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Torsten

Torsten: A Pharmacokinetic/Pharmacodynamic Model Library for Stan

User Manual

Torsten Version 0.84 for Stan Version 2.17.1

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Individuals

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1. Introduction

Stan is an open source probabilistic programing language designed primarily to do Bayesian data analysis [2]. Several of its features make it a powerful tool to specify and fit complex models. Notably, its language is extremely flexible and its No U-Turn Sampler (NUTS), an adaptative Hamiltonian Monte Carlo algorithm, has proven more efficient than commonly used Monte Carlo Markov Chains (MCMC) samplers for complex high dimensional problems [3]. Our goal is to harness these innovative features and make Stan a better software for pharmacometrics modeling. Our efforts are twofold:

- (1) We contribute to the development of new mathematical tools, such as functions that support differential equations based models, and implement them directly into Stan's core language.
- (2) We develop Torsten, an extension with specialized pharmacometrics functions.

Throughout the process, we work very closely with the Stan Development Team. We have benefited immensely from their mentorship, advice, and feedback. Just like Stan, Torsten is an open source project that fosters collaborative work. Interested in contributing? Shoot us an e-mail and we will help you help us (billg@metrumrg.com)!

Torsten is licensed under the BSD 3-clause license.

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: https://github.com/metrumresearchgroup/stan.

1.1. Installing Torsten.

Installation files are available on GitHub:

https://github.com/metrumresearchgroup/example-models.

There is currently no mechanism to install Torsten on top of your version of Stan. This is still a work in progress. In the meantime, we offer a version of Stan with Torsten built inside of it. Torsten 0.84 works with Stan 2.17.1. Torsten is built inside the Stan and Stan-math repositories and is agnostic to the interface. We offer support to install Torsten with rstan and CmdStan.

- 1.1.1. Intalling Torsten with rstan. To install RStan with Torsten, install the R package Torsten-Headers (https://github.com/metrumresearchgroup/TorstenHeaders) and run install_torsten():
 - > devtools::install github('metrumresearchgroup/TorstenHeaders')
 - > library(torstenHeaders)
 - > install_torsten()
- 1.1.2. Installing Torsten with CmdStan. You can install the CmdStan interface with Stan and Torsten using the bash file setupTorsten.sh, i.e., running sh setupTorsten.sh from the command line. Then compile CmdStan by running the command make build. If multiple CPU

cores are available you can speed up the installation by adding the -jN option where N is the number of cores to use for installation.

1.2. Overview.

Torsten is a collection of Stan functions to facilitate analysis of pharmacometric data using Stan. The current version includes:

- 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
- Mix compartment model with PK forcing function described by a linear one or two compartment model

The models and data format are based on NONMEM®¹/NMTRAN/PREDPP conventions including:

- Recursive calculation of model predictions
 - This permits piecewise constant covariate values
- Bolus or constant rate inputs into any compartment
- Handles single dose and multiple dose histories
- Handles steady state dosing histories
 - Note: The infusion time must be shorter than the inter-dose interval.
- Implemented NMTRAN data items include: TIME, EVID, CMT, AMT, RATE, ADDL, II, SS

In general, all real variables may be passed as Stan parameters. A few exceptions apply to functions which use a numerical integrator (i.e. the general and the mix compartment models). The below listed cases present technical difficulties, which we expect to overcome in Torsten's next release:

- The RATE and TIME arguments must be fixed
- In the case of a multiple truncated infusion rate dosing regimen:
 - The bioavailability (F) and the amount (AMT) must be fixed.

This library provides Stan language functions that calculate amounts in each compartment, given an event schedule and an ODE system.

1.3. Implementation details.

- Stan version 2.17.1
- All functions are programmed in C++ and are compatible with the Stan math automatic differentiation library [4]
- All functions can be called directly in a Stan file in a manner identical to other built-in functions

¹NONMEM® is licensed and distributed by ICON Development Solutions.

- One and two compartment models: hand-coded analytical solutions
- General linear compartment models with semi-analytical solutions using the built-in matrix exponential function
- General compartment models with numerical solutions 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.
- Mix compartment model: the PK forcing function is solved analytically and the forced ODE system is solved numerically.

1.4. 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)
- Allow numerical methods to handle RATE, AMT, TIME, and the bioavailability fraction (F) as parameters in all cases.
- Optimize Matrix exponential functions
 - Function for the action of Matrix Exponential on a vector
 - Hand-coded gradients
 - Special algorithm for matrices with special properties
- Fix issue that arises when computing the adjoint of the lag time parameter (in a dosing compartment) evaluated at $t_{lag} = 0$.
- Extend formal tests
 - We want more C++ Google unit tests to address cases users may encounter
 - Comparison with simulations from the R package mrgsolve and the software NONMEM®
 - Recruit non-developer users to conduct beta testing

1.5. Updates since Torsten 0.83.

- Torsten is now up to date with Stan version 2.17.1.
- Add piecewise linear interpolation function.
- Add univariate integral functions.
- Minor revisions to User Manual.

2. Using Torsten

The reader should have a basic understanding of how Stan works before reading this chapter. There are excellent resources online to get started with Stan (http://mc-stan.org/documentation/).

In this section we go through the different functions Torsten adds to Stan. It will be helpful to apply these functions to a simple example. We have uploaded code and data on https://github.com/metrumresearchgroup/example-models.

The core model functions of Torsten and their arguments are summarized in Table 1.

