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# ACCELERATED PREDICTIVE HEALTHCARE ANALYTICS WITH PUMAS, A HIGH PERFORMANCE PHARMACEUTICAL MODELING AND SIMULATION PLATFORM

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## ABSTRACT

Pharmacometric modeling establishes causal quantitative relationship between administered dose, tissue exposures, desired and undesired effects and patient's risk factors. These models are employed to de-risk drug development and guide precision medicine decisions. Recent technological advances rendered collecting real-time and detailed data easy. However, the pharmacometric tools have not been designed to handle heterogeneous, big data and complex models. The estimation methods are outdated to solve modern healthcare challenges. We set out to design a platform that facilitates domain specific modeling and its integration with modern analytics to foster innovation and readiness to data deluge in healthcare.

New specialized estimation methodologies have been developed that allow dramatic performance advances in areas that have not seen major improvements in decades. New ODE solver algorithms, such as coefficient-optimized higher order integrators and new automatic stiffness detecting algorithms which are robust to frequent discontinuities, give rise to up to 4x performance improvements across a wide range of stiff and non-stiff systems seen in pharmacometric applications. These methods combine with JIT compiler techniques and further specialize the solution process on the individual systems, allowing statically-sized optimizations and discrete sensitivity analysis via forward-mode automatic differentiation, to further enhance the accuracy and performance of the solving and parameter estimation process. We demonstrate that when all of these techniques are combined with a validated clinical trial dosing mechanism and non-compartmental analysis (NCA) suite, real applications like NLME parameter estimation see run times halved while retaining the same accuracy. Meanwhile in areas with less prior optimization of software, like optimal experi-

mental design, we see orders of magnitude performance enhancements. Together we show a fast and modern domain specific modeling framework which lays a platform for innovation via upcoming integrations with modern analytics.

**Keywords** pharmacometrics, nonlinear mixed effects models, pharmacokinetics, pharmacodynamics, differential equations, high-performance computing, parallelism, optimal design, noncompartmental analysis, stiffness, just-in-time compilation, Julia

Clinical pharmacologists, pharmacometricians, systems biologists and statisticians have been leveraging the latest advances in scientific computing to solve complex healthcare problems. The scientific analyses continue to evolve to be more complex as more data became available historically. However, the latest advances in computational science are not readily available to healthcare scientists.

Scientific computing and machine learning have continued to make strides in recent decade, improving core methodologies from multigrid preconditioners and neural network architectures all the way down to the basics of matrix multiplication. Surprisingly, one field which has remained consistent in both the methodologies and software used in practice is small systems of ordinary differential equations. The core software, such as the Hairer's widely used Runge-Kutta methods (DOPRI5, DOP853) [13] and the backwards differentiation formulae (BDF) methods derived from LSODE [16] (LSODA [15], VODE [8], and CVODE [17]), were fully developed by the 90's and have been the stable core of scientific computing ever since. However, in this manuscript we will demonstrate that this general area of computational science can see dramatic computational performance improvements by developing a new set of solver algorithms specialized for simulation of small-scale stiff and non-stiff dynamical systems.

The delay in the availability of the latest advances to healthcare scientists limits their ability to gain deep insights into why some patients do not respond to treatment, why some develop serious toxicity, risk factors for deciding on the right treatment for the right patient (precision medicine). The access to heterogenous data from laboratory measurements, radiographic scans, clinical scripts, genomic and genetic data has not fully translated into actionable science due to the lack of more efficient tools, to a large extent [12]. A related challenge has also been the lack of a unified platform to perform these advanced scientific analyses. For example, a systems biologist cannot perform nonlinear mixed effects modeling for the same project within one software. Unknowingly, this caused serious communication issues between scientists at different stages of the same project. On other hand, one scientist cannot be expected to master all software tools. An integrated platform that allows scientists to build tools in a seamless manner is urgently needed.

Fields which require high fidelity and stable estimation of parameters of such dynamical systems, such as pharmacometrics and systems biology, are frequently constrained by the calculation times required when solving large numbers of such systems [2, 7]. In this manuscript we will demonstrate how these new differential equation solvers are integrated with automatic differentiation and parameter fitting routines in a manner that decreases the time of real-world applications by an order of magnitude.

Likewise, without a bridge that connects these new solver and compiler tools to domain-specific tooling, the ability for pharmacometricians to make use of these tools is limited and requires a modeler to go to tools usually written in low level languages like C++ or Fortran. This limits the flexibility of the allowed models and decreases the speed at which the latest advancements in high performance computing become available to the practitioner. Pumas (**PharmaceUtical Modeling And Simulation**) is impacting this flow by directly packaging the latest mathematical and hardware advances inside of a pharmacometric modeling context inside of the Julia high level high performance language [6]. The driving paradigm of Pumas is to have a completely flexible core while successively simplifying interfaces through generally useful defaults. This allows for a graded approach to learning the modeling framework where beginners can simply use the defaults and expect it to match standard behavior, while keeping non-standard pharmacometric models (such as stochastic differential equations) directly accessible, optimized, and able to utilize all of the hardware compatibility tooling in a first-class manner. This is made possible because Pumas is written for the Julia programming language while being written entirely with the Julia programming language; allowing user-written extensions to flow directly from standard usage. While Julia is a high level programming language which allows Pumas to have ease of use for non-programmers, the Julia language is a just-in-time (JIT) compiled language as fast as low level languages like C or Fortran. Thus both the library and any user-written components are free from interpreter overhead imposed in languages like Python or R. Therefore, our approach with Pumas is to not shy away from using the language and its extensive package ecosystem, and instead integrate our approaches with these tools.

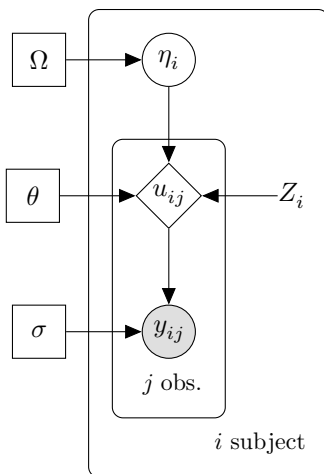


Figure 1: **Diagrammatic description of nonlinear mixed effect models.** A plate diagram of the model: rectangle nodes denote parameters, circles denote random quantities which are either latent (unfilled) or observed (filled), diamonds are deterministic given the inputs, and nodes without a border are constant.

In this paper we will describe the generalized nonlinear mixed effects models (NLME) [7] framework which Pumas utilizes for personalized precision dosing [31]. We will then showcase how the deep integration with the DifferentialEquations.jl [33] software package can allow for many domain-optimized approaches to be accessible within the context of pharmacometric models such as those seen in pharmacokinetics and pharmacodynamics (PK/PD) [3, 43]. We will demonstrate how this connection facilitates Integrated Pharmacometrics and Systems Pharmacology (iPSP) [41] by allowing the optimized solution of large sparse PBPK and QSP models within the NLME context, and showcase how alternative differential equation forms like Differential-Algebraic Equations (DAEs) can be used to stabilize a model or Stochastic Differential Equations (SDEs) can be used to generalize a model to include process noise. Features of Pumas, like integrated high-performance noncompartmental analysis (NCA), automatically parallelized visual predictive checks (VPCs), and fast tooling for optimal design of clinical experiments is all integrated into the Pumas system to allow full applications to be simple and fast. After seeing the modeling benefits of such a framework, we detail the performance benefits, showcasing acceleration over previous software in the standard ODE NLME cases while demonstrating automated parallelism. Together Pumas is a tool built for the next-generation of pharmacometric analysis that will allow for modeling and developing personalized precision medicine in areas that were previously inaccessible due to excessive computational cost.

## 1 Introduction to Nonlinear Mixed Effects Modeling with Pumas

### 1.1 Nonlinear Mixed Effects Models

Many pharmacometric studies fall into a class of models known as nonlinear mixed effects models [7]. Figure 1 gives a diagrammatic overview of this two-stage hierarchical model. In the context of pharmacometrics, the lower level model describes the drug dynamics within a subject via a differential equation while the higher level model describes how the dynamical model is different between and within subject. The fixed effects are the values  $\theta$  which are independent of the subject. One can think of the fixed effects as population typical values. With every subject  $i$  there is a set of covariate values  $Z_i$  which we can know about a subject in advance, such as their weight, height, or sex. We then allow a parameter  $\eta_i$  which is known as the random effect to be the difference between the typical value and the subject. The structural model  $g$  is the function that collates these values into the dynamical parameters for a subject  $p_i$ , i.e.:

$$p_i = g(\theta, Z_i, \eta_i) \quad (1)$$

The dynamical parameters are values such as reaction rates, drug clearance, and plasma volume which describe how the drug and patient reaction evolves over time through an ordinary differential equation (ODE):

$$u'_i = f(u_i, p_i, t) \quad (2)$$

where  $f$  is the dynamical model and  $u$  is the state variables that are being evolved, such as the drug concentrations over time. The  $j$ th observables of patient  $i$ ,  $y_{ij}$ , such as the maximum concentration or area under the curve (AUC), are derived values from the dynamical simulation through a function  $h$ :

$$y_{ij} = h_j(u_i, p_i, Z_i, \eta_i, \theta) \quad (3)$$

Lastly, measurements are taken on the derived values by assuming measurement noise of some distribution (commonly normal) around the prediction point.

