

# User Manual

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An R package for parametric  
and non-parametric modeling  
and simulation of  
pharmacokinetic and  
pharmacodynamic systems



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# Introduction

Thank you for your interest in Pmetrics! This guide provides instructions and examples to assist users of the Pmetrics R package, by the Laboratory of Applied Pharmacokinetics at the University of Southern California. Please see our website at <http://www.lapk.org> for more information.

Here are some tips for using this guide.

- Items that are hyperlinked can be selected to jump rapidly to relevant sections.
- At the bottom of every page, the text “User’s Guide” can be selected to jump immediately to the table of contents.
- Items in `courier` font correspond to R commands

## Citing Pmetrics

Please help us maintain our funding to provide Pmetrics as a free research tool to the pharmacometric community. If you use Pmetrics in a publication, you can cite it as below. In R you can always type `citation("Pmetrics")` to get this same reference.

**Neely MN, van Guilder MG, Yamada WM, Schumitzky A, Jelliffe RW. Accurate Detection of Outliers and Subpopulations With Pmetrics, a Nonparametric and Parametric Pharmacometric Modeling and Simulation Package for R. Ther Drug Monit 2012; 34:467–476.**

## Disclaimer

You, the user, assume all responsibility for acting on the results obtained from Pmetrics. The USC Laboratory of Applied Pharmacokinetics, members and consultants to the Laboratory of Applied Pharmacokinetics, and the University of Southern California and its employees assume no liability whatsoever. Your use of the package constitutes your agreement to this provision.

## System Requirements and Installation

Pmetrics and all required components will run under Mac (Unix), Windows, and Linux. There are three required software components which must be installed on your system **in this order**:

1. The statistical programming language and environment “**R**”
2. The **Pmetrics** package for R
3. **gfortran** or some other Fortran compiler

A fourth, highly recommended, but optional component is **Rstudio**, a user-friendly wrapper for R. It can be installed any time after installing R (i.e. step 1 above).

All components have versions for Mac, Windows, and Linux environments, and 32- and 64- bit processors. All are free of charge.

### R

R is a free software environment for statistical computing and graphics, which can be obtained from <http://www.R-project.org/>. Pmetrics is a library for R.

### Pmetrics

If you are reading this manual, then you have likely visited our website at <http://www.lapk.org>, where you can select the software tab and download our products, including Pmetrics.

Pmetrics is distributed as a package source file archive (.tgz for Mac, .zip for Windows, .tar.gz for Linux). **Do not open the archive.** To install Pmetrics from the R console, use the command `install.packages(file.choose())`, and navigate when prompted to the folder in which you placed the Pmetrics package archive (.zip or .tgz) file. Pmetrics will need the several additional R packages for some functions. However, you do not have to install these if you do not already have them in your R library. They should automatically be downloaded and installed when you use `PMbuild()` or the first time you use a Pmetrics function that requires them.

## Fortran

In order to run Pmetrics, a Fortran compiler is required. After you have installed Pmetrics, the first command to run is `PMbuild()` which will verify an installed fortran compiler on your system. You can get detailed instructions on how to obtain and install gfortran appropriate for your system at this page on our website: [http://www.lapk.org/Pmetrics\\_install.php](http://www.lapk.org/Pmetrics_install.php).

## Rstudio

A text editor that can link to R is useful for saving scripts. Both the Windows and Mac versions of R have rudimentary text editors that are stable and reliable. Numerous other free and paid editors can also do the job, and these can be located by searching the internet. We prefer [Rstudio](#), which we strongly recommend.

## What This Manual Is Not

We assume that the user has familiarity with population modeling and R, and thus this manual is not a tutorial for basic concepts and techniques in either domain. We have tried to make the R code simple, regular and well documented. A very good free online resource for learning the basics of R can be found at <http://www.statmethods.net/index.html>. We recognize that initial use of a new software package can be complex, so please feel free to contact us at any time, preferably through the Pmetrics forum at <http://www.lapk.org> or directly by email at [contact@lapk.org](mailto:contact@lapk.org).

This manual is also not intended to be a theoretical treatise on the algorithms used in IT2B or NPAG. For that the user is directed to our website at [www.lapk.org](http://www.lapk.org).

## Getting Help and Updates

There is an active LAPK forum available from our website at <http://www.lapk.org> with all kinds of useful tips and help with Pmetrics. Register (separately from your LAPK registration) and feel free to post! Within R, you can also use `help("command")` or `?command` in the R console to see detailed help files for any Pmetrics command. Many commands have examples included in this documentation and you can execute the examples with `example(command)`. Note that, here, quotation marks are unnecessary around `command`. You can also type `PMmanual()` to launch this manual from within Pmetrics as well as a catalogue of all Pmetrics functions. Finally, `PMnews()` will display the Pmetrics change log.

Pmetrics will check for updates automatically every time you load it with `library(Pmetrics)`. If an update is available, it will provide a brief message to inform you. You can then use `PMupdate()` to update Pmetrics from within R, without having to visit our website. You will be prompted for your LAPK user email address and password. When bugs arise in Pmetrics, you may see a start up message to inform you of the bug and a patch can be installed by the command `PMpatch()` if available. Note that patches must be reinstalled with this command every time you launch Pmetrics, until the bug is corrected in the next version.

As of version 1.0.0 Pmetrics has graphical user interface (GUI) capability for many functions. Using `PMcode("function")` will launch the GUI in your default browser. While you are interacting with the GUI, R is "listening" and no other activity is possible. The GUI is designed to generate Pmetrics R code in response to your input in a friendly, intuitive environment. That code can be copied and pasted into your Pmetrics R script. You can also see live plot previews with the GUI. All this is made possible with the 'shiny' package for R.

Currently, the following GUIs are available: `PMcode("run")` and `PMcode("plot")`. More are coming!

# Pmetrics Components

There are three main software programs that Pmetrics controls.

- IT2B is the ITerative 2-stage Bayesian parametric population PK modeling program. It is generally used to estimate parameter ranges to pass to NPAG. It will estimate values for population model parameters under the assumption that the underlying distributions of those values are normal or transformed to normal.
- NPAG is the Non-parametric Adaptive Grid software. It will create a non-parametric population model consisting of discrete support points, each with a set of estimates for all parameters in the model plus an associated probability (weight) of that set of estimates. There can be at most one point for each subject in the study population. There is no need for any assumption about the underlying distribution of model parameter values.
- The simulator is a semi-parametric Monte Carlo simulation software program that can use the output of IT2B or NPAG to build randomly generated response profiles (e.g. time-concentration curves) for a given population model, parameter estimates, and data input. Simulation from a non-parametric joint density model, i.e. NPAG output, is possible, with each point serving as the mean of a multivariate normal distribution, weighted according to the weight of the point. The covariance matrix of the entire set of support points is divided equally among the points for the purposes of simulation.

Pmetrics has groups of R functions named logically to run each of these programs and to extract the output. Again, these are extensively documented within R by using the `help` (command) or `?command` syntax.

- `ITrun`, `ITparse`, `ERRrun`
- `NPrun`, `NPparse`
- `PMload`, `PMsave`, `PMreport`
- `SIMrun`, `SIMparse`

For IT2B and NPAG, the “run” functions generate batch files, which when executed, launch the software programs to do the analysis. `ERRrun` is a special implementation of IT2B designed to estimate the assay error polynomial coefficients from the data, when they cannot be calculated from assay validation data (using `makeErrorPoly()`) supplied by the analytical laboratory. The batch files contain all the information necessary to complete a run, tidy the output into a date/time stamped directory with meaningful subdirectories, extract the information, generate a report, and a saved Rdata file of parsed output which can be quickly and easily loaded into R. On Mac (Unix) systems, the batch file will automatically launch in a Terminal window. On Windows systems, the batch file must be launched manually. In both cases, the execution of the program to do the actual model parameter estimation is independent of R, so that the user is free to use R for other purposes.

For the Simulator, the “`SIMrun`” function will execute the program directly within R.

For all programs, the “parse” functions will extract the primary output from the program into meaningful R data objects. For IT2B and NPAG, this is done automatically at the end of a successful run, and the objects are saved in the output subdirectory as `IT2Bout.Rdata` or `NPAGout.Rdata`, respectively.

For IT2B and NPAG, the “`PMload`” function can be used to load the either of the above `.Rdata` files after a successful run. “`PMsave`” is the companion to `PMload` and can re-save modified objects to the `.Rdata` file.

The “`PMreport`” function is automatically run at the end of a successful NPAG and IT2B run, and it will generate an HTML page with summaries of the run, as well as the `.Rdata` files and other objects. The default browser will be automatically launched for viewing of the HTML report page. It will also generate a `.tex` file suitable for processing by a `LATEX` engine to generate a pdf report. See the [Pmetrics Outputs](#) section.

Within Pmetrics there are also functions to manipulate data `.csv` files and process and plot extracted data.

- Manipulate data `.csv` files: `PMreadMatrix`, `PMcheck`, `PMwriteMatrix`, `PMmatrixRelTime`, `PMwrk2csv`, `NM2PM`
- Process data: `makeAUC`, `makeCov`, `makeCycle`, `makeFinal`, `makeOP`, `makeNCA`, `makePTA`, `makeErrorPoly`
- Plot data: `plot.PMcov`, `plot.PMcycle`, `plot.PMfinal`, `plot.PMmatrix`, `plot.PMop`, `plot.PMsim`, `plot.PMdiag`, `plot.PMpta`
- Model selection and diagnostics: `PMcompare`, `plot.PMop` (with residual option), `makeNPDE`, `PMstep`

- Pmetrics function defaults: PMwriteDefaults

Again, all functions have extensive help files and examples which can be examined in R by using the `help(command)` or `?command` syntax.