2.1. Example 1: Two Compartment Model.

Table 1. Core Torsten model functions and their arguments.

	function	argument	parameters		
model	name	names	in theta		
one compartment	PKModelOneCpt	time, amt, rate, ii,	CL, V_2, k_a		
model with first order		evid, cmt, addl, ss,			
absorption		theta, F, tlag			
two compartment	PKModelTwoCpt	time, amt, rate, ii,	CL, Q, V_2, V_3, k_a		
model with first order		evid, cmt, addl, ss,			
absorption		theta, F, tlag			
general linear	linOdeModel		_		
compartment model		time, amt, rate, ii, CL, V2, ka evid, cmt, addl, ss, theta, F, tlag time, amt, rate, ii, CL, Q, V2, V3, k evid, cmt, addl, ss, theta, F, tlag time, amt, rate, ii, NA: pass in conevid, cmt, addl, ss, rate matrix instructions system, F, tlag **ODE_system, nCmt, Parameters that time, amt, rate, ii, evid, cmt, addl, ss, theta, F, tlag, rel_tol, abs_tol, max_num_steps **reduced_ODE_system, CL, V2, ka, for nOde, time, amt, by the parameter rate, ii, evid, get passed to the cmt, addl, ss, theta, F, tlag, rel_tol, abs_tol, max_num_steps **reduced_ODE_system, CL, Q, V2, V3, k nOde, time, amt, lowed by the parameter, ii, evid, ters that get passed, cmt, addl, ss, the reduced ODE theta, F, tlag, tem rel_tol, abs_tol, ab			
general compartment	genOdeModel_*	- , , , ,	9		
models			passed to ODE system		
		,			
		<u>-</u>			
mix 1 compartment	mixOde1Cpt_*	_			
model					
			~ ·		
			duced ODE system		
		, ,			
mix 2 compartment	mixOde2Cpt_*		CL O Vo Vo k fol-		
model	mixodezcpc_^	-			
model					
		, , , , ,	· ·		
			V-111		
		max_num_steps			

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: plasma concentrations of parent drug (c)
- 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.

The plasma concentration (c) are simulated according to the following equations:

$$\begin{array}{rcl} \log \left(c \right) & \sim & N \left(\log \left(\widehat{c} \right), \sigma^2 \right) \\ \widehat{c} & = & f_{2cpt} \left(t, CL, Q, V_2, V_3, k_a \right) \\ \left(CL, Q, V_2, V_3, ka \right) & = & \left(5 \text{ L/h}, 8 \text{ L/h}, 20 \text{ L}, 70 \text{ L}, 1.2 \text{ h}^{-1} \right) \\ \sigma^2 & = & 0.01 \end{array}$$

and the drug concentration is given by $c = y_2/V_2$.

where the mass of drug in the central compartment (y_2) is obtained by solving the system of ordinary differential equations (ODEs):

$$y_1' = -k_a y_1$$

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

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

The data are generated using the R package *mrgsolve* [5], see TwoCptModelSimulation.R. We use this example to demonstrate use of several functions in the Torsten library.

2.2. Linear One and Two Compartment Model Function.

The one and two compartment model functions have the form:

There is no need to skip a line, but we do so to distinguish between *event* arguments and *model* arguments.

The event arguments describe the event schedule of the clinical trial. time, amt, rate, and ii are arrays of real and evid, cmt, addl, and ss arrays of integers. All arrays have the same length, which corresponds to the number of events.

Next we have the model arguments: theta contains the ODE parameters, F the bioavailability fraction in each compartment, and tlag the lag time in each compartment. The model arguments may be either one or two dimensional arrays. If they are one dimensional arrays, the parameters are constant for all events. If they are two dimensional arrays then each row contains the parameters for the interval [time[i-1], time[i]]. The number of rows should equal the number of events.

The options for *model name* are:

- PKModelOneCpt
- PKModelTwoCpt

which respectively correspond to the one and two compartment model with a first order absorption (Figure 1). An array in theta is expected to contain parameters CL, V_2 , and ka for the one compartment case, and CL, Q, V_2 , V_3 , and ka for the two compartments case, in this order. Setting ka to 0 eliminates the first-order absorption. F contains the bioavailability fraction of each compartment (non-effective if set to 1) and tlag the lag time in each compartment (non-effective if set to 0).

PKModelTwoCpt can be used to fit example 1 as shown in Figure 2.2 and TwoCptModel.stan. We are interested in evaluating the ODE parameters, stored in theta. The bioavailability fraction and the lag times on the other hand are fixed, and we therefore declare F and tlag in the **transformed data** block. Three MCMC chains of 2000 iterations were simulated. The first 1000 iteration of each chain were discarded. Thus 1000 MCMC samples per chain were used for the subsequent analyses.

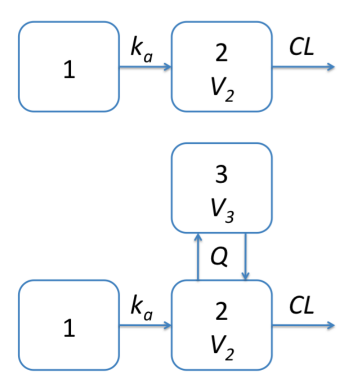


FIGURE 1. One and two compartment models with first order absorption implemented in Torsten.

Result. The MCMC history plots (Figure 3) suggest that the 3 chains have converged to common distributions for all of the key model parameters. The fit to the plasma concentration data (Figure 5) 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 estimates summarized in Table 2 and Figure 4 are consistent with the values used for simulation.

Table 2. Summary of the MCMC simulations of the marginal posterior distributions of the model parameters

-	mean	se_mean	sd	2.5%	25%	50%	75%	97.5%	n_eff	Rhat
$\overline{\text{CL}}$	4.82	0.002	0.0901	4.64	4.76	4.82	4.88	5.00	2464.73	1.00
Q	7.54	0.016	0.58	6.43	7.15	7.54	7.92	8.69	1385.75	1.00
V2	21.14	0.069	2.45	16.37	19.44	21.19	22.78	25.89	1245.64	1.00
V3	76.35	0.110	5.35	65.98	72.75	76.26	79.83	87.30	2379.15	1.00
ka	1.23	0.005	0.169	0.923	1.12	1.23	1.35	1.58	1295.01	1.00
sigma	0.108	0.000	0.012	0.0887	0.0999	0.107	0.115	0.135	1973.97	1.00

```
data {
 int<lower = 1> nt; // number of events
 int<lower = 1> nObs; // number of observation
 int<lower = 1> iObs[nObs]; // index of observation
 int 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 {
 F[1] = 1;
 F[2] = 1;
 F[3] = 1;
 tlag[1] = 0;
 tlag[2] = 0;
 tlag[3] = 0;
}
parameters {
 real<lower = 0> CL;
 real<lower = 0 > Q;
 real<lower = 0> V2;
 real<lower = 0> V3;
 real<lower = 0> ka;
 real<lower = 0> sigma;
transformed parameters {
 theta[1] = CL;
 theta[2] = Q;
 theta[3] = V2;
 theta[4] = V3;
 theta[5] = ka;
 x = PKModelTwoCpt(time, amt, rate, ii, evid, cmt, addl, ss,
                   theta, F, tlag);
 cHat = col(x, 2) ./ V2; // get concentration in the central compartment
 cHatObs = cHat[iObs]; // predictions for observed data records
```