The following showcases a classic model of Theophylline dynamics via a 1-compartment model implemented in Pumas, where patients have covariates:

$$Z_i = \begin{bmatrix} wt_i, \\ sex_i, \end{bmatrix} \quad (4)$$

a structural collocation:

$$g_i = \begin{bmatrix} Ka \\ CL \\ V \end{bmatrix} = \begin{bmatrix} \theta_1 e^{\eta_{i,1} \kappa_{i,k,1}}, \\ \theta_2 \left(\frac{wt_i}{70}\right)^{0.75} \theta_4^{sex_i} e^{\eta_{i,2}}, \\ \theta_3 e^{\eta_{i,3}}, \end{bmatrix} \quad (5)$$

internal dynamics:

$$\frac{d[\text{Depot}]}{dt} = -Ka[\text{Depot}], \quad (6)$$

$$\frac{d[\text{Central}]}{dt} = Ka[\text{Depot}] - \frac{CL}{V}[\text{Central}]. \quad (7)$$

and normally distributed measurement noise. The reason for the NLME model is that, if we have learned the population typical values,  $\theta$ , then when a new patient comes to the clinic we can guess how they are different from the typical value by knowing their covariates  $Z_i$  (with the random effect  $\eta_i = 0$ ). We can simulate between-subject variability not captured by our model by sampling  $\eta_i$  from some representative distributions of the  $\eta_i$  from our dataset (usually denoted  $\eta_i \sim N(0, \Omega)$ ). Therefore this gives a methodology for understanding and predicting drug response from easily measurable information.

```

model = @model begin
  @param begin
    θ ∈ VectorDomain(4)
    Ω ∈ PSDDomain(3)
    σ ∈ RealDomain()
  end

  @random begin
    η ~ MvNormal(Ω)
  end

  @covariates wt sex

  @pre begin
    CL = θ[1] * (wt/70)^0.75 * exp(η[1])
    V = θ[2] * (wt/70)^0.75 * θ[4]^sex * exp(η[2])
    Ka = θ[3] * exp(η[3])
  end

  @dynamics begin
    Depot' = -Ka*Depot
    Central' = Ka*Depot - Central*CL/V
  end

  @derived begin

```

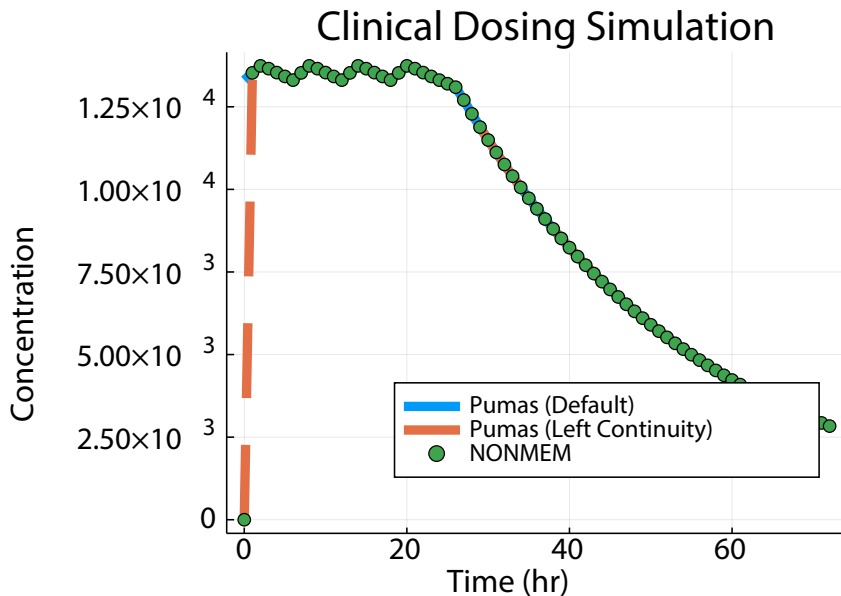


Figure 2: **Clinical dosing simulation comparison against NONMEM.** Shown is the continuous simulation trajectory outputted from Pumas using the default (right) and left continuity choices, compared against NONMEM. This figure both showcases a relative difference  $< 1 \times 10^{-10}$  except at the starting point, where the NONMEM simulation requires a pre-dose estimate of zero whereas the Pumas simulations allows a continuous output via a post-dose observation and allows for switching to the NONMEM behavior.

```
conc := Central/V
dv ~ @. Normal(Central/V, sqrt(conc^2*σ))
end
end
```

## 1.2 Clinical Dosage Regimen Modeling

In addition to nonlinear mixed effects model designations, Pumas allows for the specification of clinical dosage regimens. These dosage regimens are modeled as discontinuities to the differential equation and can be specified using a standard clinical dataset or by using the programmatic `DosageRegimen` method directly from Julia code. Figure 2 showcases the result of a Pumas simulation of this model with steady state dosing against the pharmaceutical software NONMEM [4], showcasing that Pumas recovers the same values. The test suite for the dosing mechanism is described in Appendix 5. Appendix Figures 9 and 10 demonstrate similar results across a 20 different dosage regimens using numerical ODE solving and analytical approach for accelerated handling of linear dynamics. One interesting feature of Pumas demonstrated here is the ability to specify the continuity of the observation behavior. Because of the discontinuities at the dosing times, the drug concentration at the time of a dose is not unique and one must choose the right continuous or the left continuous value. Previous pharmaceutical modeling software, such as NONMEM, utilize right-continuous values for most discontinuities but left-continuous values in the case of steady state dosing. To simplify this effect for the user, Pumas defaults to using the right-continuous values, but allows the user to change to left continuity through the `continuity` keyword argument. By using right continuity, the observation is always the one captured post-dose. Since the simulations usually occur after the dose, this makes the observation series continuous with less outlier points, a feature we believe will reduce the number of bugs in fitting due to misspecified models.

## 1.3 Maximum Likelihood and Bayesian Model Fitting

To understand how a drug interacts within the body, we can either simulate populations of individuals and dosage regimens or utilize existing patient data to estimate our parameters. Transitioning from a simulation

approach to a model estimation approach is simply the change of the verb applied to the model object. The follow code demonstrates how moving between simulation and fitting in Pumas is a natural transition:

```
## Specify the parameters to simulate from
fixeffs = (
  θ = [0.4, 20, 1.1, 2],
  Ω = diagm(0 => [0.04, 0.04, 0.04]),
  σ = 0.04
)

## Create a dosage regimen
ev = DosageRegimen(100, time=0)

## Create subject(s) with doses and covariates
s1 = Subject(id=1, events=ev, covariates=(sex=0, wt=70))
s2 = Subject(id=2, events=ev, covariates=(sex=1, wt=70))

## A population is a vector of subjects
data = [s1, s2]

## Simulate using model, data, parameters over a time range
obs = simobs(model, data, fixeffs, obstimes=0:0.1:120)

## Model Fitting
data = read_pumas("data_path", covariates=[:sex,:wt])
res = fit(model, data, fixeffs, Pumas.FOCEI()) # Fit using the FOCEI likelihood approximation

## Simulate using fitted model parameters
obs = simobs(model, data, coef(res), obstimes=0:0.1:120) # Check the fit
```

This example first generates a data set using `simobs` and then proceeds to fit the parameters of the model to data using `fit`. For fitting one currently has a choice between using:

1. FO: a first order approximation to the likelihood integral
2. FOCE: a first order conditional estimation of the likelihood integral without interaction
3. FOCEI: a first order conditional estimation of the likelihood integral with interaction
4. LaplaceI: a second order Laplace approximation of the likelihood integral with interaction
5. BayesMCMC: A Bayesian Markov Chain Monte Carlo estimation of model posteriors using the Hamiltonian Monte Carlo no-U-turn sampler (NUTS)<sup>1</sup>

We use the `Optim.jl` package [27] for the numerical optimization and default to using a safeguarded BFGS algorithm with a backtracking line search for the fixed effects and Newton's method with a trust region strategy for the empirical bayes estimates. The result of the fitting process is a `FittedPumasModel` which we call `res`. This object can be inspected to determine many quantities such as confidence intervals (as demonstrated later). Importantly, `coef(res)` returns the coefficients for the fixed effects in a `NamedTuple` that directly matches the style of `fixeffs`, and thus we demonstrate calling `simobs` on the returned coefficients as a way to check the results of the fitting process. When applied to the model of Section 1.1, this process returns almost exactly the coefficients used to generate the data, and thus this integration between the simulation and fitting models is routinely used within Pumas to generate unit tests of various methodologies.

## 2 Generalized Quantitative Pharmacology Models with Pumas

The following sections detail featured which are new to pharmaceutical modeling software or enhanced via parallelization or direct integration. In summary:

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<sup>1</sup>Prior distributions are required for this fitting method and are specified in the `param` block by using a distribution instead of a domain.

1. We show how the integrated Non-compartmental Analysis (NCA) module can be mixed with NLME simulation and estimation.
2. We show how alternative dynamical formulations, such as stochastic differential equation can be directly utilized in the NLME models.
3. We showcase how the Pumas generalized error distribution form allows for mixing discrete and continuous data and likelihoods
4. We detail the model diagnostics and validation tools
5. We demonstrate the functionality for optimal design of experiments.