## Customizing Pmetrics Options

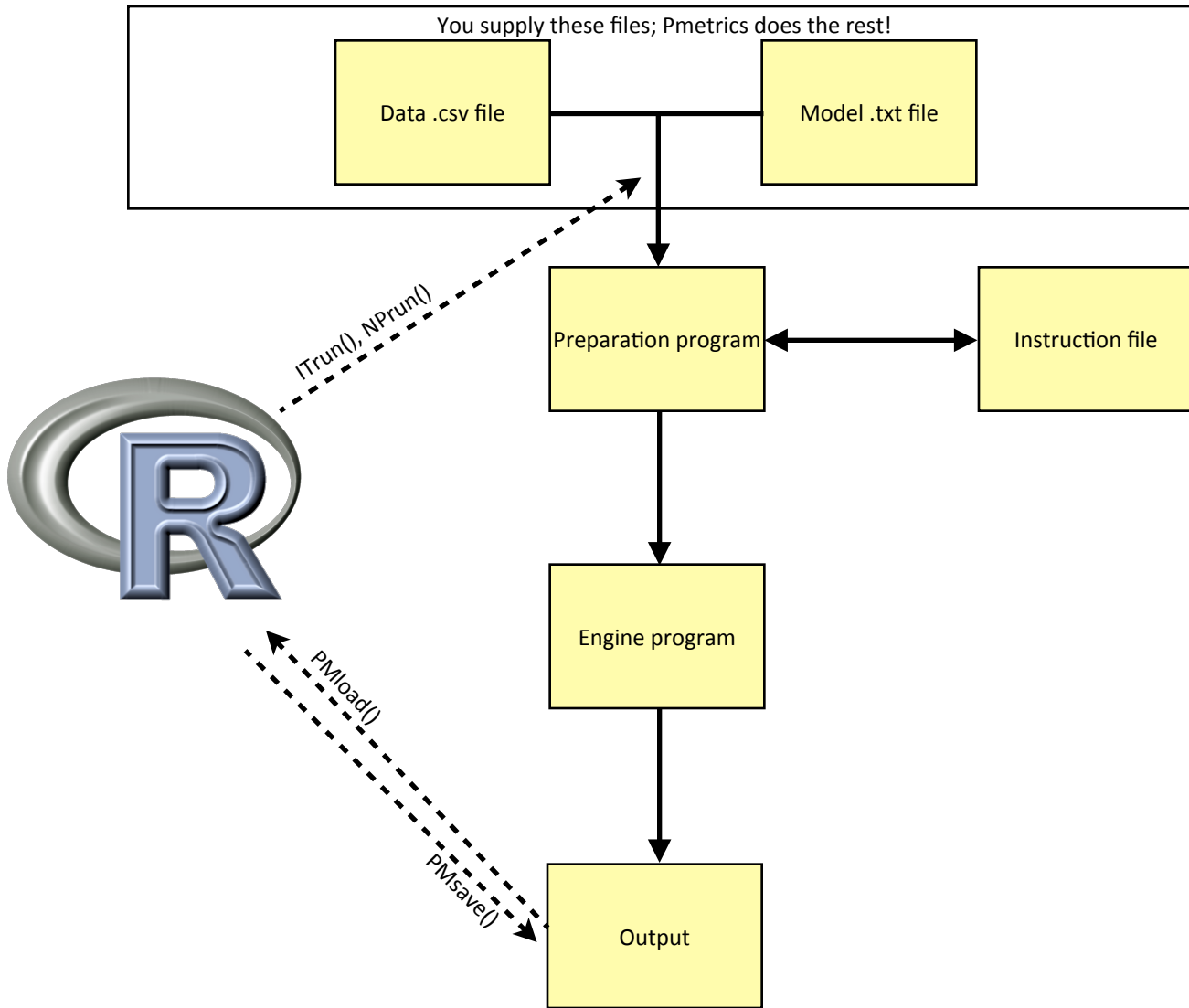
You can change global options in Pmetrics

`setPMoptions(sep, dec)` Currently you can change two options. `sep` will allow Pmetrics to read data files whose field separators are semicolons and decimal separators are commas, e.g. `setPMoptions(sep="; ", dec=", ")`. These options will persist from session to session until changed.

`getPMoptions()` will return the current options.

## General Workflow

The general Pmetrics workflow for IT2B and NPAG is shown in the following diagram.



R is used to specify the working directory containing the data .csv and model .txt files. Through the batch file generated by R, the preparation program is compiled and executed. The instruction file is generated automatically

by the contents of the data and model files, and by arguments to the `NPrun()`, `ITrun()` or `ERRrun()` commands. The batch file will then compile and execute the engine file according to the instructions, which will generate several output files upon completion. Finally, the batch file will call the R script to generate the summary report and several data objects, including the `IT2Bout.Rdata` or `NPAGout.Rdata` files which can be loaded into R subsequently using `PMload()`. Objects that are modified can be saved back to the `.Rdata` files with `PMsave()`.

Both input files (data, model) are text files which can be edited directly.

## Pmetrics Input Files

### Data .csv Files

Pmetrics accepts input as a spreadsheet “matrix” format. It is designed for input of multiple records in a concise way. **Please keep the number of characters in the file name  $\leq 8$ .**

Files are in comma-separated-values (.csv) format. Examples of programs that can save .csv files are any text editor (e.g.TextEdit on Mac, Notepad on Windows) or spreadsheet program (e.g. Excel). Click on hyperlinked items to see an explanation.

**IMPORTANT:** The order, capitalization and names of the header and the first 12 columns are fixed. All entries must be numeric, with the exception of ID and “.” for non-required placeholder entries.

#### PODATA DEC 11

#ID	EVID	TIME	DUR	DOSE	ADDL	II	INPUT	OUT	OUTEQ	C0	C1	C2	C3	COV...
GH	1	0	0	400	.	.	1	.	.	.	.	.	.	.
GH	0	0.5	.	.	.	.	.	0.42	1	0.01	0.1	0	0	
GH	0	1	.	.	.	.	.	0.46	1	0.01	0.1	0	0	
GH	0	2	.	.	.	.	.	2.47	1	0.01	0.1	0	0	
GH	4	0	0	150	.	.	1	.	.	.	.	.	.	
GH	1	3.5	0.5	150	.	.	1	.	.	0.01	0.1	0	0	
GH	0	5.12	.	.	.	.	.	0.55	1	0.01	0.1	0	0	
GH	0	24	.	.	.	.	.	0.52	1	0.01	0.1	0	0	
1423	1	0	1	400	-1	12	1	.	.	.	.	.	.	
1423	1	0.1	0	100	.	.	2	.	.	.	.	.	.	
1423	0	1	.	.	.	.	.	-99	1	0.01	0.1	0	0	
1423	0	2	.	.	.	.	.	0.38	1	0.01	0.1	0	0	
1423	0	2	.	.	.	.	.	1.6	2	0.05	0.2	-0.11	0.002	

**PODATA DEC\_11** This is the fixed header for the file and must be in the first line. It identifies the version. It is not the date of your data file.

**#ID** This field must be preceded by the “#” symbol to confirm that this is the header row. It can be numeric or character and identifies each individual. All rows must contain an ID, and all records from one individual must be contiguous. Any subsequent row that begins with “#” will be ignored, which is helpful if you want to exclude data from the analysis, but preserve the integrity of the original dataset, or to add comment lines. IDs should be 11 characters or less but may be any alphanumeric combination. **There can be at most 800 subjects per run.**

**EVID** This is the event ID field. It can be 0, 1, or 4. Every row must have an entry.

0 = observation

1 = input (e.g. dose)

2, 3 are currently unused

4 = reset, where all compartment values are set to 0 and the time counter is reset to 0. This is useful when an individual has multiple sampling episodes that are widely spaced in time with no new information gathered. This is a dose event, so dose information needs to be complete.

<b>TIME</b>	This is the elapsed time in decimal hours since the first event. It is not currently clock time (e.g. 21:30), although this is planned. Every row must have an entry, and within a given ID, rows must be sorted chronologically, earliest to latest.
<b>DUR</b>	This is the duration of an infusion in hours. If EVID=1, there must be an entry, otherwise it is ignored. For a bolus (i.e. an oral dose), set the value equal to 0.
<b>DOSE</b>	This is the dose amount. If EVID=1, there must be an entry, otherwise it is ignored.
<b>ADDL</b>	This specifies the number of additional doses to give at interval II. It may be missing for dose events (EVID=1 or 4), in which case it is assumed to be 0. It is ignored for observation (EVID=0) events. Be sure to adjust the time entry for the subsequent row, if necessary, to account for the extra doses. If set to -1, the dose is assumed to be given under steady-state conditions. ADDL=-1 can only be used for the first dose event for a given subject, or an EVID=4 event, as you cannot suddenly be at steady state in the middle of dosing record, unless all compartments/times are reset to 0 (as for an EVID=4 event).
<b>II</b>	This is the interdose interval and is only relevant if ADDL is not equal to 0, in which case it cannot be missing. If ADDL=0 or is missing, II is ignored.
<b>INPUT</b>	This defines which input (i.e. drug) the DOSE corresponds to. Inputs are defined in the model file.
<b>OUT</b>	This is the observation, or output value. If EVID=0, there must be an entry; if missing, this must be coded as -99. It will be ignored for any other EVID and therefore can be ". ". There can be at most 150 observations for a given subject.
<b>OUTEQ</b>	This is the output equation number that corresponds to the OUT value. Output equations are defined in the model file.
<b>C0, C1, C2, C3</b>	These are the coefficients for the assay error polynomial for that observation. Each subject may have up to one set of coefficients per output equation. If more than one set is detected for a given subject and output equation, the last set will be used. If there are no available coefficients, these cells may be left blank or filled with " ." as a placeholder.
<b>COV...</b>	Any column after the assay error coefficients is assumed to be a covariate, one column per covariate.