FIGURE 2. Stan language for fitting a two compartment model using the PKModelTwoCpt function (abstract)

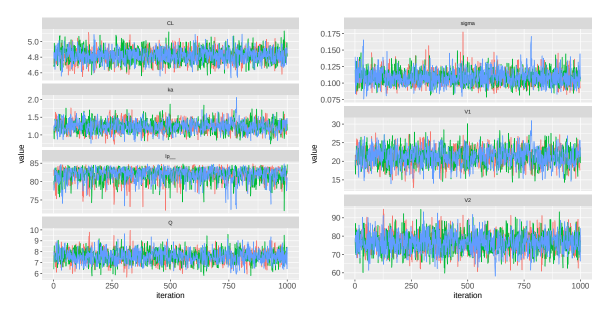


FIGURE 3. MCMC history plots for the parameters of a two compartment model with first order absorption (each color corresponds to a different chain)

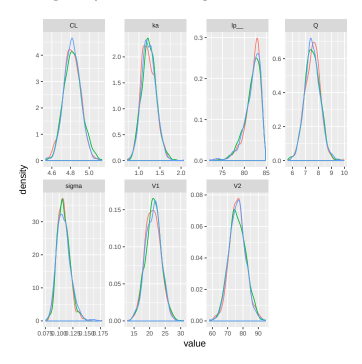


FIGURE 4. Posterior Marginal Densities of the Model Parameters of a two compartment model with first order absorption (each color corresponds to a different chain)

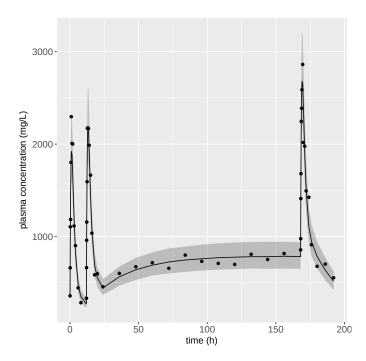


FIGURE 5. Predicted (posterior median and 90 % credible intervals) and observed plasma drug concentrations of a two compartment model with first order absorption

2.3. General Linear ODE Model Function.

A general linear ODE model refers to a model that may be described in terms of a system of first order linear differential equations with (piecewise) constant coefficients, i.e., a differential equation of the form:

$$y'\left(t\right) = Ky\left(t\right)$$

where K is a matrix. For example K for a two compartment model (equation 1) with first order absorption is:

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

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

The linear ODE model function has the form:

system can be:

- the matrix K, if the constant rate matrix is the same for all events.
- an array of constant rate matrices. The length of the array is the number of events and each element corresponds to the matrix at the interval [time[i-1], time[i]].

system contains all the ODE parameters, so we no longer need theta.

Figure 6 and LinTwoCptModel.stan illustrate the use of linOdeModel for fitting a two compartment model with first order absorption.

FIGURE 6. Stan language for fitting a two compartment model using the linOdeModel function (abstract)

2.4. General ODE Model Function.

Torsten may be used to fit models described by a system of user-specified first-order ODEs, i.e., differential equations of the form:

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

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

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

The general ODE model functions have the form:

where ODE_system specifies f(t, y(t)), which the user defines inside the **functions** block (see section 19.2 of the Stan reference manual for details and Figure 7 for an example). The user does NOT include the rates in their definition of f. Torsten automatically corrects the derivatives when the rates are non-zero.

nCmt is the number of compartments (or, equivalently, the number of ODEs) in the model. rel_tol, abs_tol, and max_step are tuning parameters for the ODE integrator: respectively the relative tolerance, the absolute tolerance, and the maximum number of steps.

The options for model_name are:

- generalOdeModel_rk45
- generalOdeModel_bdf

They respectively call the built-in Runge-Kutta 4th/5th order (rk45) integrator, recommended for non-stiff ODEs, and the Backward Differentiation (BDF) integrator, recommended for stiff ODEs. Which value to use for the tuning parameters depends on the integrator and the specifics of the ODE system. Reducing the tolerance parameters and increasing the number of steps make for a more robust integrator but can significantly slow down the algorithm. The following can be used as a starting point: rel_tol = 1e-6, abs_tol = 1e-6 and max_step = 1e+6 for the rk45 integrator and rel_tol = 1e-10, abs_tol = 1e-10 and max_step = 1e+8 for the bdf integrator². Users should be prepared to adjust these values. For additional information, see the Stan User's Manual [6, section 21.6].

A few notable restrictions apply to generalOdeModel_*:

- rate and time cannot be passed as parameters.
- In the case of a multiple truncated infusion rate dosing regimen:
 - The bioavailability (F) and the amount (amt) cannot be passed as parameters.

²These are the default tuning parameters for integrate_ode_rk45() and integrate_ode_bdf(). Torsten functions do not have a default values for these parameters. The user must explicitly pass the tuning parameters to generalOde-Model_*().

FIGURE 7. Stan language for fitting a two compartment model using the generalOdeModel_rk45 function (abstract)

```
functions {
 # define ODE system for two compartment model
 real[] twoCptModelODE(real t,
                       real[] y,
                        real[] theta,
                        real[] dummy_real,
                        int[] dummy_int) {
   real Q = theta[1];
   real CL = theta[2];
   real V2 = theta[3];
   real V3 = theta[4];
   real ka = theta[5];
   real k12 = Q / V2;
   real k21 = Q / V3;
   real k10 = CL / V2;
   real y[3];
   dydt[1] = -ka * y[1];
   dydt[2] = ka * y[1] - (k10 + k12)*y[2] + k21*y[3];
   dydy[3] = k12 * y[2] - k21 * y[3];
   return dydt;
transformed parameters {
 theta[1] = CL;
 theta[2] = Q;
 theta[3] = V1;
 theta[4] = V2;
 theta[5] = ka;
 x = generalCptModel_rk45(twoCptModelODE, 3,
                          time, amt, rate, ii, evid, cmt, addl, ss,
                          theta, F, tlag,
                          1e-8, 1e-8, 1e8);
```

These restrictions also apply to mixOde#Cpt_* functions, discussed in the next section.

