-chain paralleleisation to reduce simulation time. When run from cmdstan, each MPI run generates one chain, and we use 4 MPI runs to generate 4 chains.

```
# chain 1
mpiexec -n nproc ./FribergKarlsson sample adapt delta=0.95 data
\hookrightarrow file=fribergkarlsson.data.R init=fribergkarlsson.init.R random
\hookrightarrow seed=8765 id=1 output file=output.1.csv
# chain 2
mpiexec -n nproc ./FribergKarlsson sample adapt delta=0.95 data
  file=fribergkarlsson.data.R init=fribergkarlsson.init.R random
    seed=8765 id=2 output file=output.2.csv
# chain 3
mpiexec -n nproc ./FribergKarlsson sample adapt delta=0.95 data

→ file=fribergkarlsson.data.R init=fribergkarlsson.init.R random
\rightarrow seed=8765 id=3 output file=output.3.csv
# chain 4
mpiexec -n nproc ./FribergKarlsson sample adapt delta=0.95 data

→ file=fribergkarlsson.data.R init=fribergkarlsson.init.R random
   seed=8765 id=4 output file=output.4.csv
```

**5.10.2.** Results. Table 5.2 summarizes the sampling and some diagnostics output. estimation reflects the real value of the parameters (Table 5.2 and Figure 5.17. Similar to the previous example, PPCs shown in Figure 5.18 and 5.19 indicate the model is a good fit.

Table 5.2. Summary of the MCMC simulations of the marginal posterior distributions of the model parameters for the Friberg-Karlsson population model example.

variable	mean	median	$\operatorname{sd}$	mad	q5	q95	rhat	${\operatorname{ess}}_{\operatorname{bulk}}$	$\operatorname{ess_{tail}}$
CLHat	9.539	9.535	0.522	0.487	8.692	10.401	1.006	971.369	1655.449
QHat	15.401	15.386	1.018	1.000	13.742	17.090	1.000	2263.843	2447.006
V1Hat	37.396	37.360	2.244	2.228	33.762	41.058	1.001	1936.476	2372.815
V2Hat	101.698	101.394	6.503	6.119	91.529	112.538	1.001	2580.227	2592.925
kaHat	1.997	1.997	0.074	0.074	1.873	2.115	1.001	7056.877	2993.406
mttHat	113.681	113.204	11.506	10.910	95.807	133.514	1.001	4255.900	3269.646
circOHat	4.760	4.752	0.241	0.229	4.375	5.163	1.002	3774.920	2783.663
omega[1]	0.223	0.217	0.047	0.042	0.160	0.307	1.000	1751.864	2235.607
omega[2]	0.339	0.329	0.073	0.067	0.239	0.473	1.001	2363.843	2607.056
omega[3]	0.264	0.256	0.057	0.051	0.186	0.367	1.002	2128.660	2018.425
omega[4]	0.257	0.249	0.056	0.051	0.182	0.361	1.003	2293.877	2937.673
omega[5]	0.177	0.169	0.112	0.118	0.019	0.376	1.000	1550.483	2045.025
omega[6]	0.188	0.183	0.044	0.041	0.127	0.269	1.000	2377.698	2965.713
omega[7]	0.409	0.394	0.256	0.259	0.045	0.865	1.003	1386.987	2015.873
gamma	0.171	0.168	0.035	0.033	0.121	0.235	1.000	8809.668	3189.676
$_{ m sigma}$	0.097	0.096	0.003	0.003	0.093	0.101	1.002	5436.508	2899.706
$\operatorname{sigmaNeut}$	0.106	0.105	0.012	0.011	0.088	0.127	1.000	2809.059	3031.605
alphaHat	2.24e-4	2.19e-4	3.97e-05	3.80 e-05	1.66e-4	2.96e-4	1.000	5138.105	2807.328
	CLHat	QHa	t	V1Hat	V2	2Hat	kaHa	at	
					$\wedge$				
					/ `				
					<u> </u>				
8 9	9 10 11	11 13 15	17 19 30	35 40 45	80 100	120 140	1.8 1.9 2.0	2.1 2.2	
	mttHat	circ0H	[nt	alphaHat	ome	ugo[1]	omaga	.[2]	
mttHat				арпана	omega[1]		omega[2]		
				/ \					
				/ \				Cha	in
80 100	0 120 140 160	4.0 4.5 5.0	) 5.5	2e-04 3e-04 4e-	04 02 0	3 0.4 0.5	0.2 0.4 (		
80 100 120 140 160									
C	omega[3]	omega	[4]	omega[5]	ome	ga[6]	omega		3
	λ		Q	7			r A		4
	1								
					/ \				
0.2	0.3 0.4 0.5	0.2 0.3 0.4 0	0.5 0.6	0.2 0.4 0.6 0.8	0.1 0.2	0.3 0.4	0.5 1.0	1.5	
	gamma	sigm	a ,	sigmaNeut					
/		/ \		/ \					
0.100.1	150.200.250.30	0.0900.0950.1	1000.105 0.08	30.100.120.140.1	6				

FIGURE 5.17. Posterior marginal densities of the model parameters of the Friberg-Karlsson population model.

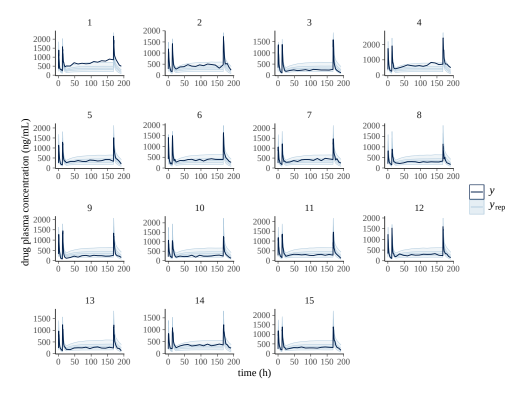


FIGURE 5.18. Predicted (50%, 90% credible interval and median) and observed individual drug plasma concentration.

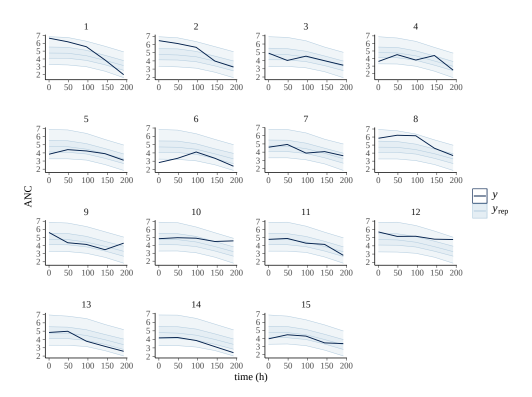


FIGURE 5.19. Predicted (50%, 90% credible interval and median) and observed individual Neutrophil counts.

# Compiling constants

Several constants are used in Torsten's makefile. These constants can be used in cmdstan/make/loca file, or use set\_make\_local command in cmdstanr.

- TORSTEN\_MPI=1 turns on within-chain parallel sation of MPI-enable functions. To use this option one must also point TBB\_CXX\_TYPE to proper C compiler. See also Section 4.7 and 4.9.
- TORSTEN\_CVS\_JAC\_AD=1 makes BDF and Adams integrator use Stan's automatic differentiation to calculate Jacobian matrix in nonlinear solver [6].

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