## 2.1 Integrated Non-compartmental Analysis (NCA)

NCA is a set of common analysis methods performed in all stages of the drug development program, pre-clinical to clinical, that has strict rules and guidelines for properly calculating diagnostic variables from experimental measurements [11]. In order to better predict the vital characteristics, Pumas includes a fully-featured NCA suite which is directly accessible from the nonlinear mixed effects modeling suite. Appendix Figures 11 and 12 demonstrate that Pumas matches industry standard software such as PKNCA [9] and Phoenix on a set of 13104 scenarios and 78,624 subjects as described in Section 5. The following code showcases the definition of a NLME model where the observables are derived quantities calculated through the NCA suite, demonstrating the ease at which a validated NCA suite can be coupled with the simulation and estimation routines.

```
m_diffEq = @model begin
  @param begin
    θ ∈ VectorDomain(5, lower=zeros(5), init=ones(5))
    Ω ∈ PSDDomain(2)
    σ_prop ∈ RealDomain(init=0.1)
  end

  @random begin
    η ~ MvNormal(Ω)
  end

  @pre begin
    Ka = θ[1]
    CL = θ[2]*exp(η[1])
    V = θ[3]*exp(η[2])
    lags = [0,θ[4]]
    bioav = [1,θ[5]]
  end

  @covariates isPM Wt

  @dynamics begin
    Depot' = -Ka*Depot
    Central' = Ka*Depot - (CL/V)*Central
  end

  @derived begin
    cp := @. 1000*(Central / V)
    nca := @nca cp
    auc = NCA.auc(nca)
    thalf = NCA.thalf(nca)
    cmax = NCA.cmax(nca)
  end
end
```

## 2.2 Alternative Differential Equations via A Function-Based Interface: Stochastic, Delay, and Differential-Algebraic Equations for Pharmacology

While ODEs are a common form of dynamical model, in many circumstances additional features are required in order for the realism of the model to adequately predict the behavior of the process. For example, many biological processes are inherently stochastic, and thus an extension to ordinary differential equations, known as stochastic differential equations, takes into account the continuous randomness of biochemical processes [34, 36, 22, 46]. Many biological effects are delayed, in which case a delay differential equation which allows for an effect to be determined by a past value of the system may be an appropriate model [19, 23]. Together, Pumas allows for the definition of dynamical systems of the following non-standard forms:

1. Split and Partitioned ODEs (Symplectic integrators, IMEX Methods)
2. Stochastic ordinary differential equations (SODEs or SDEs)
3. Differential algebraic equations (DAEs)
4. Delay differential equations (DDEs)
5. Mixed discrete and continuous equations (Hybrid Equations, Jump Diffusions)

where each can be simulated with high-performance adaptive integrators with specific method choices stabilized for stiff equations. The following code demonstrates the definition solving of a stochastic differential equation model with steady state dosing via a high strong order adaptive SDE solver specified using an alternative interface known as the Pumas function-based interface. This interface is entirely defined with standard Julia functions, meaning that any tools accessible from Julia can be utilized in the pharmacological models through this context.

```
using StochasticDiffEq

θ = [
    1.5, #Ka
    1.0, #CL
    30.0 #V
]

p = ParamSet((θ=VectorDomain(3, lower=zeros(4), init=θ), Ω=PSDDomain(2)))

function randomfx(p)
    ParamSet((η=MvNormal(p.Ω),))
end

function pre_f(params, randoms, subject)
    function pre(t)
        θ = params.θ
        η = randoms.η
        Ka = θ[1],
        CL = θ[2]*exp(η[1]),
        V = θ[3]*exp(η[2])
    end
end

function f(du,u,p,t)
    Depot,Central = u
    du[1] = -p.Ka*Depot
    du[2] = p.Ka*Depot - (p.CL/p.V)*Central
end

function g(du,u,p,t)
    du[1] = 0.5u[1]
    du[2] = 0.1u[2]
end
```



```
prob = SDEProblem(f,g,nothing,nothing)

init_f(col,t) = [0.0,0.0]

function derived_f(col,sol,obstimes,subject, param, randeffs)
    V = col.V
    Σ = col.Σ
    central = sol(obstimes;idxs=2)
    conc = @. central / V
    dv = @. Normal(conc, conc*Σ)
    (dv=dv,)
end

function observed_f(col,sol,obstimes,samples,subject)
    samples
end

model = Pumas.PumasModel(p,randomfx,pre_f,init_f,prob,derived_f,observed_f)

param = init_param(model)
randeffs = init_randeffs(model, param)

data = Subject(events = DosageRegimen([10, 20], ii = 24, addl = 2, ss = 1:2,
                                     time = [0, 12], cmt = 1))
sol = solve(model,data,param,randeffs,alg=SRIW1())
plot(sol)
```

### 2.3 Generalized Error Distribution Abstractions

Traditional nonlinear mixed effects models formulate the error distribution handling in a different manner, focusing on the perturbation via the error against the estimate. Mathematically, this amounts to having an  $\epsilon$  measurement noise, i.e. defining the observables as:

$$y_{ij} = h_j(u_i, p_i, Z_i, \eta_i, \theta) + \epsilon_{ij} \quad (8)$$

Instead of using the perturbation format, Pumas allows for users to specify the likelihood directly via any arbitrary `Distributions.jl` [5] distribution, with the mean of the distribution corresponding to the prediction. This has some immediate advantages. For one, many distributions are not directly amenable to the perturbation form. For example, one may wish to model a discrete observable, such as a pain score, probabilistically based on clinical outputs. A discrete likelihood function, like a Poisson distribution, can thus be given to described the predicted outputs like in the following example:

```
poisson_model = @model begin
    @param begin
        tvbase ∈ RealDomain(init=3.0, lower=0.1)
        d50 ∈ RealDomain(init=0.5, lower=0.1)
        Ω ∈ PSDDomain(fill(0.1, 1, 1))
    end

    @random begin
        η ~ MvNormal(Ω)
    end

    @covariates dose

    @pre begin
        baseline = tvbase*exp(η[1])
        Dose = dose
        D50 = d50
    end
end
```

```
end

@derived begin
  dv ~ @. Poisson(baseline*(1-Dose/(Dose + D50)))
end
end
```

This type of model could not be described with continuous perturbations and thus would have to be approximated in other scenarios. In addition, since any distribution is possible in this format, users can extend the modeling schema to incorporate custom distributions. Discrete Markov Chain models, or Time to Event models for example can be implemented as a choice of a custom distribution, thus making it easy to extend the modeling space directly from standard language use.

## 2.4 Integrated and Parallelized Model Diagnostics and Validation

After running the estimation procedure, it is important to have a wide suite of methods to post-process the results and validate the predictions. Model diagnostics are used to check if the model fits the data well. Pumas provides a comprehensive set of diagnostics tooling. A number of residual diagnostics are available as well as shrinkage estimators. Additionally, for model validation, Visual Predictive Checks (VPCs) [18] can be used for a variety of models in a performant and robust manner.

The diagnostics tooling comprises of:

1. Residuals: Populations residuals take the inter-individual variability into account as opposed to Individual residuals which are correlated due to the lack of accounting for inter-individual variability.

The following Population residuals are available in Pumas<sup>2</sup>:

- Normalized Prediction Distribution Errors (NPDE) [26]
- Weighted Residuals (WRES)
- Conditional Weighted Residuals (CWRES)
- Conditional Weighted Residuals with Interaction (CWRESI)

Similarly the following Individual residuals are also available -

- Individual Weighted Residuals (IWRES)
- Individual Conditional Weighted Residuals (ICWRES)
- Individual Conditional Weighted Residuals with Interaction (ICWRESI)
- Expected Simulation based Individual Weighted Residuals (EIWRES)

2. Population Predictions: Population prediction are defined as the difference between the observations and the model expectation for subject  $i$ , i.e.  $y_i - \mathbf{E}[y_i]$  where the  $\mathbf{E}[y_i]$  are the population predictions. The population predictions, EPRED, PRED, CPRED and CPREDI are defined as the  $\mathbf{E}[y_i]$  from equations of NPDE, WRES, CWRES and CWRESI respectively.

3. Individual Predictions: The individual residuals are defined as the difference between the observations and the model prediction for subject  $i$ , i.e.  $y_i - f_i(\theta, \eta_i^*, Z_i, t_i)$  where the  $f_i(\theta, \eta_i^*, Z_i, t_i)$  are the individual predictions. The individual predictions IPRED, CIPRED and CIPREDI are defined as the  $f_i(\theta, \eta_i^*, Z_i, t_i)$  from equations of IWRES, ICWRES and ICWRESI. The individual expected prediction (EIPRED) is defined as the individual mean conditioned on  $\eta_k$ , i.e.  $\mathbf{E}[y_i|\eta^k]$ , from equation of EIWRES.

4. Akaike Information Criterion (AIC) and Bayesian Information Criterion (BIC)

5.  $\eta$  Shrinkage and  $\epsilon$  Shrinkage [35]

The above discussed diagnostics can be evaluated through the `inspect` function and further exported as a `DataFrame`. Additionally, the `infer` function is available for computing the covariance matrix of the population parameters thus giving the the 95% confidence intervals and the standard errors for the parameter estimates. As an example below we call `infer` and `inspect` on the `FittedPumasModel` object `res` obtained as the result of `fit` call in earlier section.

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<sup>2</sup>Defined in [28] unless mentioned otherwise

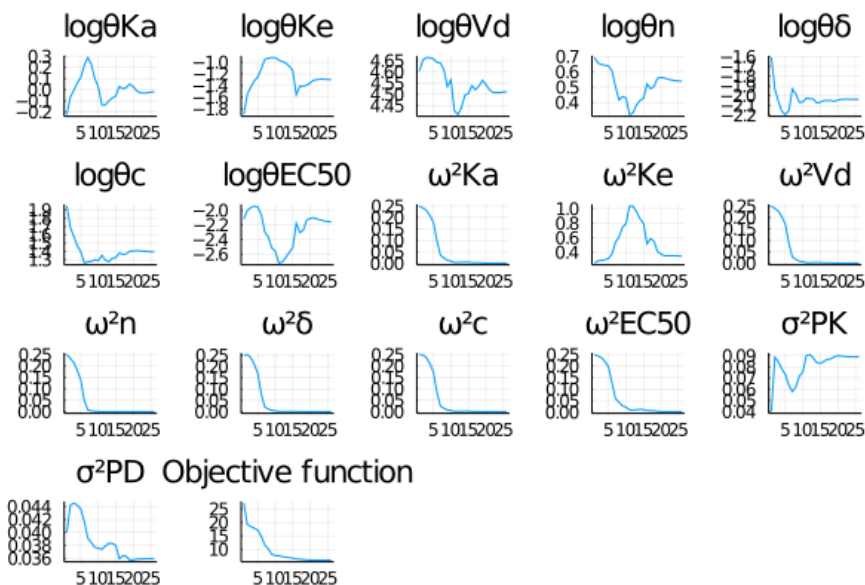


Figure 3: Convergence Plot for the HCV model

```
resinfer = infer(res)
DataFrame(resinfer)
resinspect = inspect(res)
DataFrame(resinspect)
```

Additionally several out of the box visualizations are available for practitioners to evaluate the model fit are are mainly provided by the PumasPlots.jl package. Convergence plot for fitting obtained with the `convergencefunction` for the HCV model is shown in the following Figure 3.

```
convergence(res)
```