2.5. Mixed ODE Model Function.

In certain cases, an ODE system can be divided in two subsystems:

$$y'_1 = f_1(t, y_1)$$

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

where y_1 , y_2 , f_1 , and f_2 are vector-valued functions, and y_1' is independent of y_2 . This structure arises in PK/PD models, where y_1 describes a forcing PK function and y_2 the PD effects. If y_1 has an analytical solution, we can construct a *mixed solver*, which analytically solves y_1 and numerically integrates y_2 . This approach leads to an appreciable gain in computational efficiency. In the example of a Friberg-Karlsson semi-mechanistic model [1], we observe an average speedup of $\sim 47 \pm 18\%$ when using the mix solver in lieu of the numerical integrator [?]. Torsten supports the mixed solver for cases where y_1 solves the ODEs for a One or Two Compartment model with a first-order absorption.

The mix ODE model functions have the form:

where reduced_ODE_system specifies the system we 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 (Figure 8).

FIGURE 8. Stan language for defining a reduced ODE system

Again, the user does not specify the rates. Torsten automatically corrects the derivatives for non-zero rates.

The options for modelName are:

- mixOde1CptModel_rk45
- mixOde1CptModel_bdf

- mixOde2CptModel_rk45
- mixOde2CptModel_bdf

These four functions correspond to all the permutations we can obtain when using a forcing One or Two Compartment function, and the Runge-Kutta 4th/5th order (rk45) or Backward Differentiation (BDF) integration method. The mixed ODE functions can be used to compute the steady state solutions supported by the general ODE model functions.

Restrictions regarding which arguments may be passed as parameters for generalOdeModel_* also apply to mixOde#CptModel_*.

We cannot apply the mixed solver to the Two Compartment example we have been using so far. Instead, we will consider the model which motivated the implementation of the method in the first place.

2.6. Example 2: Friberg-Karlsson Semi-Mechanistic Model [1].

In this second example, we add to our two Compartment model a PD effect, described by a system of nonlinear ODEs.

Neutropenia is observed in patients receiving an ME-2 drug. Our goal is to model the relation between neutrophil counts and drug exposure. Using a feedback mechanism, the body maintains the number of neutrophils at a baseline value (Figure 9). While in the patient's blood, the drug impedes the production of neutrophils. As a result, the neutrophil count goes down. After the drug clears out, the feedback mechanism kicks in and brings the neutrophil count back to baseline.

Friberg-Karlsson Model for drug-induced myelosuppression (ANC)

$$\log(ANC_i) \sim N(\log(Circ), \sigma_{ANC}^2)$$

$$Circ = f_{FK}(MTT, Circ_0, \alpha, \gamma, c)$$

$$(MTT, Circ_0, \alpha, \gamma, ktr) = (125, 5.0, 3 \times 10^{-4}, 0.17)$$

$$\sigma_{ANC}^2 = 0.001$$

where c is the drug concentration in the blood we get from the Two Compartment model, and Circ is obtained by solving the following system of nonlinear ODEs:

$$y'_{\text{prol}} = k_{\text{prol}} y_{\text{prol}} (1 - E_{\text{drug}}) \left(\frac{Circ_0}{y_{\text{circ}}}\right)^{\gamma} - k_{\text{tr}} y_{\text{prol}}$$

$$y'_{\text{trans1}} = k_{\text{tr}} y_{\text{prol}} - k_{\text{tr}} y_{\text{trans1}}$$

$$y'_{\text{trans2}} = k_{\text{tr}} y_{\text{trans1}} - k_{\text{tr}} y_{\text{trans2}}$$

$$y'_{\text{trans3}} = k_{\text{tr}} y_{\text{trans2}} - k_{\text{tr}} y_{\text{trans3}}$$

$$y'_{\text{circ}} = k_{\text{tr}} y_{\text{trans3}} - k_{\text{tr}} y_{\text{circ}}$$

where $E_{druq} = \alpha c$.

The ODEs specifying the Two Compartment Model (equation 1) do not depend on the PD ODEs (equation 2) and can be solved analytically by Torsten. We therefore specify our model using a

mixed solver function. We do not expect our system to be stiff and use the Runge-Kutta 4th/5th order integrator (Figures 10 and 11).

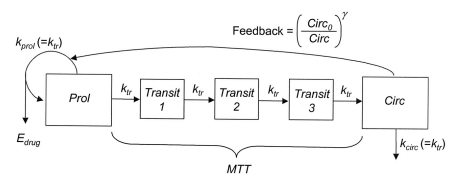


FIGURE 9. Friberg-Karlsson semi-mechanistic Model [1]

FIGURE 10. Stan language to define a reduced ODE system

```
functions {
 \ensuremath{\text{\#}} define reduced ODE system for two compartment model
 real[] FK_ODE(real t,
                real[] y,
               real[] y_pk,
               real[] theta,
               real[] dummy_real,
               int[] dummy_int) {
   real V2 = theta[3];
   ## PK variables
   real VC = parms[3];
   ## PD variables
   real MTT = parms[6];
   real circ0 = parms[7];
   real alpha = parms[8];
   real gamma = parms[9];
   real ktr = 4 / MTT;
   real prol = y[1] + circ0;
   real transit1 = y[2] + circ0;
   real transit2 = y[3] + circ0;
   real transit3 = y[4] + circ0;
   real circ = fmax(machine_precision(), y[5] + circ0);
   real conc = y_pk[2] / VC;
   real Edrug = alpha * conc;
   real dydt[5];
   conc = y_pk[2] / VC;
   Edrug = alpha * conc;
   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;
 }
}
```

FIGURE 11. Stan language for fitting a Friberg-Karlsson model using mixOde2CptModel_rk45

```
functions {
 real[] FK_ODE {
 }
transformed date {
int nOde = 5;
transformed parameters {
vector[nt] cHat;
 vector[nObsPK] cHatObs;
 vector[nt] neutHat;
 vector<lower = 0>[nObsPD] neutHatObs;
 real theta[nParms]; # ODE parameters
 matrix[nt, nCmt] x;
 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 = mixOde2CptModel_rk45(FK_ODE, nOde,
                          time, amt, rate, ii, evid, cmt, addl, ss,
                          theta, F, tlag,
                          1e-6, 1e-6, 1e+6);
 cHat = x[, 2] / VC;
 neutHat = x[, 8] + circ0;
 cHatObs = cHat[iObsPK];
 neutHatObs = neutHat[iObsPD];
```