## Manipulation of CSV files

There are several functions in Pmetrics which are useful for either converting other formats into Pmetrics data files, or checking Pmetrics data files for errors and fixing some of them automatically.

`PMreadMatrix(filename, ...)` This function simply reads *filename* and creates a PMmatrix object in memory which can be plotted (see `?plot.PMmatrix`) or otherwise analyzed.

`PMcheck(PMmatrix|filename, model, ...)` This function will check a .csv file named *filename* or a PMmatrix data frame containing a previously loaded .csv file (the output of `PMreadMatrix`) for errors which would cause the analysis to fail. If a model file is provided, and the data file has no errors, it will also check the model file for errors. If it finds errors, it will generate a new errors.xlsx file with all errors highlighted and commented so that you can find and correct them easily. See `?PMcheck` for details in R.

`PMwriteMatrix(data.frame, filename, ...)` This function writes an appropriate data.frame as a new .csv file. It will first check the data.frame for errors via the `PMcheck()` function above, and writing will fail if errors are detected. This can be overridden with `override=T`.

`PMmatrixRelTime()` This function converts dates and clock times of specified formats into relative times for use in the NPAG, IT2B and Simulator engines. See `?PMmatrixRelTime` for details.

`PMwrk2csv()` This function will convert old-style, single-drug USC\*PACK .wrk formatted files into Pmetrics data .csv files. Details are available with `?PMwrk2csv` in R.

`PMmb2csv()` This function will convert USC\*PACK .mb files into Pmetrics data .csv files. Details are available with `?PMmb2csv` in R.

`NM2PM()` Although the structure of Pmetrics data files are similar to NONMEM, there are some differences. This function attempts to automatically convert to Pmetrics format. It has been tested on several examples, but there are probably NONMEM files which will cause it to crash. Running `PMcheck()` afterwards is a good idea. Details can be found with `?NM2PM` in R.

## Model Files

Model files for Pmetrics are ultimately Fortran text files with a header version of TSMULT... As of Pmetrics version 0.30, we have adopted a very simple user format that Pmetrics will use to generate the Fortran code automatically for you. Version 0.4 additionally eliminates the previously separate instruction file. A model library is available on our website at <http://www.lapk.org/pmetrics.php>.

**Naming your model files.** The default model file name is "model.txt," but you can call them whatever you wish. However, **please keep the number of characters in the model file name ≤ 8**. When you use a model file in `NPrun()`, `ITrun()`, `ERRrun()`, or `SIMrun()`, Pmetrics will make a Fortran model file of the same name, temporarily renaming your file. At the end of the run, your original model file will be in the /inputs subfolder of the run folder, and the generated Fortran model file will be called "model.for" and moved to the /etc subfolder of the run folder. If your model is called "mymodel.txt", then the Fortran file will be "mymodel.for".

You can still use appropriate Fortran model files directly, but we suggest you keep the .for extension for all Fortran files to avoid confusion with the new format. If you use a .for file as your model, you will have to specify its name explicitly in the `NPrun()`, `ITrun`, `ERRrun()`, or `SIMrun()` command, since the default model name again is "model.txt." If you use a .for file directly, it will be in the /inputs subfolder of the run folder, not in /etc, since you did not use the simpler template as your model file.

**Structure of model files.** The new model file is a text file with 11 blocks, each marked by "#" followed by a header tag.

[#PRImary variables](#)

[#COVariates](#)

[#SECcondary variables](#)

[#BOLus inputs](#)

[#INITial conditions](#)

[#F \(bioavailability\)](#)

[#LAG time](#)

[#DIFFerential equations](#)

[#OUTputs](#)

[#ERRor](#)

[#EXTra](#)

For each header, only the capital letters are required for recognition by Pmetrics. The blocks can be in any order, and header names are case-insensitive (i.e. the capitalization here is just to show which letters are required). Fortran is also case-insensitive, so in variable names and expressions case is ignored. Details of each block are next, followed by a [complete example](#).

**Important:** Sometimes it is important to preserve spacing and formatting in Fortran code that you might insert into blocks, particularly the #EXTRA block. If you wish to do this, insert [format] and [/format] before and after any code that you wish to reproduce verbatim with spacing in the fortran model file.

**Comments:** You can insert comments into your model text file by starting a line with a capital "C" followed by a space. These lines will be removed/ignored in the final fortran code.

## Primary variables

Primary variables are the model parameters that are to be estimated by Pmetrics or are designated as fixed parameters with user specified values. It should be a list of variable names, one name to a line. Variable names should be 11 characters or fewer. Some variable names are [reserved](#) for use by Pmetrics and cannot be used as primary variable names. **The number of primary variables must be between 2 and 32, with at most 30 random or 20 fixed.**

On each row, following the variable name, include the range for the parameter that defines the search space. These ranges behave slightly differently for NPAG, IT2B, and the simulator.

- For all engines, the format of the limits is *min, max*. A single value will indicate that the parameter is to be fixed but unknown in the population, i.e. the value is taken as the starting point for the optimization, but the final value will depend on the model and data and will be the same across the population. A single value followed by an “!” will indicate that this value is to be held constant (i.e. fixed and known) across the population, and not to be estimated.
- For **NPAG**, when *min/max* limits are specified, they are absolute, i.e. the algorithm will not search outside this range.
- For **IT2B**, the range defines the Bayesian prior distribution of the parameter values for cycle 1. For each parameter, the mean of the Bayesian prior distribution is taken as the middle of the range, and the standard deviation is *xsig\*range* (see [IT2B runs](#)). Adding a plus sign (+) to a line will prevent that parameter from being assigned negative values. NPAG and the simulator will ignore the pluses as the ranges are absolute for these engines. Note that prior to version 1.5.0, this used to be an exclamation point (!) but to be consistent throughout the model file, the exclamation point is now used when fixed values are desired.
- The **simulator** will ignore the ranges with the default value of NULL for the *limits* argument. If the simulator *limits* argument is set to NA, which will mean that these ranges will be used as the limits to truncate the simulation (see [Simulator Runs](#)).

Example:

```
#Pri
KE, +0, 5
V, 0.01, 100
KA, 0, 5
KCP, 5
KPC, 0 , 5
Tlag1, 0, 2
IC3, 0, 10000
FA1, 0.5!
```

In this example, KE has a range of 0 to 5, which will be absolute for NPAG and the simulator (if limits=NA), but defines the prior distribution for KE if using IT2B. The “+” limits KE to the positive real numbers for IT2B and is ignored in NPAG. KCP is set initially to 5 regardless of the engine, but will be estimated and the same in all subjects. FA1 is held constant (not estimated) at 0.5 for all subjects.

## Covariates

Covariates are subject specific data, such as body weight, contained in the data .csv file. The covariate names, which are the column names in the data file, can be included here for use in secondary variable equations. The order should be the same as in the data file and although the names do not have to be the same, we strongly encourage you to make them the same to avoid confusion.

Covariates are applied at each dose event. The first dose event for each subject must have a value for every covariate in the data file. By default, missing covariate values for subsequent dose events are linearly interpolated between existing values, or carried forward if the first value is the only non-missing entry. To suppress interpolation and carry forward the previous value in a piece-wise constant fashion, include an exclamation point (!) in any declaration line.

**Note** that any covariate relationship to any parameter may be described as the user wishes by mathematical equations and Fortran code, allowing for exploration of complex, non-linear, time-dependent, and/or conditional relationships.

Example:

```
#Cov  
wt  
cyp  
IC!
```

where IC will be piece-wise constant and the other two will be linearly interpolated for missing values.

### Secondary variables

Secondary variables are those that are defined by equations that are combinations of primary, covariates, and other secondary variables. If using other secondary variables, define them first within this block. Equation syntax must be Fortran. It is permissible to have conditional statements, but because expressions in this block are translated into variable declarations in Fortran, expressions other than of the form "X = function(Y)" must be prefixed by a "&" and contain only variables which have been previously defined in the Primary, Covariate, or Secondary blocks. Note that prior to version 1.5.0, the continuation symbol was "+" before each line, but to avoid confusion with the use of "+" in the Primary block for IT2B models, and to be consistent with Fortran notation, the "&" is used henceforth.

Example:

```
#Sec  
CL = Ke * V * wt**0.75  
& IF(cyp .GT. 1) CL = CL * cyp
```

### Bolus inputs

By default, inputs with DUR (duration) of 0 in the data .csv file are "delivered" instantaneously to the model compartment equal to the input number, i.e. input 1 goes to compartment 1, input 2 goes to compartment 2, etc. This can be overridden with NBOLUS(input number) = compartment number.

Example:

```
#Bol  
NBCOMP(1) = 2
```

### Initial conditions

By default, all model compartments have zero amounts at time 0. This can be changed by specifying the compartment amount as X(.) = expression, where "." is the compartment number. Primary and secondary variables and covariates may be used in the expression, as can conditional statements in Fortran code. An "&" continuation prefix is not necessary in this block for any statement, although if present, will be ignored.

Example:

```
#Ini  
X(2) = IC*V (i.e. IC is a covariate with the measured trough concentration prior to an observed dose)  
X(3) = IC3 (i.e. IC3 is a fitted amount in this unobserved compartment)
```

In the first case, the initial condition for compartment 2 becomes the value of the IC covariate (defined in #Covariate block) multiplied by the current estimate of V during each iteration. This is useful when a subject has been taking a drug as an outpatient, and comes in to the lab for PK sampling, with measurement of a concentration immediately prior to a witnessed dose, which is in turn followed by more sampling. In this case, IC or any other covariate can be set to the initial measured concentration, and if V is the volume of compartment 2, the initial condition (amount) in compartment 2 will now be set to the measured concentration of drug multiplied by the estimated volume for each iteration until convergence.