In Pumas we provide the ability to run Visual Predictive Checks with `vpc` function. For continuous models VPCs are computed using the Quantile Regression based approach discussed in [20]. The syntax for computing and plotting the VPCs is shown below followed by the Figure 4 of a stratified VPC plot in Pumas.

```
vpc_res = vpc(res, stratify_by = [:Wt])
using Plots
plot(vpc_model)
```

## 2.5 Optimal Design of Experiments

Optimal design of clinical trials has increasingly played a central role in increasing the effectiveness of such studies while minimizing costs [25]. To facilitate the advancement of optimal design methodologies into stan-

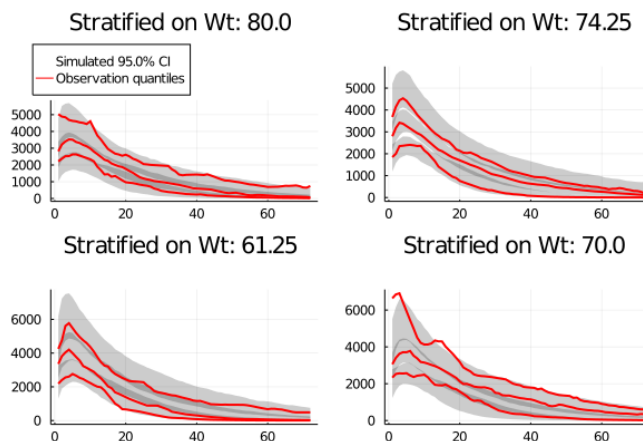


Figure 4: **Visual Predictive Check**

standard clinical trial workflows, Pumas includes functionality for high performance calculations of the Fisher Information Matrix (FIM) which is central to optimal design methodologies. The function `informationmatrix` takes in a `FittedPumasModel`, the output of the `fit` estimation function, and computes either the expected or observed FIM dependent on a keyword argument `expected` using a normality approximation [40]. The following demonstrates the workflow for performing parameter estimation with FO and calculating the expected FIM in the Warfarin Pk model [29]:

```
data = read_pumas(example_data("warfarin"))

model = @model begin

  @param begin
    θ1 ∈ RealDomain(lower=0.0, init=0.15)
    θ2 ∈ RealDomain(lower=0.0, init=8.0)
    θ3 ∈ RealDomain(lower=0.0, init=1.0)
    Ω ∈ PDiagDomain(3)
    σ ∈ RealDomain(lower=0.0001, init=sqrt(0.01))
  end

  @random begin
    η ~ MvNormal(Ω)
  end

  @pre begin
    Tvcl = θ1
    TvV = θ2
    Tvka = θ3
    CL = Tvcl*exp(η[1])
    Vc = TvV*exp(η[2])
    Ka = Tvka*exp(η[3])
  end

  @dynamics Depots1Central1
```

```
@vars begin
  conc = Central / Vc
end

@derived begin
  dv ~ @. Normal(log(conc),  $\sigma$ )
end

param = ( $\theta_1$  = 0.15,
          $\theta_2$  = 8.0,
          $\theta_3$  = 1.0,
          $\Omega$  = Diagonal([0.07, 0.02, 0.6]),
          $\sigma$  = sqrt(0.01))

ft = fit(model, data, param, Pumas.FO())
fim = informationmatrix(ft)
```

Note that the derivative calculations used within the FIM calculation are the discrete sensitivity analysis derivatives derived via forward-mode automatic differentiation as described in Section 3.2.

### 3 High Performance and Stability-Enhanced Model Fitting

#### 3.1 Fine-Tuning Differential Equation Solver Behavior to Model Features

The core computation of the model fitting process utilizes the `simobs` function for generating solutions to the differential equation during the likelihood approximation, meaning that every step of the optimization is solving thousands of the same small differential equation representing different possible parameter configurations amongst all subjects in a clinical trial. Thus, while in isolation these small ODEs may simulate very fast, real-world NLME model fitting with large numbers of subjects consistently arrives at workflows which take hours to days or weeks with the majority of the time due to the cost of solving small ODEs. Thus practical workflows of industry pharmacologists would be heavily impacted if the speed of these systems could be dramatically decreased.

Pumas recognizes the crux of the computational issue and thus has many new features for optimizing the internal solve of the fitting process. The options for controlling the solvers are same between the simulation and estimation workflows. The full gamut of options from `DifferentialEquations.jl` are exposed to allow users to control the solvers as much as possible. This allows for specializing the solver behavior on the known characteristics of the functions and its solution. For example, the concentrations modeled in the ODEs need to stay positive in order for the model to be stable, but numerical solvers of ODEs do not generally enforce this behavior which can cause divergences in the optimization process. In Pumas, one can make use of advanced strategies [38] like rejecting steps out of the domain by using `isoutofdomain` or using the `PositiveDomain` callback.

However, a more immediate effect of this connection is the ability to choose between a large set of highly optimized integration methods. Table 1 shows timing results of Pumas on pharmacokinetic (PK) and pharmacokinetic/pharmacodynamic (PK/PD) models using both the native `DifferentialEquations.jl` methods and some classic C++ and Fortran libraries and demonstrates a performance advantage averaging around 3x between the best Julia-based method against the best wrapped C++ or Fortran solver method. This table also demonstrate a few different dimensions by which this performance advantage is achieved. First of all, the `DifferentialEquations.jl` library uses different Runge-Kutta methods which are derived to have asymptotically better error qualities for the same amount of work [42, 44], with a much larger effect for the high order (7th order) integrator. Secondly, `DifferentialEquations.jl` uses a tuned PI-adaptive timestepping method [39] which is able to stabilize the solver and increase the timesteps, thus decreasing the total amount of work to integrate the equation. Lastly, Pumas is able to utilize the JIT compiler to compile a form of the differential equation solver that utilizes stack-allocated arrays and is specific to the size and ODE function. While this optimization only applies to small ODE systems (the optimization is no longer beneficial at around 10 or more ODEs), many pharmacometric models fall into this range of problems and these benchmarks demonstrate that it can have a noticeable effect on the solver time.

ODE Solver vs Time (seconds)	OC	OC(LT)	MR	MR(LT)	HCV	HCV(LT)
AutoTsit5-Rosenbrock23 (SArray)	0.000057	0.000216	0.000051	0.000222	0.000133	0.000542
Tsit5 (SArray)	<b>0.000038</b>	0.000173	<b>0.000034</b>	0.000186	<b>0.000110</b>	0.000475
Vern7 (SArray)	0.000050	<b>0.000140</b>	0.000053	<b>0.000141</b>	0.000160	<b>0.000423</b>
AutoTsit5-Rosenbrock23 (Array)	0.000211	0.000359	0.000233	0.000420	0.000360	0.000787
Tsit5 (Array)	0.000043	0.000207	0.000052	0.000222	0.000138	0.000634
Vern7 (Array)	0.000068	0.000169	0.000075	0.000166	0.000207	0.000522
CVIDE (Array)	0.000091	0.000294	0.000119	0.001456	0.000400	0.001075
DOP853 (Array)	0.000184	0.000647	0.000191	0.000765	0.000410	0.001829
DOPRI (Array)	0.000205	0.000330	0.000239	0.000397	0.000464	0.001364

Table 1: **Effect of specialized ODE solvers on forward simulation of small pharmacometric ODE models.** Shown is the effect of the ODE solver choice on the speed of the forward pass of common pharmacometric models. OC for the one-compartment model of Section 1.1, MR for the multiple response model described in Section 5, and HCV for the hepatitis C model described in Section 5. Times are all shown in seconds. By default the tolerances for the solvers was  $1 \times 10^{-3}$  for the relative tolerance and  $1 \times 10^{-6}$  for the absolute tolerance, while LT stands for low tolerance with  $1 \times 10^{-8}$  for the relative tolerance and  $1 \times 10^{-12}$  for the absolute tolerance (representative of tolerances used when simulating vs when simulating). Solutions were checked against a tolerance  $1 \times 10^{-14}$  reference solution to ensure the actual errors were within the same order of magnitude. The automatic stiffness detection AutoTsit5-Rosenbrock23 method is a combination between Tsit5 [42] and a Rosenbrock method Rosenbrock23 [37], while Tsit5 and Vern7 [44] are explicit Runge-Kutta methods from the DifferentialEquations.jl library [33]. CVIDE is from the Sundials library [17], while DOP853 and DOPRI are from the Hairer Fortran methods suite [13]. The Array methods are generically compiled for heap-allocated mutable arrays, where the SArray versions are specially optimized for the specific size of the ODE using Julia’s JIT compiler.

ODE Solver	PBPK	PBPK(LT)
AutoVern7-Rodas5	0.168	<b>0.695</b>
Rodas5	<b>0.143</b>	1.072
KenCarp4	0.865	3.167
RadauIIA5	0.183	1.973
CVIDE	0.505	1.472

Table 2: **Effect of specialized ODE solvers on forward simulation the Voriconazole physiologically-based pharmacokinetic (PBPK) model [48].** Shown is the effect of the ODE solver choice on the speed of the forward pass of the PBPK model with steady state dosing. Given the high degree of variance in the stiff ODE solver accuracies at the same tolerances, the tolerances were aligned using a reference solution computed at  $1 \times 10^{-14}$  tolerances, and high tolerance was chosen to be the tolerance pair by power of 10 which achieve a true error closest to  $1 \times 10^{-5}$  and for low tolerance the pair closes to  $1 \times 10^{-9}$ . The automatic stiffness detection method AutoVern7-Rodas5 is a combination between Vern7 and Rodas5, while Rodas5 [45], KenCarp4 [21], and RadauIIA5 [14] are implicit methods from the DifferentialEquations.jl library [33]. CVIDE is from the Sundials library [17]

One additional advantage of this tweak-ability is the ease to span multiple domains. Physiologically-based pharmacokinetic (PBPK) models are typically larger stiff ODE-based models which incorporates systems-type mechanistic modeling ideas to enhance the model’s predictive power [24]. Table 2 demonstrates the performance advantage of the native Julia methods that are unique to Pumas on a 15 stiff ODE PBPK model with steady state dosing, demonstrating a 2x-3x performance advantage over the classic CVIDE method used in many other pharmacometric modeling suites.