2.7. Univariate integral.

Based on the ODE solver capability in Stan, Torsten is able to calculate the integral of a univariate function. Following the naming pattern of the ODE solvers, the integral of function f is given by

```
integral = univariate\_integral\_rk45(f, t0, t1, theta, x_r, x_i)
```

using 4th-order Runge-Kutta integrator (for nonstiff problems), or by

```
integral = univariate_integral_bdf(f, t0, t1, theta, x_r, x_i)
```

using BDF integrator (for stiff problems). Here f is a scalar-value integrand, t0 and t1 is the left and right limit of the integral interval, respectively. θ contains parameters. x_r and x_i are real and integer data, respectively.

The integrand function f must follow the following form (Figure 12).

FIGURE 12. Stan language for defining a univariate integrand

Figure 13 shows an example using univariate_integral_rk45 to calculate the integral of a quadratic function.

2.8. Piecewise linear interpolation.

Torsten provides a function 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. The x values must be in increasing order. All 3 arguments may be data or parameters.

The Stan function linear interpolation implements the following function.

$$y_{\text{out}} = \begin{cases} y_1, & x_{\text{out}} < x_1 \\ y_i + \frac{y_{i+1} - y_i}{x_{i+1} - x_i} (x_{\text{out}} - x_i), & x_{\text{out}} \in [x_i, x_{i+1}) \\ y_n, & x_{\text{out}} \ge x_n \end{cases}$$
where
$$x = \{x_1, x_2, \dots, x_n\}$$

$$y = \{y_1, y_2, \dots, y_n\}$$

$$x_{i+1} > x_i \ \forall \ i$$

The following function signatures are currently implemented:

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

FIGURE 13. Stan language for defining a univariate integrand

```
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);
```

Use of linear interpolation is illustrated in a Stan model shown in Figure 14 for fitting a piecewise linear function to a data set consisting of a set of x, y pairs. Complete code for an example using that model is available on GitHub: https://github.com/metrumresearchgroup/example-models. The example is named testInterp2.

FIGURE 14. Stan language model illustrating use of the linear_interpolation function.

```
data{
 int nObs;
 real xObs[nObs];
 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);
model{
 xSimplex ~ dirichlet(rep_vector(1, nx - 1));
 y ~ 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);
```

3. Additional Examples

Code for examples can be found on GitHub: https://github.com/metrumresearchgroup/example-models.

All the files to run a model are stored under the directory that bears the model's name. There are four files per example:

- <model name>.stan
- <model name>.data.R
- <model name>.init.R
- <model name>Simulation.R

data.R contains the data we fit the model to and init.R an initial estimate of the parameters. These two files are generated using Simulation.R. The R folder contains R scripts to compile and run the models, as well as code to output diagnostic plots and statistics.

3.1. Example 3: Effect Compartment Population Model.

Let us expand example 1 to a population model fitted to the combined data from phase I and phase IIa studies. 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:

Population Model for Plasma Drug Concentration (c).

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

Furthermore we add a fourth compartment in which we measure a PD effect (Figure 15).

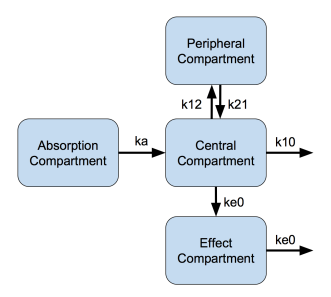


FIGURE 15. Effect Compartment Model

Effect Compartment Model for PD response (R).

$$\begin{array}{rcl} R_{ij} & \sim & N\left(\widehat{R}_{ij}, \sigma_{R}^{2}\right) \\ \widehat{R}_{ij} & = & \frac{E_{max}c_{eij}}{EC_{50j} + c_{eij}} \\ c'_{e \cdot j} & = & k_{e0j}\left(c_{\cdot j} - c_{e \cdot j}\right) \\ \log\left(EC_{50j}, k_{e0j}\right) & \sim & N\left(\log\left(\widehat{EC}_{50}, \widehat{k}_{e0}\right), \Omega_{R}\right) \\ \left(E_{max}, \widehat{EC}_{50}, \widehat{k}_{e0}\right) & = & (100, 100.7, 1) \\ \Omega_{R} & = & \left(\begin{array}{cc} 0.2^{2} & 0 \\ 0 & 0.25^{2} \end{array}\right), \quad \sigma_{R} = 10 \end{array}$$

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: 1.25, 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 0.083, 0.167, 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)

FIGURE 16. Stan language for fitting an effect compartment model using linOdeModel (abstract)

```
transformed parameters {
 for(j in 1:nSubjects){
 Omega = quad_form_diag(rho, omega);
 for(j in 1:nSubjects){
    \begin{array}{lll} \text{CL[j]} &= \exp(\log \text{theta[j, 1]}) & \text{(weight[j] / 70)} \\ ^0.75; \\ \text{Q[j]} &= \exp(\log \text{theta[j, 2]}) & \text{(weight[j] / 70)} \\ \text{V1[j]} &= \exp(\log \text{theta[j, 3]}) & \text{weight[j] / 70;} \\ \end{array} 
   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[j] / V1[j];

k21 = Q[j] / V2[j];
   ke0[j] = exp(logKe0[j]);
   EC50[j] = exp(logEC50[j]);
   K = rep_matrix(0, 4, 4);
   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],] = linOdeModel(time[start[j]:end[j]],
                                            amt[start[j]:end[j]]
                                             rate[start[j]:end[j]],
                                             ii[start[j]:end[j]]
                                             evid[start[j]:end[j]],
                                             cmt[start[j]:end[j]],
                                             addl[start[i]:end[i]],
                                             ss[start[j]:end[j]],
                                             K, F, tlag);
   cHat[start[j]:end[j]] = 1000 * x[start[j]:end[j], 2] ./ V1[j];
   ceHat[start[j]:end[j]] = 1000 * x[start[j]:end[j], 4] ./ 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];
```

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 linOdeModel. 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 linOdeModel within a loop over the individual subjects rather than over the individual observations (Figure 16.