In the second case, the initial condition for compartment 3 becomes another variable, IC3 defined in the #Primary block, to fit in the model, given the observed data.

## F (bioavailability)

Specify the bioavailability term, if present. Use the form FA(.) = expression, where "." is the input number. Primary and secondary variables and covariates may be used in the expression, as can conditional statements in Fortran code. An "&" continuation prefix is not necessary in this block for any statement, although if present, will be ignored.

Example:

```
#F  
FA(1) = FA1
```

## Lag time

Specify the lag term, if present, which is the delay after an absorbed dose before observed concentrations. Use the form TLAG(.) = expression, where "." is the input number. Primary and secondary variables and covariates may be used in the expression, as can conditional statements in Fortran code. An "&" continuation prefix is not necessary in this block for any statement, although if present, will be ignored.

Example:

```
#Lag  
TLAG(1) = Tlag1
```

## Differential equations

Specify a model in terms of ordinary differential equations, in Fortran format. XP(.) is the notation for  $dX(.)/dt$ , where "." is the compartment number. X(.) is the amount in the compartment. **There can be a maximum of 20 such equations.**

Example:

```
#Dif  
XP(1) = -KA*X(1)  
XP(2) = RATEIV(1) + KA*X(1) - (KE+KCP)*X(2) + KPC*X(3)  
XP(3) = KCP*X(2) - KPC*X(3)
```

RATEIV(1) is the notation to indicate an infusion of input 1 (typically drug 1). The duration of the infusion and total dose is defined in the [data.csv](#) file. **Up to 7 inputs are currently allowed.** These can be used in the model file as RATEIV(1), RATEIV(2), etc. The compartments for receiving the inputs of oral (bolus) doses are defined in the #Bolus block.

## Outputs

Output equations, in Fortran format. Outputs are of the form Y(.) = expression, where "." is the output equation number. Primary and secondary variables and covariates may be used in the expression, as can conditional statements in Fortran code. An "&" continuation prefix is not necessary in this block for any statement, although if present, will be ignored. **There can be a maximum of 6 outputs.** They are referred to as Y(1), Y(2), etc. These equations may also define a model explicitly as a function of primary and secondary variables and covariates.

Examples:

```
#Out  
Y(1) = X(2)/V
```

#OUT

```
RES = B(1) * KIN/(KIN-KOUT) * (EXP(-KOUT*TPD)-EXP(-KIN*TPD))
```

$Y(1) = RES/VD$

This second example is known as the Bateman equation for a model with linear absorption (KIN) into and elimination (KOUT) from a central compartment, and a time post-dose (TPD) or lag time. Here B(1) is the oral bolus dosing vector for drug 1, and VD is the volume of the central compartment.

### Error

This block contains all the information Pmetrics requires for the structure of the error model. In Pmetrics, each observation is weighted by  $1/\text{error}^2$ . There are two choices for the error term:

1. error = SD \* gamma
2. error =  $(SD^2 + \lambda^2)^{0.5}$  (Note that lambda is only available in NPAG currently).

where SD is the standard deviation (SD) of each observation [obs], and gamma and lambda are terms to capture extra process noise related to the observation, including mis-specified dosing and observation times.

SD is modeled by a polynomial equation with up to four terms:  $C_0 + C_1[\text{obs}] + C_2[\text{obs}]^2 + C_3[\text{obs}]^3$ . The values for the coefficients should ideally come from the analytic lab in the form of inter-run standard deviations or coefficients of variation at standard concentrations. You can use the Pmetrics function `makeErrorPoly()` to choose the best set of coefficients that fit the data from the laboratory. Alternatively, if you have no information about the assay, you can use the Pmetrics function `ERRrun()` to estimate the coefficients from the data. Finally, you can use a generic set of coefficients. We recommend that as a start,  $C_0$  be set to half of the lowest concentration in the dataset and  $C_1$  be set to 0.15.  $C_2$  and  $C_3$  can be 0.

In the multiplicative model, gamma is a scalar on SD. In general, well-designed and executed studies will have data with gamma values approaching 1. Poor quality, noisy data will result in gammas of 5 or more. Lambda is an additive model to capture process noise, rather than the multiplicative gamma model.

To specify the model in this block, the first line needs to be either `L=[number]` or `G=[number]` for a lambda or gamma error model. The [number] term is the starting value for lambda or gamma. Good starting values for lambda are 1 times  $C_0$  for good quality data, 3 times  $C_0$  for medium, and 5 or 10 times  $C_0$  for poor quality. Note, that  $C_0$  should generally not be 0, as it represents machine noise (e.g. HPLC or mass spectrometer) that is always present. For gamma, good starting values are 1 for high-quality data, 3 for medium, and 5 or 10 for poor quality. If you include an exclamation point (!) in the declaration, then lambda or gamma will be fixed and not estimated. Note that you can only fix lambda currently to zero.

The next line(s) contain the values for  $C_0$ ,  $C_1$ ,  $C_2$ , and  $C_3$ , separated by commas. There should be one line of coefficients for each output equation. By default Pmetrics will use values for these coefficients found in the data file. If none are present or if the model declaration line contains an exclamation point (!) the values here will be used.

Example 1: estimated lambda, starting at 0.4, one output, use data file coefficients but if missing, use 0.1,0.1,0,0

```
#Err  
L=0.4  
0.1,0.1,0,0
```

Example 2: fixed gamma of 2, two outputs, use data file coefficients but if missing, use 0.1,0.1,0,0 for the first output, but use 0.3, 0.1, 0, 0 for output 2 regardless of what is in the data file.

```
#Err  
G=2!  
0.1,0.1,0,0  
0.3,0.1,0,0!
```

## Extra

This block is for advanced Fortran programmers only. Occasionally, for very complex models, additional Fortran subroutines are required. They can be placed here. The code must specify complete Fortran subroutines which can be called from other blocks with appropriate call functions. As stated earlier, sometimes it is important to preserve spacing and formatting in Fortran code that you might insert into blocks, particularly the #EXTRA block. If you wish to do this, insert [format] and [/format] before and after any code that you wish to reproduce verbatim with spacing in the fortran model file.

## Reserved Names

The following cannot be used as primary, covariate, or secondary variable names. They can be used in equations, however.

Reserved Variable	Function in Pmetrics
ndim	internal
t	time
x	array of compartment amounts
xp	array of first derivative of compartment amounts
rpar	internal
ipar	internal
p	array of primary parameters
r	input rates
b	input boluses
npl	internal
numeqt	output equation number
ndrug	input number
nadd	covariate number
rateiv	intravenous input for inputs when DUR>0 in data files
cv	covariate values array
n	number of compartments
nd	internal
ni	internal
nup	internal
nuic	internal
np	number of primary parameters

nbcomp	bolus compartment array
psym	names of primary parameters
fa	bioavailability
tlag	lag time
tin	internal
tout	internal

## Complete Example

Here is a complete example of a model file, as of Pmetrics version 0.40 and higher:

```
#Pri
KE, 0, 5
V0, 0.1, 100
KA, 0, 5
Tlag1, 0, 3
#Cov
wt
C this weight is in kg

#Sec
V = V0*wt

#Lag
TLAG(1) = Tlag1

#Out
Y(1) = X(2)/V

#Err
L=0.4
0.1,0.1,0,0
```

### Notes:

By omitting a #Diffeq block with ODEs, Pmetrics understands that you are specifying the model to be solved algebraically. In this case, at least KE and V must be in the Primary or Secondary variables. KA, KCP, and KPC are optional and specify absorption, and transfer to and from the central to a peripheral compartment, respectively.

The comment line “C this weight is in kg” will be ignored.

## Brief Fortran Tutorial

Much more detailed help is available from <http://www.cs.mtu.edu/~shene/COURSES/cs201/NOTES/fortran.html>.

Arithmetic Operator	Meaning
+	addition
-	subtraction
*	multiplication
/	division
**	exponentiation

Relational Operator	Alternative Operator	Meaning
<	.LT.	less than
<=	.LE.	less than or equal
>	.GT.	greater than
>=	.GE.	greater than or equal
==	.EQ.	equal
/=	.NE.	not equal

Selective Execution	Example
IF (logical-expression) one-statement	IF (T >= 100) CL = 10
IF (logical-expression) THEN statements END IF	IF (T >= 100) THEN CL = 10 V = 10 END IF
IF (logical-expression) THEN statements-1 ELSE statements-2 END IF	IF (T >= 100) THEN CL = 10 ELSE CL = CL END IF

# How to use R and Pmetrics

## Setting up a Pmetrics project

When beginning a new modeling project, it is convenient to use the command `PMtree("project name")`. This command will set up a new directory in the current working directory named whatever you have included as the "project name". For example, a directory called "DrugX" will be created by `PMtree("DrugX")`. Beneath this directory, several subdirectories will be also created: **Rscript**, **Runs**, **Sim**, and **src**. The **Rscript** subdirectory will contain a skeleton R script to begin Pmetrics runs in the new project. The **Runs** subdirectory should contain all files required for a run (described next) and it will also contain the resulting numerically ordered run directories created after each Pmetrics NPAG or IT2B run. The **Sim** subdirectory can contain any files related to simulations, and the **src** subdirectory should contain original and manipulated source data files. Of course, you are free to edit this directory tree structure as you please, or make your own entirely.