### 3.2 Fast and Accurate Likelihood Hessian Calculations via Automatic Differentiation

When performing maximum likelihood estimation or Bayesian estimation with a gradient-based sampler like Hamiltonian Monte Carlo, the limiting step is often the calculation of the gradient of the likelihood. Finite difference calculations are not efficient since every calculation of a perturbation involves a numerical solve of the ODE system and either one or two perturbations are required for each model parameter for forward and central differencing respectively. Furthermore, the finite difference approximation of a derivative is an unstable process [10] which in some cases can result in very inaccurate gradients when combined numerical solutions to ODEs.

	AD Almquist	FD Almquist	FD simple
Relative error	<b>9.52e-7</b>	0.811	0.00047
Time in seconds	<b>0.431</b>	0.641	18.6

Table 3: **Timings and relative errors for the gradient computations of the marginal likelihood in a non-linear fixed effects model with time varying coefficients in the ODE.** Shown are the timings of the three likelihood gradient calculation methods along with their relative error. AD stands for automatic differentiation while FD stands for finite differentiation, demonstrating both the efficiency and accuracy gains obtained through by utilizing the automatic differentiation derived discrete sensitivity analysis.

The performance of the derivative calculations for the marginal likelihoods can be improved by utilizing a formulation with the sensitivity equations due to [1]. While very efficient, the method relies on second order derivatives in a way that makes the process unstable and accurate derivatives are required for process to work well. Instead of utilizing the traditional form of the sensitivity equations, Pumas generates an implementation of discrete forward sensitivity analysis by utilizing dual number arithmetic through the differential equation solver (cite ForwardDiff). Our group has previously shown that this discrete sensitivity analysis via automatic differentiation outperforms traditional sensitivity analysis since it allows the compiler more freedom to optimize the generated code for passes like single instruction, multiple data (SIMD) auto-vectorization [32].

Table 3.2 demonstrates the effect on run time and accuracy of using the sensitivity method from [1] with finite difference and automatic differentiation based derivatives respectively as well as a simple but expensive finite difference based gradient computation. The error is computed relative to a solution computed with 256 bit precision floating point numbers. All ODE solutions are computed to a relative tolerance of  $10^{-8}$ . The results show that the simple finite difference based gradient doesn't have too large an error but is slow while the faster sensitivity based method has a large relative error. The error is so large that gradient based optimization is no longer practical. In contrast, the sensitivity based gradient computation based on automatic differentiation loses almost no precision and is even faster than the finite difference based gradient because it requires fewer evaluations of the objective function.

### 3.3 Accelerated Maximum Likelihood Fitting

Maximum likelihood directly corresponds to repeated calculation of likelihood gradients. Given the performance advantages that are obtained due to the ODE solver and discrete sensitivity analysis advantages over previous software, one would predict that the model estimation routines would see a similar benefit as derived from these components. Appendix 5 describes the full set of benchmark models for maximum likelihood estimation. In Figure 5 we demonstrate this is the case by calculating the elapsed run time to estimate 6 models in both Pumas and NONMEM. We note that the Pumas fitting mechanism has a higher fixed cost which effects the results on the smallest models, but on the two larger models we see more than a 2x performance advantage for Pumas, similar to the difference between the non-fully optimized ODE solver and the alternative array-based algorithms and implementations. Figure 6 demonstrates that this performance advantage over NONMEM extends to larger nonlinear models, showcasing the general applicability of the performance enhancements seen in Pumas. In all of the occasions the resulted in similar likelihoods.

### 3.4 Automated Parallelism and Full-Application Benchmarks

Pumas is able to automatically parallelize the solution of the NLME model solution across subjects. 2 forms of parallelism are currently available:

1. Multithreaded parallelism for shared memory machines
2. Multiprocessing for distributed memory architectures

Multithreaded is enabled by default and no extra steps are required for this parallelism to occur. Meanwhile the multiprocessing allows a user to utilize cores across any cluster with passwordless SSH, which includes architectures like traditional high-performance computing (HPC) clusters and cloud architectures like Azure. Figure 7 demonstrates the good scaling of multithreading on a PK/PD maximum likelihood estimation. The scaling does not meet the theoretical optimal properties of linear scaling because there are parts of the fitting process that cannot be parallelized.

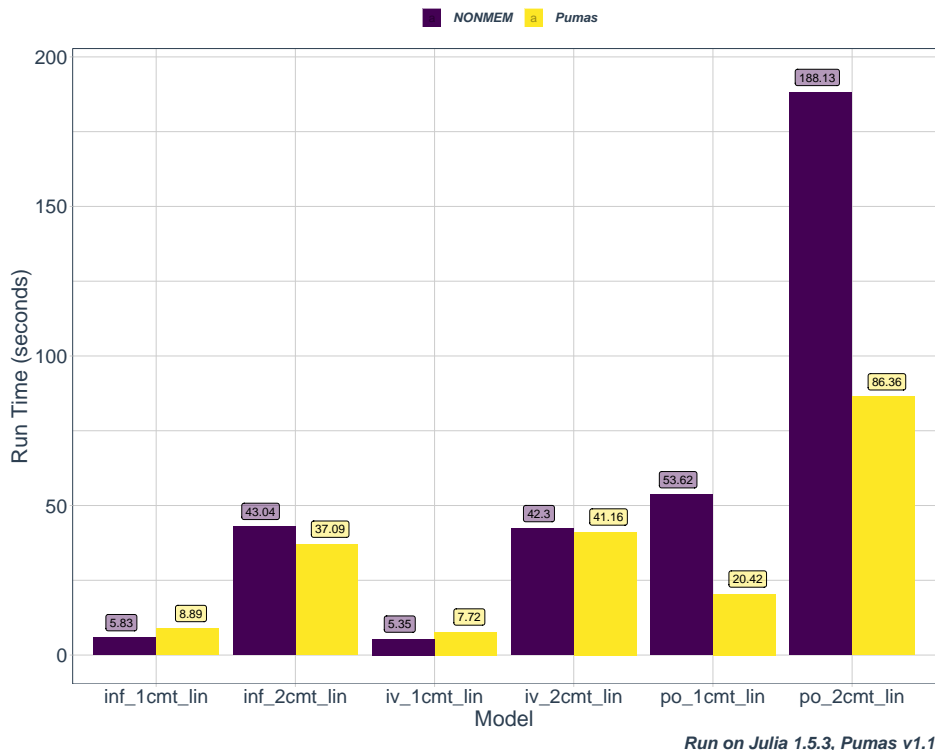


Figure 5: **Maximum likelihood estimation results.** Shown is the run time of a fit of the various Pk/Pd models

### 3.5 High Performance Non-Compartmental Analysis (NCA)

In the previous sections we demonstrated that the integrated NCA suite reproduced the results of industry-standard tools, demonstrating the correctness of the implementation over 13104 scenarios. In addition to determining the correctness, we calculated the run times. The full analysis tool 3 hours and 16 minutes in PKNCA while in Pumas the run time was 56 seconds, demonstrating a 210x acceleration. This speed is particularly useful when 1,000s of clinical trials are simulated during the planning of a large patient trial.

### 3.6 Accelerated Optimal Design of Experiments

In order to assess the effectiveness of accelerated differential equation solving on optimal design workflows, we tested the calculation time of the FIM on the Warfarin PK model (Section 2.5) and the HCV PK/PD model Section 5 against the PopED Population Optimal Experimental Design framework [30]. Figure 8 showcases that Pumas calculates the FIM approximately 20x faster than PopED on the Warfarin model and approximately 220x faster on the HCV model. Much of the acceleration can be attributed to the `AutoVern7(Rodas5())` ODE solver used in the Pumas version of the FIM calculation as opposed to the `ode45` method internally used by PopED. This demonstrates how central improvements to the solving of small ODEs can give large advantages to real applications built on top of such methodologies.

## 4 Conclusion

Pumas pulls together a diverse array of pharmacometrics problems into a single platform and utilizes the JIT optimization to directly specialize the internal solvers. We have showcased how the Pumas platform achieves across the board performance improvements from preclinical to clinical analyses over existing pharmacometrics tools. This more broadly illustrates how integrating optimizing compilers into dynamic tool chains can improve performance over traditional approaches which do not specialize on the problem's scale. Even some-



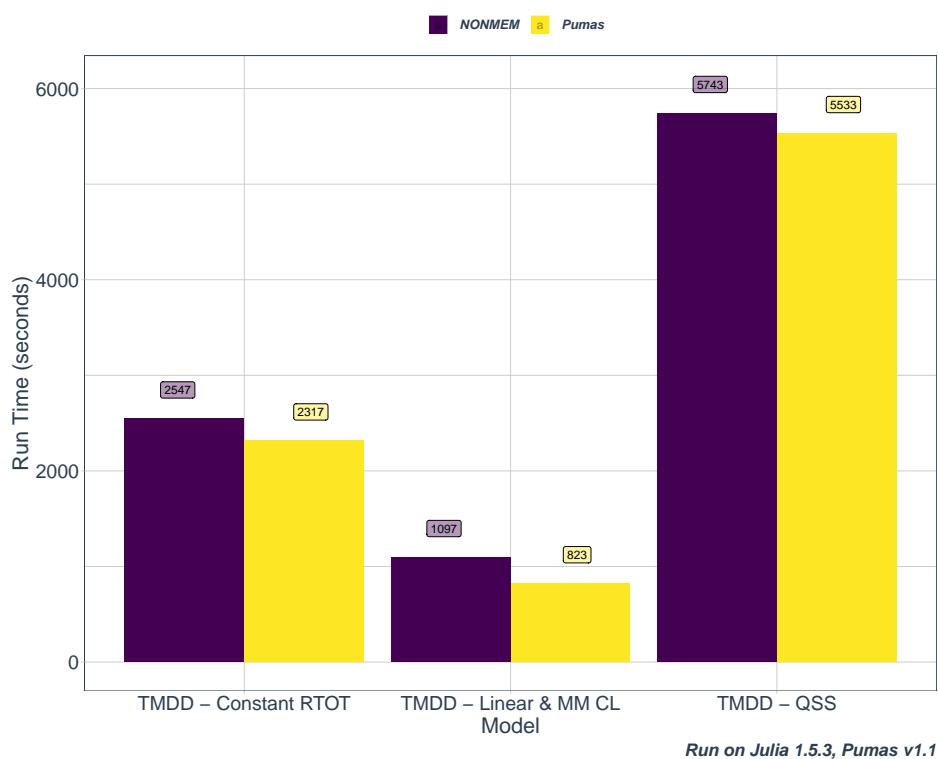


Figure 6: **Runtime comparisons of non-linear mechanistic TMDD models.** Shown is the run time of a fit of the Michaelis-Menten (MM) model, Constant  $R_{tot}$  model and Rapid binding (QE) and quasi steady-state (QSS) models