Results. We use the same diagnosis tools as for the previous example. The MCMC history plots (Figure 17) suggest the 4 chains have converged to common distributions. We note some minor auto-correlations for lp_{-} (the log posterior) and for IIV parameters: specifically $\Omega_{ke_{-}0}$ and ρ . The correlation matrix ρ does not explicitly appear in the model, but it is used to construct Ω , which

parametrizes the PK IIV. The fits to the plasma concentration (Figure 19) are in close agreement with the data, notably for the sparse data case (phase IIa study). The fits to the PD data (Figure 20) look good, though the data is more noisy. The model reflects the noise by producing larger credible intervals. The estimated values of the parameters are consistent with the values used to simulate the data (Table 3 and Figure 18).

TABLE 3. Summary of the MCMC simulations of the marginal posterior distributions of the model parameters for the effect compartment model example.

	mean	se_mean	sd	2.5%	25%	50%	75%	97.5%	$n_{-}eff$	Rhat
CLHat	10.523	0.003	0.201	9.712	9.958	10.096	10.231	10.483	4000.000	0.999
QHat	14.867	0.014	0.357	14.182	14.620	14.862	15.106	15.563	678.208	1.007
V1Hat	34.188	0.067	1.089	31.940	33.494	34.214	34.918	36.251	267.748	1.016
V2Hat	103.562	0.076	2.925	98.031	101.600	103.455	105.472	109.583	488.296	1.001
kaHat	1.930	0.004	0.077	1.771	1.880	1.933	1.982	2.076	334.888	1.014
ke0Hat	1.050	0.001	0.044	0.967	1.020	1.051	1.078	1.137	164.741	1.000
EC50Hat	104.337	0.040	2.100	100.169	102.909	104.345	105.768	108.351	744.041	1.000
sigma	0.099	0.000	0.002	0.095	0.097	0.099	0.100	0.103	906.342	1.002
sigmaResp	10.156	0.003	0.197	9.779	10.023	10.154	10.286	10.552	4000.000	1.000
omega[1]	0.270	0.000	0.016	0.241	0.259	0.269	0.280	0.302	4000.000	1.001
omega[2]	0.231	0.001	0.021	0.192	0.217	0.230	0.245	0.275	531.512	1.006
omega[3]	0.219	0.002	0.031	0.158	0.199	0.218	0.238	0.281	158.198	1.017
omega[4]	0.267	0.001	0.026	0.218	0.249	0.266	0.284	0.319	684.870	1.001
omega[5]	0.285	0.002	0.037	0.214	0.259	0.284	0.309	0.361	284.545	1.009
omegaKe0	0.271	0.003	0.047	0.183	0.239	0.271	0.303	0.363	217.350	1.007
omegaEC50	0.213	0.001	0.021	0.174	0.199	0.213	0.227	0.255	190.193	1.000

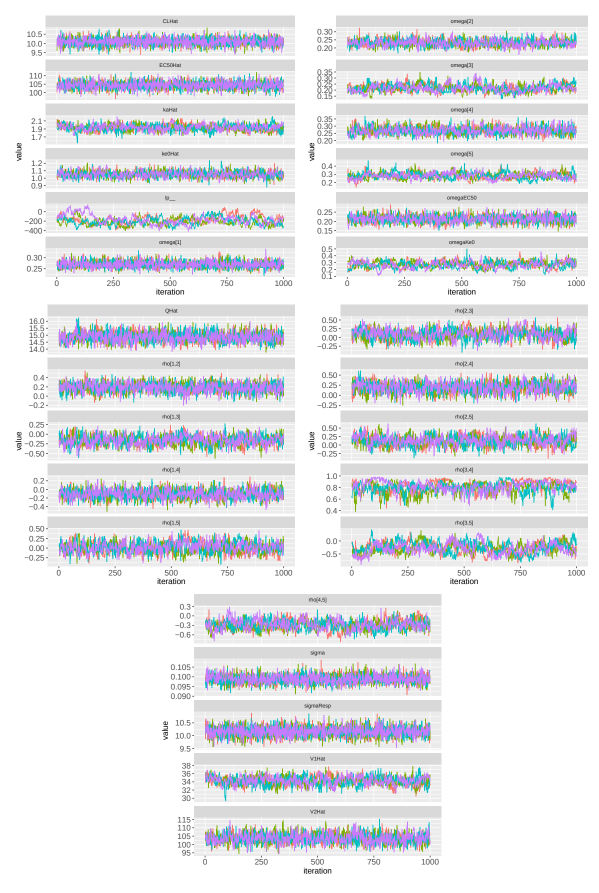


FIGURE 17. MCMC history plots for the parameters of an Effect Compartment Model (each color corresponds to a different chain) for example 2

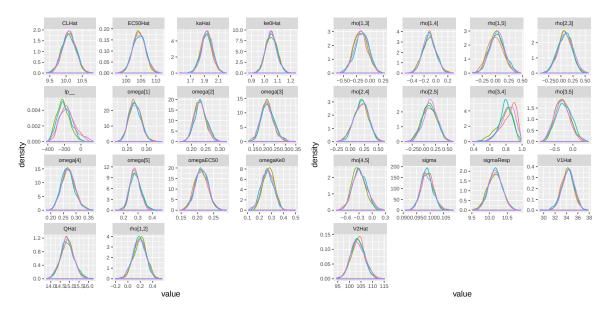


FIGURE 18. Posterior Marginal Densities of the Model Parameters of an Effect Compartment Model (each color corresponds to a different chain) for example 2

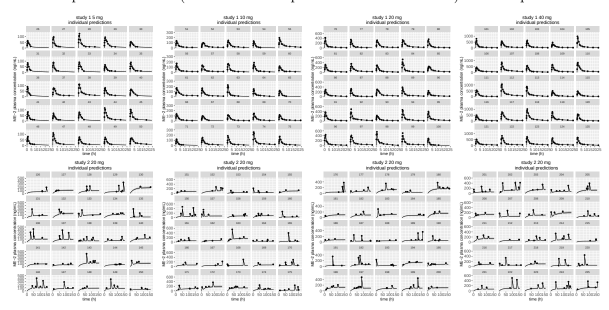


FIGURE 19. Predicted (posterior median and 90 % credible intervals) and observed plasma drug concentrations for example 2 for an Effect Compartment Model



FIGURE 20. Predicted (posterior median and 90 % credible intervals) and observed PD Response for example 2

3.2. Example 4: Friberg-Karlsson Semi-Mechanistic Population Model.

We now return to example 2 and extend it to a population model. While we recommend using the mixed solver, for completeness we'll show how to specify the model with the generalOdeModel_* function. We leave it as an exercise to the reader to rewrite the model with mixOde2CptModel_*.