## Getting the required files to run Pmetrics

When you wish to execute a Pmetrics run, you must ensure that appropriate Pmetrics model .txt and data .csv files are in the working directory, i.e. the Runs subdirectory of the project directory. R can be used to help prepare the data .csv file by importing and manipulating spreadsheets (e.g. `read.csv()`). The Pmetrics function `PMcheck()` can be used to check a .csv file or an R data frame that is to be saved as a Pmetrics data .csv file for errors. It can also check a model file for errors in the context of a datafile, e.g. covariates that do not match. `PMcheck(..., fix=T)` attempts to automatically rid data files of errors. The function `PMwriteMatrix()` can be used to write the R data object in the correct format for use by IT2B, NPAG, or the Simulator.

You can also download sample data and scripts from the [Pmetrics downloads](#) section of our website. Edit prior versions of model files to make new model files.

## Using scripts to control Pmetrics

As you will see in the skeleton R script made by `PMtree()` and placed in the Rscript subdirectory, if this is a first-time run, the R commands to run IT2B or NPAG are as follows. Recall that the "#" character is a comment character.

```
library(Pmetrics)
#Run 1 - add your run description here
setwd("working directory")
NPrun() #for NPAG or ITrun() for IT2B
```

The first line will load the Pmetrics library of functions. The second line sets the working directory to the specified path. The third line generates the batch file to run NPAG or IT2B and saves it to the working directory.

**NOTE:** On Mac systems, the batch file will be automatically launched in a Terminal window. On Windows systems, the batch file must be launched manually by double clicking the *npscript.bat* or *itscript.bat* file in the working directory.

`ITrun()` and `NPrun()` both return the full path of the output directory to the clipboard. By default, runs are placed in folders numbered sequentially, beginning with "1".

Now the output of IT2B or NPAG needs to be loaded into R, so the next command does this.

```
PMload(run_number)
```

Details of these commands and what is loaded are described in the R documentation (`?PMload`) and in the following section. The `run_number` should be included within the parentheses to be appended to the names of loaded R objects, allowing for comparison between runs, e.g. `PMload(1)`. Finally, at this point other Pmetrics commands can be added to the script to process the data, such as the following.

```

plot(final.1)
plot(cycle.1)
plot(op.1,type="pop") or plot(op.1$pop1)
plot(op.1) #default is to plot posterior predictions for output 1
plot(op.1,type="pop",resid=T)

```

Of course, the full power of R can be used in scripts to analyze data, but these simple statements serve as examples.

If you do not use the `PMtree()` structure, we suggest that the R script for a particular project be saved into a folder called “Rscript” or some other meaningful name in the working directory. Folders are not be moved by the batch file. Within the script, number runs sequentially and use comments liberally to distinguish runs, as shown below.

```

library(Pmetrics)

#Run 1 - Ka, Kel, V, all subjects
setwd("working directory")
NPrun() #assumes model="model.txt" and data="data.csv"
PMload(1)
...

```

Remember in R that the command `example(function)` will provide examples for the specified function. Most Pmetrics functions have examples.

### Using Shiny to control Pmetrics

Shiny is an R package made by the same group who produces Rstudio. While you are learning to use Pmetrics, you can use a graphical user interface that teaches you how to build code. Currently we have implemented Shiny interfaces for NPAG, IT2B and many plots in Pmetrics.

To use these Shiny interfaces, type `PMcode("x")`, where x is the function you wish. Currently you have two choices: “run” and “plot”.

`PMcode("run")` will launch the following dialog window in your default browser. As you make your selections, you will see R code generated which you can copy back to your script and execute. Simply close the browser window when you are finished.

The screenshot shows the NPAG Shiny interface with the 'Run' tab selected. The interface includes a dropdown menu for selecting a run type (NPAG or IT2B), tabs for Data, Model, and Run, and a code editor area. The code editor contains the following R code:

```

Data Model Run
template.csv
Covariate names: wt, age, ic2, ic3
Use Shift or CTRL (Windows) or CMD (Mac) + Click to (de)select subjects.
 Include subjects
 Exclude subjects
1 2 3
Number of drug inputs: 1
Salt fraction for each input, separated by commas:
1
AUC Interval in hours:
24

```

Copy and paste the code below into your R script to execute the run and get results back into R:  
Note: If you accepted the default value for an argument, it is not necessary to include that argument in the call to NPrun and it has been omitted here, following standard R practice.

```

NPrun(data="template.csv")
PMload(1)

```

**Current Model**

Primary Variables			Covariates		Secondary Variables	
name	min	max	name	type	VM = Vmax0 * wt**0.75	V = Vc0 * wt
Ka	0.00	15.00	wt	Interpolated		
Vmax0	0.00	60.00	age	Interpolated		
Km	0.00	60.00				
Vc0	0.00	5.00				
FA1	0.10	1.00				
KCP	0.00	15.00				
KPC	0.00	15.00				

Bolus Inputs		Initial Conditions		Lag Time	
None		None		None	

Differential Equations		Outputs		Error Model	
X(1)' = -Ka*X(1)		Y(1) = X(2)/V		declaration	type
XP(2) = Ka*X(1)-RATEIV(1)-VM/(Vm*V-X(2))*X(2)-KCP*X(2)-KPC*X & (3)				Lambda	Estimated, Initial=0.4
XP(3) = KCP*X(2) - KPC*X(3)				0.02,0.1,0,0	If not in datafile

You can select from an NPAG or IT2B run, and then use the Data, Model, and Run tabs to navigate to different options. With a valid Pmetrics model and data file loaded, you will see a summary of the model in the Current Model window.

If you use PMcode("plot") you will see the dialog window below. It will allow you to select various Pmetrics objects that have been already loaded with PMload(), and offer you many plot options, as well as the code to generate you selected plot.

## Pmetrics Data Objects

After a successful IT2B or NPAG run, an R datafile is saved in the output subdirectory of the newly created numerically ordered folder in the working directory. After IT2B, this file is called "IT2Bout.Rdata", and after NPAG it is called "NPAGout.Rdata". As mentioned in the previous section, these data files can be loaded by ensuring that the Runs folder is set as the working directory, and then using the Pmetrics commands PMload (run\_num).

There are several Pmetrics data objects contained within the Rdata files which are loaded with PMload(), making these objects available for plotting and other analysis.

### Objects loaded by PMload (run\_num)

Objects	Variables	Comments
op (class: PMop, list)	\$id	Subject identification
	\$time	Observation time in relative decimal hours
	\$obs	Observation
	\$pred	Predicted value
	\$pred.type	Type of prediction, i.e. based on the population parameter values or Bayesian posterior parameter values
	\$icen	Median (default) or mean of the parameter distributions used to calculate the predicted values.
	\$outeq	
	\$block	Dosing block, usually 1 unless data file contains EVID=4 dose reset events, in which case each such reset within a given ID will increment the dosing block by 1 for that ID
	\$obsSD	Calculated standard deviation (error) of the observation based on the assay error polynomial
	\$d	Difference between pred and obs
	\$ds	Squared difference between pred and obs
	\$wd	\$d, weighted by the \$obsSD
	\$wds	\$ds, weighted by the \$obsSD

Objects	Variables	Comments
final (class: PMfinal, list)	\$popPoints	(NPAG only) Data.frame of the final cycle joint population density of grid points with column names equal to the name of each random parameter plus \$prob for the associated probability of that point
	\$popMean	The final cycle mean for each random parameter distribution
	\$popSD	The final cycle standard deviation for each random parameter distribution
	\$popCV	The final cycle coefficient of variation for each random parameter distribution
	\$popVar	The final cycle variance for each random parameter distribution
	\$popCov	The final cycle covariance matrix for each random parameter distribution
	\$popCor	The final cycle correlation matrix for each random parameter distribution
	\$popMedian	The final cycle median for each random parameter distribution
	\$popRanFix	The final cycle values for all fixed but random parameters
	\$postPoints	(NPAG only) Data frame of the Bayesian posterior parameter points for each of the first 100 subjects, with the following columns: <b>id</b> : subject ID <b>point</b> : point number for that subject <b>parameters</b> : parameters in the model <b>prob</b> : probability of each point in the posterior for each patient
	\$postMean	A matrix of means of posterior distributions for each subject and parameter
	\$postSD	A matrix of SDs of posterior distributions for each subject and parameter
	\$postVar	A matrix of variances of posterior distributions for each subject and parameter
	\$postCov	An array of covariance matrices for posterior parameter values for each subject

Objects	Variables	Comments
	\$postCor	An array of correlation matrices for posterior parameter values for each subject
	\$gridpts	(NPAG only) The initial number of support points
	\$ab	Matrix of boundaries for random parameter values. For NPAG, this is specified by the user prior to the run; for IT2B, it is calculated as a user specified multiple of the SD for the parameter value distribution
cycle (class: PMcycle, list)	\$names	Vector of names of the random parameters
	\$ll	Matrix of cycle number and -2*Log-likelihood at each cycle
	\$gamlam	A matrix of cycle number and gamma or lambda at each cycle (see item #16 under <a href="#">NPAG Runs</a> below for a discussion of gamma and lambda)
	\$mean	A matrix of cycle number and the mean of each random parameter at each cycle, normalized to initial mean
	\$sd	A matrix of cycle number and the standard deviation of each random parameter at each cycle, normalized to initial standard deviation
	\$median	A matrix of cycle number and the median of each random parameter at each cycle, normalized to initial standard deviation
	\$aic	A matrix of cycle number and Akaike Information Criterion at each cycle
	\$bic	A matrix of cycle number and Bayesian (Schwartz) Information Criterion at each cycle
cov (class: PMcov, data.frame)	\$id	Subject identification
	\$time	Time for each covariate entry
	covariates...	Covariate values for each subject at each time, extracted from the raw data file

Objects	Variables	Comments
	parameters...	Mean, median, or mode of Bayesian posterior distribution for each random parameter in the model. Mode summaries are available for NPAG output only, and the default is median. Values are recycled for each row within a given subject, with the number of rows driven by the number of covariate entries
	\$icen	Median (default) or mean of the covariates and parameter value distributions.
pop (class: PMpop, data.frame)  post (class: PMpost, data.frame)  NPAG only	\$id	Subject identification
	\$time	Time of each prediction at a frequency specified in the NPrun () command, with a default of 12 minutes.
	\$icen	Median (default) or mean of the parameter distributions used to calculate the predicted values.
	\$pred	Population prior (PMpop) or Bayesian posterior (PMpost) predictions for each output equation
	\$outeq	Output equation for each prediction
	\$block	Same as for PMop objects above
NPdata (class: NPAG, list)  ITdata (class: IT2B, list)		Raw data used to make the above objects. Please use ?NPparse or ?ITparse in R for discussion of the data contained in these objects
mdata (class: PMmatrix, data.frame)	See <a href="#">Pmetrics Input Files</a> .	Your original raw data file.
NPDE (class: PMnpde, list)  This object will only be present if you have run makeNPDE () after a run is completed.	Use the command str (NPDE . x) in R, where x is the run number	This object contains the information to perform graphical and numerical analysis of normalized prediction distribution errors. It is a method of <a href="#">internal model validation</a> .