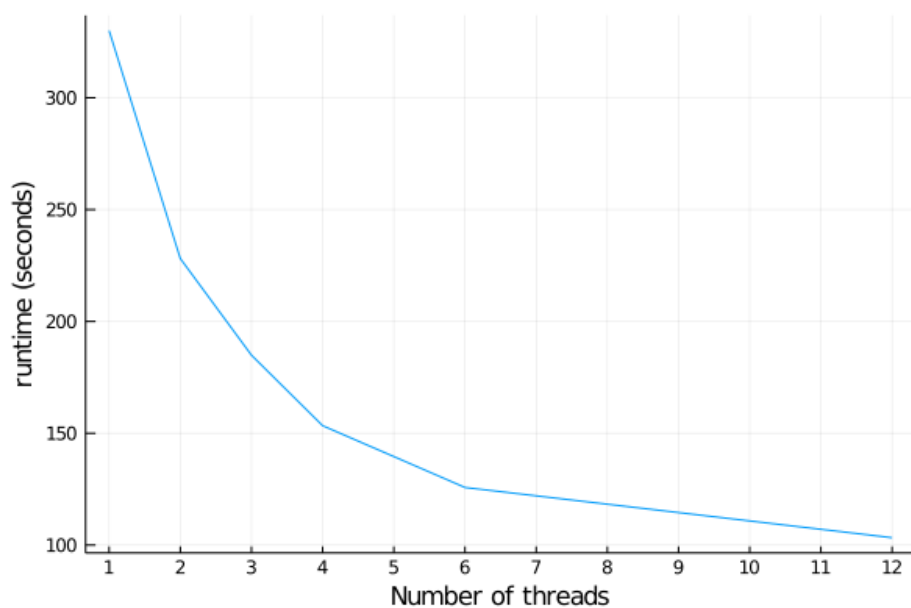


Figure 7: **Scaling of multithreaded fitting of a HCV model.** Shown is the run time of a fit of a simultaneous PK/PD model HCV model described in 5 with multithreading with 1, 2, 3, 4, 6, 8 and 12 threads.

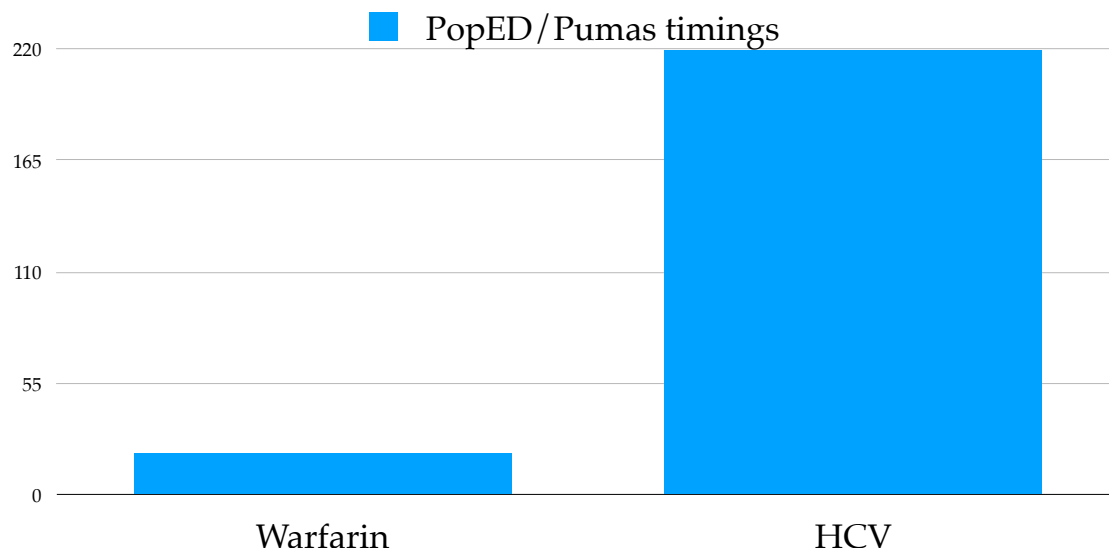


Figure 8: **Accelerated Optimal Design of Experiments.** Shown is the relative time to compute the Fisher Information Matrix (FIM) between PopED [30] and Pumas on the Warfarin and HCV models. The FIM was calculated 1000 times in order to reduce noise in the timings.

thing as standardized as an ODE solver can be improved. Pumas is still early in its development and will continue to increase its performance as it demonstrates new features to the pharmacometrics community.

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## Appendix

### Model Definitions

#### Multiple Response Model

$$\frac{d[\text{Ev1}]}{dt} = -K_e[\text{Ev1}] \quad (9)$$

$$\frac{d[\text{Cent}]}{dt} = K_e[\text{Ev1}] - \frac{[\text{Cent}]}{V_c} \left( CL + \frac{V_{max}}{K_m + [\text{Cent}]} + Q \right) + Q \frac{[\text{Periph}]}{V_p} \quad (10)$$

$$\frac{d[\text{Periph}]}{dt} = Q \frac{[\text{Cent}]}{V_c} - Q \frac{[\text{Periph}]}{V_p} \quad (11)$$

$$\frac{d[\text{Resp}]}{dt} = K_{in} \left( 1 - \frac{IMAX \left( \frac{[\text{Cent}]}{V_c} \right)^\gamma}{IC_{50}^\gamma + \left( \frac{[\text{Cent}]}{V_c} \right)^\gamma} \right) - K_{out}[\text{Resp}] \quad (12)$$

where:

$$K_e = 1 \quad (13)$$

$$CL = 1 \quad (14)$$

$$V_c = 20 \quad (15)$$

$$Q = 2 \quad (16)$$

$$V_p = 10 \quad (17)$$

$$K_{in} = 10 \quad (18)$$

$$K_{out} = 2 \quad (19)$$

$$IC_{50} = 2 \quad (20)$$

$$IMAX = 1 \quad (21)$$

$$\gamma = 1 \quad (22)$$

$$V_{max} = 0 \quad (23)$$

$$K_m = 2 \quad (24)$$

#### HCV Model

The "HCV Model" is a multiple response PKPD model from [29] that models the effect of a pegylated interferon dose given as a 24h infusion once a week for 4 weeks. Twelve simulated samples are obtained for the PK and PD at times  $t = [0.0, 0.25, 0.5, 1.0, 2.0, 3.0, 4.0, 7.0, 10.0, 14.0, 21.0, 28.0]$ . The internal dynamics are:

$$\frac{d[X]}{dt} = -k_a[X], \quad X(0) = 0 \quad (25)$$

$$\frac{d[A]}{dt} = k_a[X] - k_e[A], \quad A(0) = 0 \quad (26)$$

$$\frac{d[T]}{dt} = s - [T] \cdot (e[W] + d), \quad T(0) = \frac{c\delta}{pe} \quad (27)$$

$$\frac{d[I]}{dt} = e[W][T] - \delta[I], \quad I(0) = \frac{sep - dc\delta}{p\delta e} \quad (28)$$

$$\frac{d[W]}{dt} = p \left( 1 - \frac{[C]^n}{[C]^n + EC_{50}^n} \right) [I] - c[W], \quad W(0) = \frac{sep - dc\delta}{c\delta e} \quad (29)$$

where  $C(t) = A(t)/V_d$ . The PK variable  $C$  is modelled with an additive normal distribution and the PK variable  $W$  is modelled as being log-normal. We use the parameters reported in the original paper.

## Dosing Regimen Verification

The event handling of Pumas and the accuracy of simulation under different kinds of events was evaluated systematically. The examples for this testing were adapted from `mrgsolve` test suite<sup>3</sup>. The dosing and sampling scenarios were setup in R's `mrgsolve` package which were then used to simulate concentration time profile for a single subject in NONMEM and Pumas. The listing of all the tested scenarios is presented below:

1. Infusion (10 mg/hr) into the central compartment with 4 doses given every 12 hours
2. Infusion (10 mg/hr) into the central compartment with lag time (5 hr) with 4 doses given every 12 hours
3. Infusion (10 mg/hr) into the central compartment with lag time (5 hr) and bioavailability (0.4) with 4 doses given every 12 hours
4. Infusion (10 mg/hr) into the central compartment at steady state (ss)
5. Infusion (10 mg/hr) into the central compartment at steady state (ss) with 81% bioavailability, where frequency of events (ii) is less than the infusion duration (DUR)
6. Infusion (10 mg/hr) into the central compartment at steady state (ss) with 100% bioavailability, where frequency of events (ii) is less than the infusion duration (DUR)
7. Infusion (10 mg/hr) into the central compartment at steady state (ss) with 100% bioavailability, where frequency of events (ii) is a multiple of infusion duration (DUR)
8. Infusion (10 mg/hr) into the central compartment at steady state (ss) with 41% bioavailability, where frequency of events (ii) is exactly equal to the infusion duration (DUR)
9. Infusion (10 mg/hr) into the central compartment at steady state (ss) with 100% bioavailability, where frequency of events (ii) is exactly equal to the infusion duration (DUR)
10. Oral dose at steady state with lower bioavailability of 41%
11. Oral dose at steady state with lower bioavailability of 41% and 5 hour lag time
12. Zero order infusion followed by first order absorption into gut
13. Zero order infusion into central compartment specified by duration parameter
14. First order bolus into central compartment at ss followed by an  $ss=2$  (superposition ss) dose at 12 hours
15. First order bolus into central compartment at ss followed by an  $ss=2$  (superposition ss) dose at 12 hours followed by reset  $ss=1$  dose at 24 hours
16. Two parallel first order absorption models
17. Mixed zero and first order absorption

## Maximum Likelihood Tests

The Pumas-NLME parameter estimation algorithms, specifically *FOCEI*() were compared with NONMEM in a richly-sampled data setting with multiple models.