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

$$\begin{split} \log(ANC_{ij}) &\sim N(Circ_{ij}, \sigma_{ANC}^2) \\ \log(MTT_j, Circ_{0j}, \alpha_j) &\sim N\left(\log\left(\widehat{MTT}, \widehat{Circ_0}, \widehat{\alpha}\right), \Omega_{ANC}\right) \\ \left(\widehat{MTT}, \widehat{Circ_0}, \widehat{\alpha}, \gamma\right) &= (125, 5, 2, 0.17) \\ \Omega_{ANC} &= \begin{pmatrix} 0.2^2 & 0 & 0 \\ 0 & 0.35^2 & 0 \\ 0 & 0 & 0.2^2 \end{pmatrix}, \quad \sigma_{ANC} = 0.1 \\ \Omega_{PK} &= \begin{pmatrix} 0.25^2 & 0 & a0 & 0 & 0 \\ 0 & 0.4^2 & 0 & 0 & 0 \\ 0 & 0 & 0.25^2 & 0 & 0 \\ 0 & 0 & 0 & 0.4^2 & 0 \\ 0 & 0 & 0 & 0.25^2 \end{pmatrix} \end{split}$$

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. Note that from a computational perspective, this is a much more difficult problem than the one we dealt with in the previous example. The nonlinear nature of the ODEs forces us to use a numerical solver, which is significantly slower than the linear methods we have employed so far. Because the ODE system of interest is non-stiff, we use the rk45 version of genOdeModel (Figures 21 and 22).

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 weakly informative priors on the PK parameters and strongly informative priors on the PD parameters.

Since it takes a long time to run the model, we only use 100 iterations per chain, and study what we can learn from this less than optimal scenario. It is worth noting that Stan, because of its highly efficient MCMC sampler, still does a reasonable job estimating the posterior distribution.

FIGURE 21. Stan language for coding an ODE system describing a Friberg-Karlsson Mechanism

```
real[] twoCptNeutModelODE(real t,
              real[] x,
              real[] parms,
              real[] rdummy,
              int[] idummy) {
real CL = parms[1];
real Q = parms[2];
real V2 = parms[3];
real V3 = parms[4];
real ka = parms[5];
real mtt = parms[6];
real circ0 = parms[7];
real gamma = parms[8];
real alpha = parns[9];
real k10 = CL / V2;
real k12 = Q / V2;
real k21 = Q / V3;
real ktr = 4 / mtt;
real dxdt[8];
real conc:
real EDrug:
real transit1:
real transit2:
real transit3;
real circ;
real prol;
dxdt[1] = -ka * x[1];
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 implementing a modeled
                                                      // 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;
```

Results. The MCMC history plots are not as convincing as in the previous examples, mostly because the number of iterations is small (100 versus 1000 in the previous example) (Figure 23. It does however look as though the chains are converging to a common distribution, and we see little auto-correlation (in particular, we expect that if we had run the model for 1000 iterations, we would obtain the desired "fuzzy caterpillar" look). The model fits the data, and the credible interval reflect the noise in the data (Figure 25). The parameters estimation reflects the real value of the parameters (Table 4 and Figure 24).

FIGURE 22. Stan language for fitting a Friberg-Karlsson model using genCptModel_rk45 (abstract)

```
transformed parameters {
 for(i in 1:nSubjects) {
   parms[1] = thetaM[i, 1] * (weight[i] / 70)^0.75; # CL
parms[2] = thetaM[i, 2] * (weight[i] / 70)^0.75; # Q
parms[3] = thetaM[i, 3] * (weight[i] / 70); # V1
parms[4] = thetaM[i, 4] * (weight[i] / 70); # V2
    parms[5] = kaHat; # ka
   parms[6] = thetaM[i, 5]; # mtt
parms[7] = thetaM[i, 6]; # circ0
   parms[8] = gamma;
parms[9] = thetaM[i, 7]; # alpha
    x[start[i]:end[i]] = generalOdeModel_rk45(twoCptNeutModelODE, 8,
                                                          time[start[i]:end[i]],
                                                          amt[start[i]:end[i]],
                                                          rate[start[i]:end[i]],
                                                          ii[start[i]:end[i]],
                                                          evid[start[i]:end[i]],
                                                          cmt[start[i]:end[i]],
                                                          addl[start[i]:end[i]],
                                                          ss[start[i]:end[i]],
                                                          parms, F, tlag,
                                                          1e-6, 1e-6, 1e6);
    \texttt{cHat[start[i]:end[i]] = x[start[i]:end[i], 2] / parms[1][3]; \# divide \ by \ V1}
    neutHat[start[i]:end[i]] = x[start[i]:end[i], 8] + parms[1][7]; # Add baseline
  cHatObs = cHat[iObsPK];
  neutHatObs = neutHat[iObsPD];
```

TABLE 4. Summary of the MCMC simulations of the marginal posterior distributions of the model parameters for the Friberg-Karlsson model example.