Objects	Variables	Comments
sim (class: PMsim, list)  This object will only be present if you have run makeNPDE () after a run is completed.	\$obs	A data frame with \$id, \$time, \$out, \$outeq columns containing simulated observations at each time and output equation number in the template data file. If simulations from multiple template subjects have been combined (see <a href="#">Simulator Runs</a> ), then \$id will be of the form x.y, where x is the simulation number, and y is the template number.
	\$amt	A data frame with \$id, \$time, \$out, \$comp columns containing simulated amounts in each compartment.
	\$parValues	A data frame with \$id, ... columns containing the parameter value sets for each simulated subject with “...” signifying the columns named according to the names of the random parameters in the model
	\$totalSets	The total number of simulated sets of parameters, which may be greater than the requested number if limits were specified in the simulation (see <a href="#">Simulator Runs</a> ).
	\$totalMeans	The means of the parameter values in the total simulated sets which can be used as a check of the adequacy of the simulation to reproduce the requested mean values when limits were applied. The final truncated set will likely not have the requested mean values.
	\$totalCov	The covariances of the parameter values in the total simulated sets which can be used as a check of the adequacy of the simulation to reproduce the requested covariance values when limits were applied. The final truncated set will likely not have the requested covariance values.

Since R is an object oriented language, to access the observations in a **PMop** object, for example, use the following syntax: op\$post1\$obs.

Note that you will place an integer corresponding to the run number within the parentheses of the loading functions, e.g. PMload(1), which will suffix all the above objects with that integer, e.g. op.1, final.1, NPdata.1. This allows several models to be loaded into R simultaneously, each with a unique suffix, and which can be compared with the PMcompare () command (see [Model Diagnostics](#) below).

# Making New Pmetrics Objects

Once you have loaded the raw (**NPdata** or **ITdata** and **mdata**) or processed (**op**, **final**, **cycle**, **pop**, **post**) data objects described above with `PMload(run_num)`, should you wish to remake the processed objects with parameters other than the defaults, you can easily do so with the `make` family of commands. For example, the default for **PMop** observed vs. predicted objects is to use the prediction based on the median of the population or posterior distribution. If you wish to use the mean of the distribution, remake the **PMop** object using `makeOP()`. If you wish to see all the cycle information in a **PMcycle** object, not omitting the first 10% of cycles by default, remake it using `makeCycle()`.

For all of the following commands, the data input is either **NPdata** or **ITdata**, with additional function arguments specific to each command. Accessing the help for each function in R will provide further details on the arguments, defaults and output of each command.

Command	Description	R help
makeAUC	Make a data.frame of class <b>PMauc</b> containing subject ID and AUC from a variety of inputs including objects of <b>PMop</b> , <b>PMSim</b> , <b>PMpop</b> , <b>PMpost</b> or a suitable data.frame	?makeAUC
makeCov	Generate a data.frame of class <b>PMcov</b> with subject-specific covariates extracted from the data .csv file. This object can be plotted and used to test for covariates which are significantly associated with model parameters.	?makeCov
makeCycle	Create a PMcycle object described in the previous section.	?makeCycle
makeFinal	Create a PMfinal object described in the previous section.	?makeFinal
makeOP	Create a PMop object described in the previous section.	?makeOP

Command	Description	R help
makeNCA	<p>Create a data.frame (class <b>PMnca</b>) with the output of a non-compartmental analysis using <b>PMmatrix</b> or <b>PMpost</b> data objects as input. The <b>PMnca</b> object contains several columns.</p> <ul style="list-style-type: none"> <li>• <b>id</b>: Subject identification</li> <li>• <b>auc</b>: Area under the time-observation curve, using the trapezoidal approximation, from time 0 until the second dose, or if only one dose, until the last observation</li> <li>• <b>aumc</b>: Area under the first moment curve</li> <li>• <b>k</b>: Slope by least-squares linear regression of the final 6 log-transformed observations vs. time</li> <li>• <b>auclast</b>: Area under the curve from the time of the last observation to infinity, calculated as [Final obs]/k</li> <li>• <b>aumclast</b>: Area under the first moment curve from the time of the last observation to infinity</li> <li>• <b>aucinf</b>: Area under the curve from time 0 to infinity, calculated as auc + auclast</li> <li>• <b>aumcinf</b>: Area under the first moment curve from time 0 to infinity</li> <li>• <b>mrt</b>: Mean residence time, calculated as 1/k</li> <li>• <b>cmax</b>: Maximum predicted concentration after the first dose</li> <li>• <b>tmax</b>: Time to cmax</li> <li>• <b>cl</b>: Clearance, calculated as dose/ausinf</li> <li>• <b>vdss</b>: Volume of distribution at steady state, calculated as cl*mrt</li> <li>• <b>thalf</b>: Half life of elimination, calculated as ln(2)/k</li> <li>• <b>dose</b>: First dose amount for each subject</li> </ul>	?makeNCA
makeErrorPoly	<p>This function plots first, second, and third order polynomial functions fitted to pairs of observations and associated standard deviations for a given output assay. In this way, the standard deviation associated with any observation may be calculated and used to appropriately weight that observation in the model building process. Observations are weighted by the reciprocal of the variance, or squared standard deviation. Output of the function is a plot of the measured observations and fitted polynomial curves and a list with the first, second, and third order coefficients.</p>	?makeErrorPoly

Command	Description	R help
makePTA	<p>This function performs a Probability of Target Attainment analysis for a set of simulated doses and time-concentration profiles. Targets (e.g. Minimum Inhibitory Concentrations), the type of target attainment (i.e. %time above target, Cmax:target , AUC:target, Cmin:target, or Cx:target, where x is any time point), and the success threshold (e.g. %time &gt; 0.7 or Cmax:target &gt; 10) can all be specified. Output is a list (class <b>PMpta</b>) with two objects</p> <ul style="list-style-type: none"> <li><b>Results:</b> A data frame with the following columns: <i>simnum</i> is the number of the simulation; <i>id</i> is the simulated profile number within each simulation; <i>target</i> is the specified target; and <i>pdi</i> is the target pharmacodynamic index, e.g. time &gt; target, auc:target, etc.</li> <li><b>Outcome:</b> A data frame summarizing the results with the following columns: <i>simnum</i> and <i>target</i> are as for results; <i>prop.success</i> column has the proportion with a <i>pdi</i> &gt; success, as specified in the function call; <i>pdi.mean</i> and <i>pdi.sd</i> columns have the mean and standard deviation of the target pharmacodynamic index (e.g. proportion end-start above target, ratio of Cmax to target) for each simulation and target. If targets was specified via makePTAtarget to be a sampled distribution, then the target column will be missing from the outcome table.</li> </ul> <p><b>PMpta</b> objects can be summarized with <code>summary(x)</code> and plotted with <code>plot(x)</code>.</p>	?makePTA
makePop makePost (NPAG only)	<p>These functions create data.frames of class <b>PMpop</b> and <b>PMpost</b>, respectively.</p> <p>The <b>PMpop</b> or <b>PMpost</b> object contains several columns as described in the previous section.</p>	?makePop ?makePost

Command	Description	R help
makeNPDE	<p>This function is a Pmetrics wrapper to the autoNPDE function in the NPDE package of Comets et al (automatically loaded with Pmetrics) that will generate an <b>PMnpde</b> object which is a list of NpdeObjects (one for each output equation). NpdeObjects contain normalized prediction distribution errors. For a given NPAG or IT2B run number and output equation, this function will iterate through the data .csv file, using each subject as a template to simulate nsim new individuals from the population prior. It is HIGHLY recommended to use the default value of 1000 for nsim for the most valid calculation of NPDE. More than this could take a long time to execute. The mean population values will be used for each parameter and the covariance matrix. Errors may arise if extreme or negative concentrations are simulated from excessively large covariance matrices.</p> <p>Because considerable time may be necessary to make the NpdeObject, it will be added as an <b>NPDE</b> item to the NPAGout.Rdata or IT2Bout.Rdata objects so that it will be loaded the next time <code>PMload()</code> is run.</p> <p>Additionally, the combined simulations for all the subjects in the dataset will be saved as a <b>sim</b> item in the NPAGout.Rdata or IT2B.Rdata objects.</p>	?makeNPDE ?NPDE::autoNPDE ?NPDE::NpdeObject

## Summarizing Pmetrics Objects

There are summary commands available for several Pmetrics objects, as detailed below. All objects can be summarized by the R command `summary(x)`, where *x* is the object you wish to summarize.