After a single administration 19 samples were collected over 72 hours via simulations for 4 different dose levels of 30 subjects each. A total of 6 test cases were generated that were a combination of:

- one- or two-compartmental disposition [26]
- oral (first-order absorption), intravenous (IV) bolus, or IV infusion administration.

The  $\Omega$ 's, representing the between subject variability were set to 30% CV, implemented as diagonal covariance matrix. A proportional residual error with 20% CV was chosen to capture the error distribution. For the structural model, all one-compartment models had a population  $V_c$  of 70 L, and all two-compartment models had an additional peripheral volume ( $V_p$ ) of 40 L. For all oral absorption models,  $K_a$  was set to 1.0 h<sup>-1</sup>. All models with linear elimination had a  $CL$  of 4.0 L/h. All two-compartment models had inter-compartmental clearance ( $Q$ ) set to 4.0 L/h.

<sup>3</sup><https://github.com/mrgsolve/nmtests/blob/master/nmtest7.md>

## TMDD Model Tests

While the maximum likelihood tests above compared the standard compartmental models, we also evaluated the performance of highly non-linear mechanistic models in the class of Target Mediated Drug Disposition (TMDD). Three models from this class were selected

1. Michaelis-Menten (MM) model
2. Constant  $R_{tot}$
3. Rapid binding (QE) and quasi steady-state (QSS) models

Data were simulated from a Full TMDD model in 224 subjects over a range of doses with a total of 3948 observations. The simulated data were then estimated using the reduced models listed above using NONMEM and Pumas

## NCA Implementation Verification

Simulations were performed in R with scenarios consisting of 1-, 2-, and 3-compartment models with typical parameters and  $\pm 4$ -fold from those typical parameters on the ratio of absorption rate ( $K_a$ ) to elimination rate ( $K_{el}$ ), ratio of peripheral volume of distribution 1 and 2 ( $V_{p1}$  and  $V_{p2}$ ) to central volume of distribution ( $V_c$ ), intercompartmental clearance between  $V_c$  and  $V_{p1}$  or  $V_{p2}$  ( $Q_{cp1}$  and  $Q_{cp2}$ ), and ratio of clearance ( $CL$ ) to  $V_c$  all models, as the parameters apply; with and without target-mediated drug disposition (TMDD); and oral and intravascular bolus dosing. All models were simulated with 4%, 10%, and 20% proportional residual error. Each model was simulated with 6 subjects. This yielded a total of 13104 scenarios and 78624 subjects simulated.

Each subject was then grouped with all other subjects in its simulation scenario, and 5, 10, and 20% of concentration measurements were set to below the limit of quantification (LOQ). NCA was to be performed on each of those LOQ scenarios in PKNCA [9], Pumas, and Phoenix<sup>4</sup> (a total of 707616 NCA intervals with calculations). Comparisons were made between the results of those NCA calculations performed on a single machine.

Five parameters, were chosen as the metrics for comparison as they included both observed and derived parameters:

1.  $AUC_{last}$
2.  $C_{max}$
3.  $T_{max}$
4.  $Half - life$
5.  $AUC_{inf}(pred)$

Eighteen percent of subjects were randomly selected from the 13104 scenarios to form a subset of 2367 subjects. This smaller subset was used to perform the NCA calculations across the three software while ensuring to maintain the same default options. All analysis were conducted on the same laptop and discussed at ACoP 2019 [47].

Results from all three software for the key NCA parameters match. Most results matched within  $\pm 0.1\%$  between all software. The difference between PKNCA /Pumas and Phoenix in half-life appears to be the result of PKNCA/Pumas selecting the best fit first and then filtering for decreasing slope while Phoenix first considers only consecutive sets of points that generate a descending slope and then selects the final set of points with the best regression adjusted R squared. None of the different results would be reported in a typical reporting workflow as all  $r^2$  values were  $< 0.7$ .

## Additional Figures and Tables

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<sup>4</sup><https://www.certara.com/software/phoenix-nlme/>

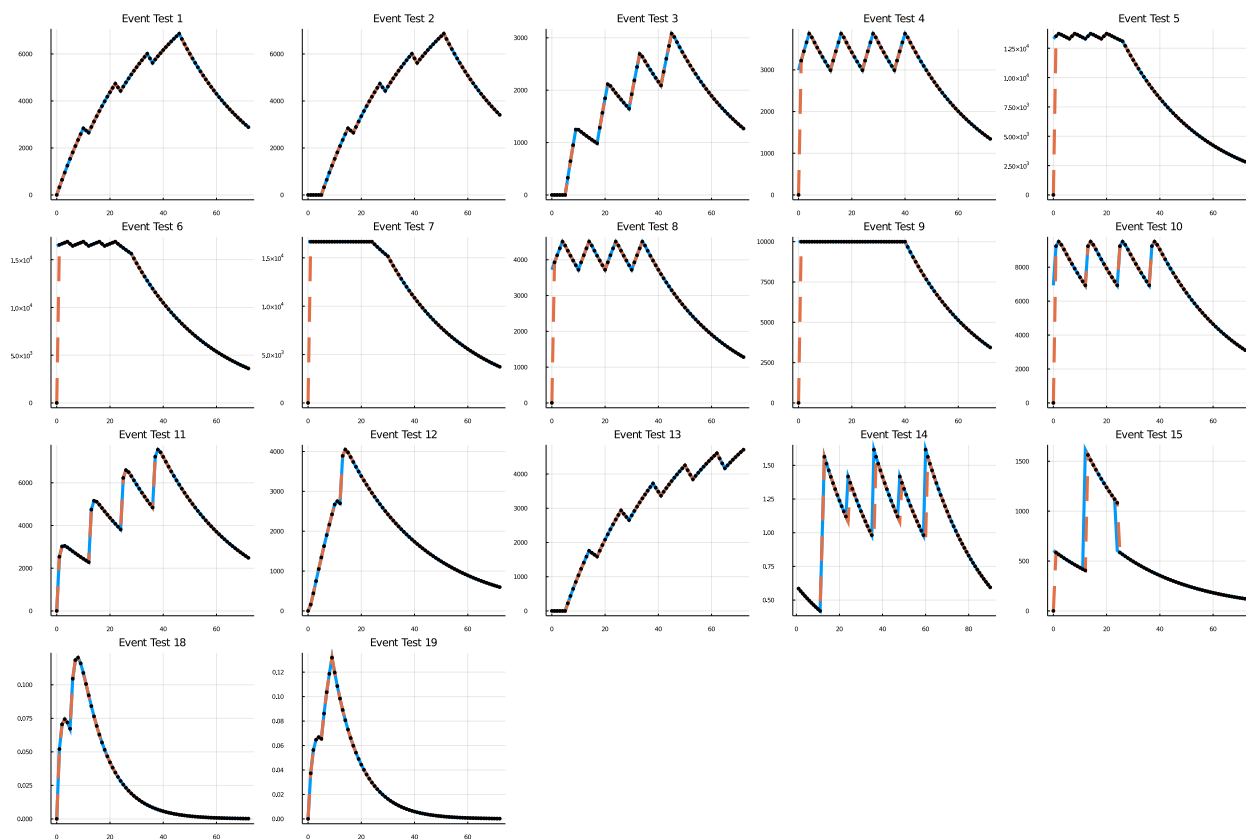


Figure 9: **Clinical dosing simulation verification against NONMEM for analytical solutions.** Shown are the clinical dosing simulation verification of the models described in Section 5 using analytical solutions of the differential equation mixed with the event handling system.