	mean	se_mean	$_{\mathrm{sd}}$	2.5%	25%	50%	75%	97.5%	n_eff	Rhat
CL	9.986	0.009	0.174	9.641	9.872	9.982	10.107	10.331	400.000	0.997
Q	14.633	0.055	1.106	12.505	13.992	14.623	15.296	16.948	400.000	0.996
V1	32.909	0.174	2.439	28.203	31.186	32.836	34.762	37.750	195.828	1.008
V2	106.631	0.311	6.226	95.234	102.269	106.403	111.000	118.533	400.000	0.999
ka	1.882	0.012	0.175	1.582	1.756	1.871	2.006	2.223	196.052	1.007
sigma	0.106	0.001	0.010	0.089	0.098	0.105	0.112	0.132	259.693	1.009
alpha	3.3E-04	1.4E-06	2.2E-05	2.9E-04	3.2E-04	3.3E-04	3.5E-04	3.8E-04	247	1.01
mtt	132.763	0.515	6.498	120.843	128.082	132.223	136.694	146.845	159.372	1.024
circ0	5.014	0.009	0.172	4.711	4.888	5.000	5.138	5.334	400.000	1.000
gamma	0.190	0.002	0.022	0.153	0.175	0.187	0.202	0.239	139.485	1.025
sigmaNeut	0.092	0.001	0.014	0.068	0.082	0.090	0.100	0.125	161.199	1.010

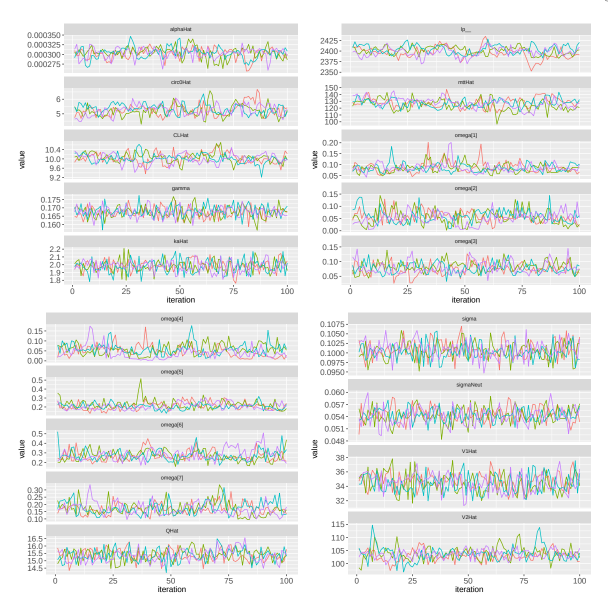


FIGURE 23. MCMC history plots for the parameters of a Friberg-Karlsson semimechanistic model (each color corresponds to a different chain) for example 3

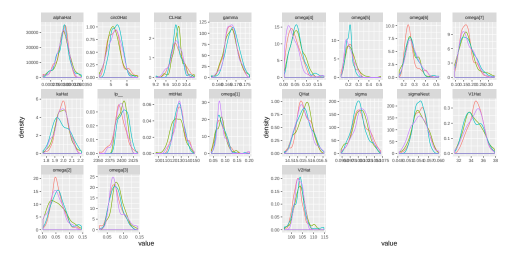


FIGURE 24. Posterior Marginal Densities of the Model Parameters of a Friberg-Karlsson semi-mechanistic model (each color corresponds to a different chain)

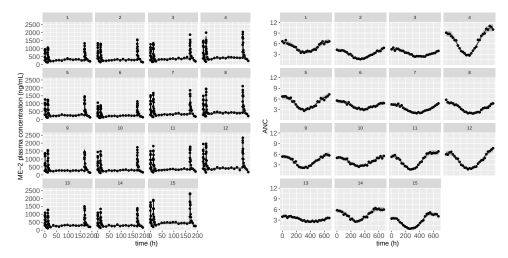


FIGURE 25. Predicted (posterior median and 90 % credible intervals) and observed plasma drug concentrations, and Neutrophil counts, for a Friberg-Karlsson semi-mechanistic model

4. Appendix

(Note: this section is still being worked on and is far from finished)

4.1. Implementing Torsten.

Stan's math library is written in C++, which offers a great deal of speed and flexibility. The Stan language provides a very handy interface that allows us to focus on statistical modeling and saves us the trouble of doing extensive coding in C++. At run time, a *make* file translates our Stan model into C++, which then gets compiled and executed. Accordingly, there are two steps to add a function to Stan: (1) write the procedure in C++, (2) expose the procedure to the language so users may use it in a Stan file.

The Stan code is open-source and available on GitHub. It is compartmentalized into several reposemath contains the mathematical functions, Stan exposes these functions. Other repose provide code to interface Stan with higher level languages, such as R and Python. Torsten exists as a forked version of math and Stan. Other reposeremain unchanged.

Regularly, we merge Stan's latest release into Torsten.

Modifications in math. All Torsten files are located in the Torsten directory, under stan/math. The code can be found on GitHub: https://github.com/metrumresearchgroup/math

Modifications in Stan. We do further modifications in Stan to expose Torsten's functions. We edit function_signatures.h to expose PKModelOneCpt, PKModelTwoCpt, and linOdeModel. The general ODE model functions are higher-order functions (i.e. they take another function as one of their arguments). They are exposed by directly modifying the grammar files, following closely the example of integrate_ode_rk45 and integrate_ode_bdf.

The code can be found on GitHub: https://github.com/metrumresearchgroup/stan.

References

- [1] Friberg, L.E. and Karlsson, M.O. Mechanistic models for myelosuppression. *Invest New Drugs* 21 (2003):183–194.
- [2] Carpenter, B., Gelman, A., Hoffman, M., Lee, D., Goodrich, B., Betancourt, M., Brubaker, M.A., Guo, J., Li, P., Riddell, A. et al. Stan: A probabilistic programming language. *Journal of Statistical Software* 20 (2016):1–37.
- [3] Hoffman, M.D. and Gelman, A. The no-u-turn sampler: Adaptively setting path lengths in hamiltonian monte carlo. *Journal of Machine Learning Research* (2014):1593–1623.
- [4] Carpenter, B., Hoffman, M.D., Brubaker, M.A., Lee, D., Li, P. and Betancourt, M.J. The stan math library: Reverse-mode automatic differentiation in c++. arXiv 1509.07164. (2015).
- [5] Baron, K.T., Hindmarsh, A.C., Petzold, L.R., Gillespie, W.R., Margossian, C.C. and Pastoor, D. mrgsolve: Simulate from ODE-Based Population PK/PD and Systems Pharmacology Models. Metrum Research Group (2017). R package version 0.8.9.
 - URL https://cran.r-project.org/package=mrgsolve
- [6] Stan Development Team. Stan Modeling Language Users Guide and Reference Manual, Version 2.17.0 (2017). http://mc-stan.org/.