### PMcov

```
summary(x, icen = "median")
?summary.PMcov
```

Summarize a PMcov object by creating a data frame with each subject's covariate values and Bayesian posterior parameter values, summarized according to icen. Default is "median" covariate values and Bayesian posterior parameter values, but could be "mean".

#### Example

```
data(PMex1)
summary(cov.1, "mean")
id   time  wt      africa age   gender height  Ka          Ke          V          Tlag1

```



### Example

```
data(PMex1)
summary(final.1)
```

par	type	quantile	value
Ka	WtMed	0.025	5.19E-01
Ka	WtMed	0.5	7.05E-01
Ka	WtMed	0.975	8.48E-01
Ka	MAWD	0.025	5.19E-02
Ka	MAWD	0.5	1.59E-01
Ka	MAWD	0.975	3.07E-01
Ke	WtMed	0.025	3.51E-02
Ke	WtMed	0.5	4.39E-02
Ke	WtMed	0.975	6.48E-02
Ke	MAWD	0.025	4.94E-03
Ke	MAWD	0.5	1.48E-02
Ke	MAWD	0.975	2.18E-02
V	WtMed	0.025	6.47E+01
V	WtMed	0.5	7.25E+01
V	WtMed	0.975	1.00E+02
V	MAWD	0.025	6.25E+00
V	MAWD	0.5	2.07E+01
V	MAWD	0.975	3.73E+01
Tlag1	WtMed	0.025	6.88E-01
Tlag1	WtMed	0.5	1.11E+00
Tlag1	WtMed	0.975	1.79E+00
Tlag1	MAWD	0.025	1.70E-01
Tlag1	MAWD	0.5	5.02E-01
Tlag1	MAWD	0.975	7.89E-01

In this example, the weighted median for Tlag1 is 1.11, with a 95% CI around the weighted median of 0.0688 to 1.79. The median absolute weighted difference (MAWD) is 0.502 with a 95% CI of 0.17 to 0.789.

### **PMmatrix**

```
summary(x, formula, FUN, ..., include, exclude)
?summary.PMfinal
```

This function will summarize a Pmetrics data file, which is of class `PMmatrix` when loaded by `PMreadMatrix()` or `PMload()`. The simplest is to summarize just the object.

For example:

```
data(PMex1)
summary(mdata.1)
```

```
Number of subjects: 20
Number of inputs: 1
Number of outputs: 1
Total number of observations (outeq 1): 139, with 0 (0.000%) missing
Number of covariates: 5
```

```
THE FOLLOWING ARE MEAN (SD), MIN TO MAX
-----
```

#### INPUTS

```
Number of doses per subject (input 1): 6.000 (0.000), 6.000 to 6.000
Dose per subject (input 1): 585.000 (45.189), 450.000 to 600.000
```

#### OUTPUTS

```
Number of obs per subject (outeq 1): 6.950 (0.224), 6.000 to 7.000
Observation per subject (outeq 1): 7.241 (3.799), 1.860 to 20.150
```

#### COVARIATES

```
wt: 54.538 (7.173), 43.000 to 66.500
africa: 1.000 (0.000), 1.000 to 1.000
age: 27.035 (7.717), 19.000 to 54.000
gender: 0.749 (0.434), 0.000 to 1.000
height: 167.792 (7.562), 150.000 to 181.000
-----
```

```
Note: See help(summary.PMmatrix) for accessing specific items by name.
```

An object of class `summary.PMmatrix` is also returned. This object is a list with the following items.

- **nsub** The number of subjects.
- **ndrug** The number of drug inputs.
- **numeqt** The number of output equations.
- **nobsXouteq** The number of observations by output equation.
- **missObsXouteq** The number of missing observations by output equation.
- **ncov** The number of covariates.
- **covnames** The covariate names.
- **ndoseXid** The number of doses per input per subject.
- **nobsXid** The number of observations per output equation per subject.
- **doseXid** The doses per input per subject.
- **obsXid** The observations per output per subject.
- **formula** The results of including a formula.

To include a formula, use standard R notation for the formula and specify the aggregating function with FUN.

#### Example

To calculate average dose in mg/kg per subject...

```
summ <- summary(mdata.1, formula = I(dose/wt)~id, FUN="mean")
summ$formula
id    I(dose/wt)
```

```

1  12.847966
2  9.022556
3  12.847966
4  11.811024
5  9.118541
6  9.230769
7  11.605416
8  11.71875
9  10.909091
10 11.516315
11 10.619469
12 12.526096
13 9.917355
14 10.135135
15 10.465116
16 9.31677
17 10.948905
18 10.158014
19 12
20 10.169492

mean(summ$formula[,2]) #choose the second column
[1] 10.84424

```

## PMop

```

summary(x, digits = max(3,getOption("digits") - 3), pred.type = "post", icen =
"median", outeq = 1)
?summary.PMop

```

This will summarize a PMop object that contains observations, predictions and errors. You can specify the number of digits, predictions based on posterior or population predictions, or based on mean or median parameter distributions, for any output equation.

### Example

```

data(PMex1)
summary(op.1) #summarize based on medians of Bayesian posterior parameter values

```

Time	Obs	Pred
------	-----	------

```

Min      120.0000  1.8600  1.9648
25%     121.0250  4.4950  4.5019
Median   126.0000  6.5600  6.5996
75%     132.0000  8.8850  8.9355
Max      144.9800  20.1500 19.8025
Mean     127.6899  7.2407  7.0689
SD       7.8600   3.7992  3.6420

```

Mean prediction error: -0.1718

Mean weighted prediction error (bias): -0.103 (P=0.9718 different than 0)

Mean squared prediction error: 0.795

Root mean squared error (RMSE): 0.8916

Percent root mean squared error (%RMSE): 12.3141

Mean weighed squared prediction error: 0.992

Bias-adjusted mean squared prediction error: 0.7655

Bias-adjusted mean weighted squared prediction error (imprecision): 0.9813

This function also returns a list with three items. The first item of the list is a data frame with the minimum, first quartile, median, third quartile, maximum, mean and standard deviation for times, observations and predictions in *x*. The second is a data frame of one row whose columns contain the mean prediction error, the mean weighted prediction error (bias), the mean squared prediction error, root mean squared error (RMSE), percent root mean squared error ( squared prediction error, the bias-adjusted mean squared prediction error, and the bias- adjusted mean weighted squared prediction error (imprecision). The third is itself a list of 6 items related to the weighted prediction bias, which are the mean, the standard error of the mean, the 95% confidence interval, the t-statistic, the degrees of freedom, and the p-value (compared to a bias of 0). In the example above, the p-value is 0.9718.

## PMpta

```

summary(x, ci = 0.95, ...)
?summary.PMpta

```

This function returns a list with two named objects: **pta** (probability of target attainment) and **pdi** (pharmacodynamic index).

### pta

A data frame with the following columns: **simnum**, **target**, **prop.success**, **pdi.mean**, and **pdi.sd** **simnum** is the number of the simulation; **target** is the specified target; **success** has the proportion with a ratio > prop.success; **pdi.mean** and **pdi.sd** are the mean and standard deviation of the pharmacodynamic index (e.g. AUC/MIC) for each simulation and target.

### pdi

A data frame with the following columns: **target**, **simnum**, **lowerCI**, **median**, **upperCI**. **target** and **simnum** are as above. **lowerCI**, **median**, and **upperCI** are the lower limit, median, and upper limit of the confidence interval for the pdi whose width is specified by **ci**.

# NPAG Runs

Here we provide details of the arguments available to the `NPrun()` command. You must have a data and model file in your working directory. In the R syntax below, any argument with a value has a default equal to that value and if you wish to use that default, you do not have to specify the argument in your function call to `NPrun()`.

```
NPrun(model = "model.txt", data = "data.csv", run,
       include, exclude, ode = -4, tol = 0.01, salt,
       cycles = 100, indpts, icen = "median",
       aucint, idelta = 12, prior, overwrite = F, nocheck = F, parallel = NA)
```

1. **model** Name of a suitable model file template in the working directory or an existing (previous) run number corresponding to a folder in the current working directory that used the same model file as will be used in the current run. If this is supplied, then the model file will be copied into the current working directory for convenience. If not supplied, the default is "model.txt". This file will be converted to a fortran model file. If it is detected to already be a fortran file, then the analysis will proceed without any further file conversion. *Examples:* `NPrun()`, `NPrun(model="model2.txt")`, `NPrun(model=2)`. The first example uses all the default options. The second example uses the defaults except for the model file. The third example uses the same model as that used in Run 2, but defaults otherwise.
2. **data** Name of a suitable data file or an existing (previous) run number corresponding to a folder in the current working directory that used the same data file as will be used in the current run. If this is supplied, then previously made '.ZMQ' files will be copied into the current working directory, bypassing the need to re-convert the .csv file and speeding up the run. *Examples:* `NPrun(data="data2.csv")`, `NPrun(data=2)`. The first example uses the defaults except for the data file. The second example uses the same data file as that used in Run 2, but defaults otherwise.
3. **run** Specify the run number of the output folder. Default if missing is the next available number. *Examples:* `NPrun()`, `NPrun(run=2)`. The first example uses all the default options and the run number will be automatically assigned to "3" if runs 1 and 2 already exist. The second example uses the defaults except for the run number. If runs 1 and 3 exist and you have deleted run 2 because you wish to re-run it, then this syntax will place the output in a folder labeled "2". If you haven't already deleted the old run folder, you can set **overwrite #15** below, to True.
4. **include** Vector of subject id values in the data file to include in the analysis. The default (missing) is all. *Examples:* `NPrun(include=c(1:3, 5))`. Include only subjects with ID numbers of 1, 2, 3 and 5.
5. **exclude** Vector of subject id values in the data file to exclude in the analysis. The default (missing) is none. Be careful if you include and exclude in the same run as you may have conflicting statements. *Examples:* `NPrun(exclude=c(1:3, 5))`. Exclude subjects with ID numbers of 1, 2, 3 and 5.
6. **ode** Ordinary Differential Equation solver  $\log_{10}$  tolerance or stiffness. Default is -4, i.e. 0.0001. Higher values will result in faster runs, but parameter estimates may not be as accurate. It is ignored if the model does not use differential equations. *Examples:* `NPrun(ode=-3)`. Do the run with a stiffness of 0.001, which will be faster than the default but perhaps less accurate.
7. **tol** Tolerance for convergence of NPAG. Smaller numbers make it harder to converge. Default value is 0.01. *Examples:* `NPrun(tol=0.001)`. Do the run with a tolerance of 0.001, which will be slower than the default but perhaps more accurate.
8. **salt** Vector of salt fractions for each ndrug, default is 1 for each drug. This is not the same as bioavailability. *Examples:* `NPrun(salt=c(1, 0.9))`. This sets the salt fraction for the first drug to 1 and for the second drug to 0.9.
9. **cycles** Maximum number of NPAG cycles to run. Default is 100, but the value can be 0 or greater. If you enter an integer greater than 0, the engine will terminate at convergence or the number of cycles you specify, whichever comes first. Early in model exploration values of 10 to 100 can be useful, with larger values later in model development. In order to facilitate model comparison, however, we recommend using the same cycle limit for all early models, e.g. 100, rather than choosing 10 for one and 100 for another. If you enter 0, this is the way to test the predictive power of a model on an independent data set and a non-uniform prior must be