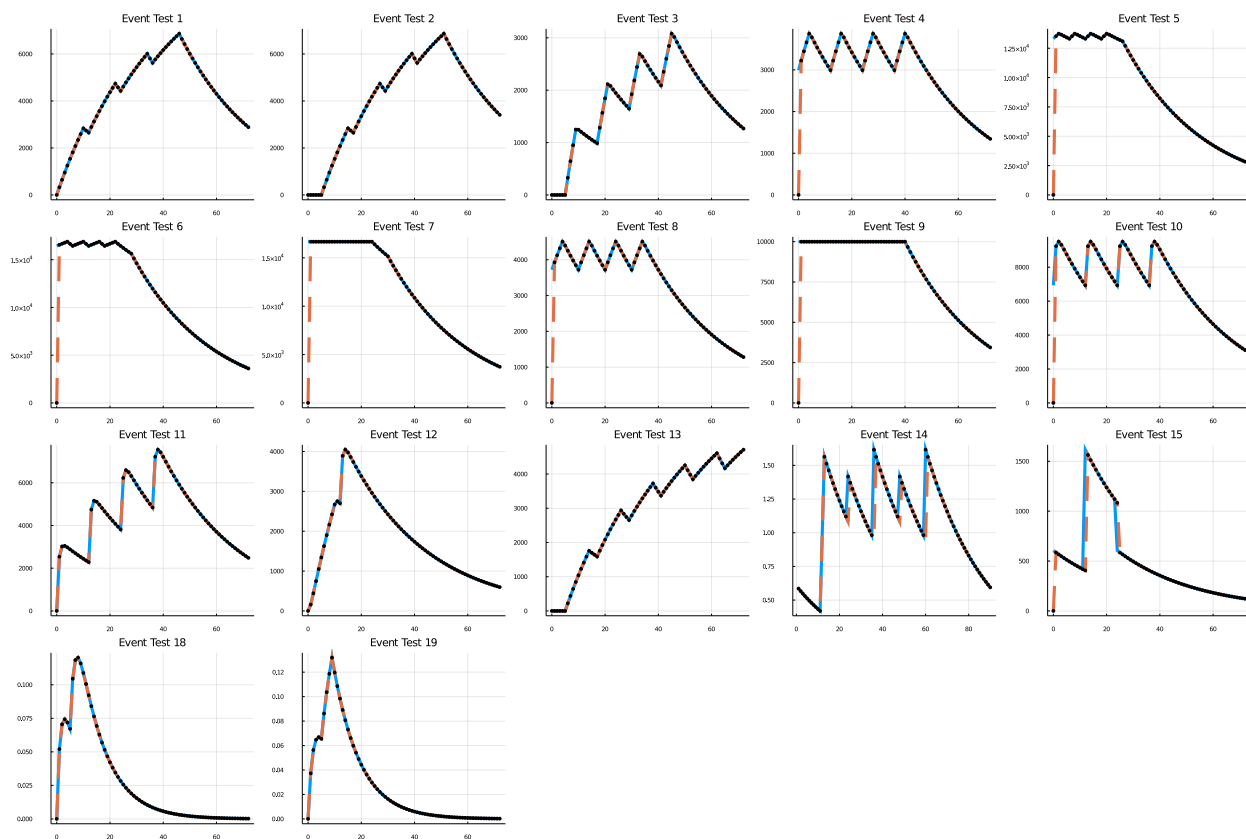


Figure 10: **Clinical dosing simulation verification against NONMEM for analytical solutions.** Shown are the clinical dosing simulation verification of the models in Section 5 using numerical approximations of the solutions of the differential equation mixed with the event handling system.

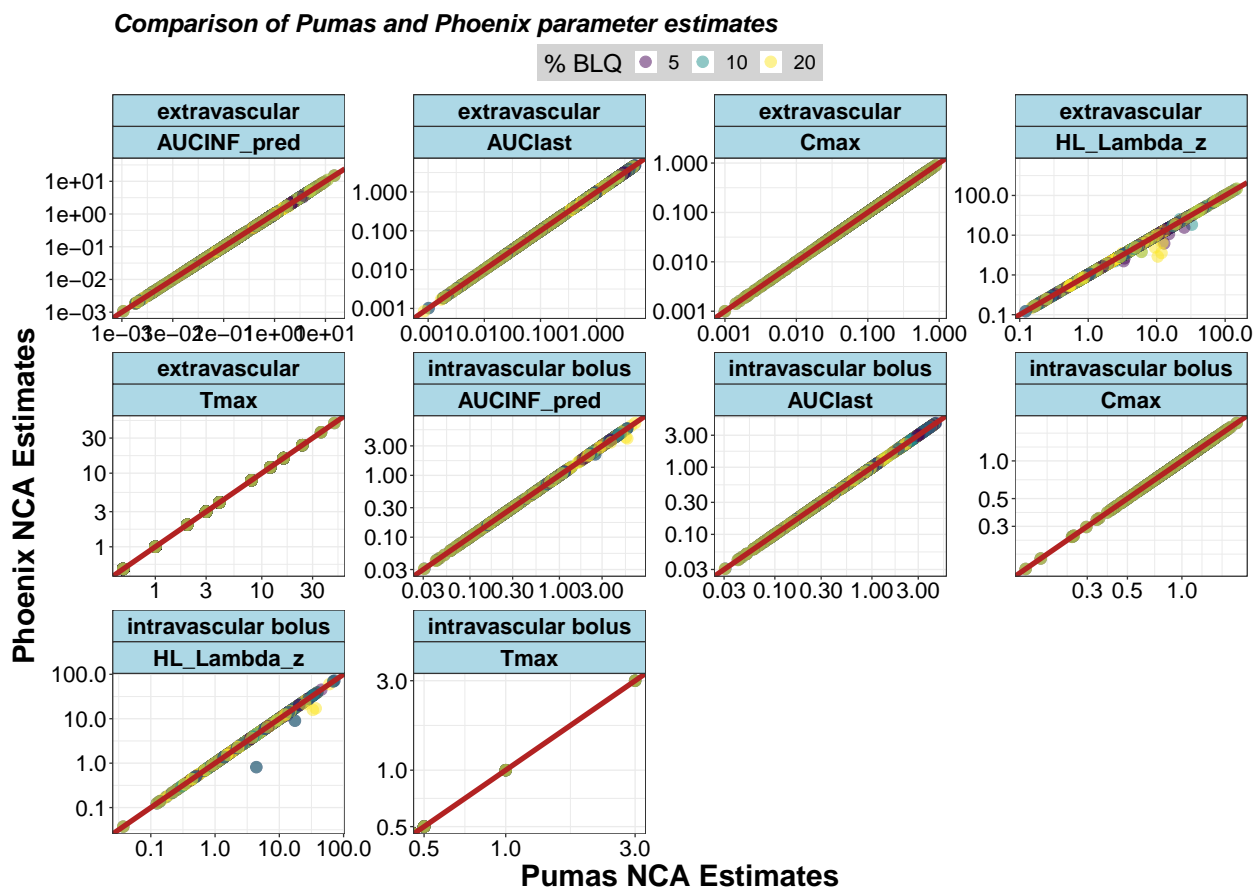


Figure 11: **Pumas and Phoenix NCA Results Comparison.** Eighteen percent of subjects were randomly selected from the 13104 scenarios to form a subset of 2367 subjects. This smaller subset was used to perform the NCA calculations across the three software while ensuring to maintain the same default options. All analysis were conducted on the same laptop.

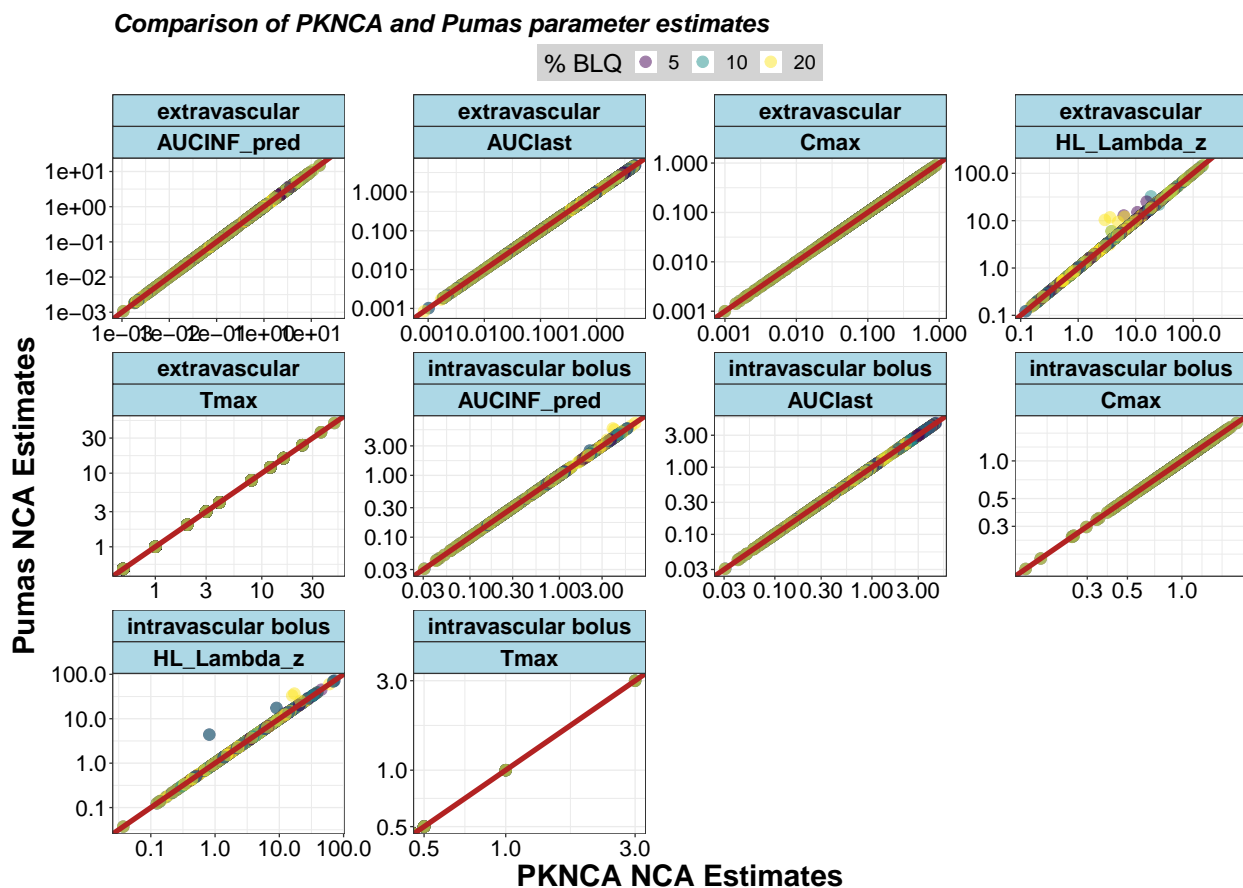


Figure 12: **Pumas and PKNCA Results Comparison.** Eighteen percent of subjects were randomly selected from the 13104 scenarios to form a subset of 2367 subjects. This smaller subset was used to perform the NCA calculations across the three software while ensuring to maintain the same default options. All analysis were conducted on the same laptop.