specified. This means that the engine will only calculate the individual Bayesian posteriors for the new subjects, using the population joint density from a previous run as a Bayesian prior. *Examples:* `NPrun(cycles=1000)`, `NPrun(data="newdata.csv",cycles=0,prior=NPdata.1)`. The first example will allow NPAG to run 1000 cycles before terminating. The second example calculates Bayesian posteriors only (i.e. NPAG is not cycling) using the default model file and some new data. Note by specifying a prior, which is required with 0 cycles, the grid points are distributed in a non-uniform manner corresponding to the prior distribution in NPdata.1 (the output of Run 1). This means that the `indpts` argument will be ignored. See also **prior** #14 below.

10. **indpts** Index of starting grid point number. Default is missing, which allows NPAG to choose depending on the number of random parameters: 1 or 2 parameters = index of 1; 3 = 3; 4 = 4, 5 = 6, 6 or more is 10+number of multiples for each parameter greater than 5, e.g. 6 = 101; 7 = 102, up to 108 for 13 or more parameters. This number corresponds to the number of grid (support) points which will initially fill the model parameter space. The larger the number of random parameters to be estimated, the more points are required. The more you choose, the slower the run will be, but results may improve. It is reasonable to choose fewer points early in model exploration and increase in later phases or if poor model fits or lack of convergence are noted. Indices of 1 to 6 correspond respectively to 2129, 5003, 10007, 20011, 40009, and 80021 points. 101 will choose 80021 points plus one additional multiple of 80021 points. *Examples:* `NPrun(indpts=102)`. This will allow NPAG to start with  $80021 + 2*80021 = 240063$  points. The unusual numbers are from early development of NPAG where primes were thought to be important.
11. **icen** Summary of parameter distributions to be used to calculate predictions. Default is "median", and other choices are "mean" or "mode". *Examples:* `NPrun(icen="mean")`. This example will use the means of the parameter distributions (rather than the medians) to calculate predictions.
12. **aucint** Interval for AUC calculations. Default is 24 hours if the number of intervals is not greater than 48; otherwise it defaults to the interval which allows for  $\leq 48$  intervals. *Examples:* `NPrun(aucint=12)`. This example will use an interval of 12 hours for AUC estimations directly by NPAG. Note that the command `makeAUC()` can be used to calculate AUCs of any interval from the output.
13. **idelta** Interval in minutes for predictions at times other than observations. Default is 12 minutes, which in general, for most models provides sufficient granularity. Smaller values can result in very large files for big populations. *Examples:* `NPrun(idelta=60)`. This generates predictions every 60 minutes.
14. **prior** Name of a suitable NPAG output object from a prior run loaded with `PMload()`, i.e. the NPdata object. A prior may be specified if the user wishes to start from a non-uniform prior distribution for the NPAG run. The default value is -99, which translates in NPAG to a uniform prior distribution. An alternative is to include a DEN0001 file from the prior NPAG run in the working directory of the new run, and specify this as the value for prior, e.g. `prior = 'DEN0001'`. *Examples:* `NPrun(data=2,model=2,prior=NPdata.2)`. This example will use the model, data, and population density files from Run 2 to continue for another 100 cycles. This is a useful way to continue a previous run that has not converged. See also **cycles** #9 above for a discussion and example of specifying a prior simply to calculate the Bayesian posterior parameter and output values for a population.
15. **overwrite** Overwrite existing run result folders. Only relevant when the **run** (#3) argument is specified. Default is FALSE. *Examples:* `NPrun(run=2,overwrite=T)`. This example will delete the previous Run 2 and replace it with the current run.
16. **nocheck** The default is false. Set to true to suppress automatic checking of data files for errors with `PMcheck()`, which depends on Java. If Java is out of date, or is not a 64-bit version on 64-bit systems, `PMcheck()` can fail. Update Java (ensure 64-bit if applicable), but use this argument as a temporary fix.
17. **parallel** As of version 1.4.0, Pmetrics can run in parallel processing mode. The default is NA, which allows Pmetrics to choose parallel mode for models with differential equations, or serial mode for algebraic models. This is because the overhead of managing the parallel threads is greater than the efficiency of algebraic processing. NOTE: Parallel processing is not possible on 32-bit systems, and this argument will always be set to false.

The NPAG run can complete in seconds for small populations with analytic solutions, or days for large populations with complex differential equations. At the end of a successful run, the results will be automatically parsed and saved to the output directory. Your default browser will launch with a summary of the run.

## IT2B Runs

Here we provide details of the arguments available to the `ITrun()` command. You must have a data and model file in your working directory. In the R syntax below, any argument with a value has a default equal to that value and if you wish to use that default, you do not have to specify the argument in your function call to `ITrun()`.

```
ITrun(model = "model.txt", data = "data.csv", run,
       include, exclude, ode = -4, salt, cycles = 100,
       tol = 0.001, xdev = 5, icen = "median",
       overwrite = F, nocheck = F)
```

1. **model** Same as for NPAG.
2. **data** Same as for NPAG.
3. **run** Same as for NPAG.
4. **include** Same as for NPAG.
5. **exclude** Same as for NPAG.
6. **ode** Same as for NPAG.
7. **salt** Same as for NPAG.
8. **cycles** Maximum number of IT2B cycles to run. Default is 100, but the value can be 1 or greater. Note that unlike NPAG, you cannot enter a value of 0. If you enter an integer greater than 0, the engine will terminate at convergence or the number of cycles you specify, whichever comes first. *Examples:* `ITrun(cycles=1000)`. The first example will allow IT2B to run 1000 cycles before terminating.
9. **tol** Same as for NPAG, except the default value is 0.001.
10. **xdev** Multiple of standard deviations for parameters to be used in NPAG as a range. Default is 5. The ranges can be found in the FROM0001 file in the /outputs folder of the run folder. IT2B is primarily used to estimate ranges for parameter values to be passed to NPAG. Because it is parametric, it is not constrained to the initial parameter ranges in the model file, unlike NPAG. *Examples:* `ITrun(xdev=3)`. Recommend parameter ranges which are equal to the mean  $\pm 3$  SD. You might choose a smaller range like this if you have log transformed your parameters.
11. **icen** Same as for NPAG.
12. **overwrite** Same as for NPAG.
13. **nocheck** Same as for NPAG.

The IT2B run can complete in seconds for small populations with analytic solutions, or days for large populations with complex differential equations. At the end of a successful run, the results will be automatically parsed and saved to the output directory. Your default browser will launch with a summary of the run.

## ERR Runs

Here we provide details of the arguments available to the `ERRrun()` command. This function is identical to `ITrun()` but the purpose is to estimate the assay error coefficients ( $C_0, C_1, C_2, C_3$ ) from the data if you have no idea what they are. Note that the estimates you get will be dependent on the model. You must have a data and

model file in your working directory. In the R syntax below, any argument with a value has a default equal to that value and if you wish to use that default, you do not have to specify the argument in your function call to `ERRrun()`.

```
ERRrun(model = "model.txt", data = "data.csv", run,
include, exclude, ode = -4, salt, cycles = 100,
search = "cursory", tol = 0.001, xdev = 5, overwrite = F)
```

1. **model** Same as for NPAG, IT2B.
2. **data** Same as for NPAG, IT2B.
3. **run** Same as for NPAG, IT2B.
4. **include** Same as for NPAG, IT2B.
5. **exclude** Same as for NPAG, IT2B.
6. **ode** Same as for NPAG, IT2B.
7. **salt** Same as for NPAG, IT2B.
8. **cycles** Maximum number of ERR cycles to run. Default is 100, but the value can be 1 or greater. Note that unlike NPAG, you cannot enter a value of 0. If you enter an integer greater than 0, the engine will terminate at convergence or the number of cycles you specify, whichever comes first. *Examples:* `ERRrun(cycles=1000)`. The first example will allow ERR to run 1000 cycles before terminating.
9. **search** Depth of the search for the coefficients. Default is "cursory", but can be "medium" or "extensive", which take progressively longer times to converge, but are more accurate.
10. **tol** Same as for IT2B.
11. **xdev** Same as for IT2B.
12. **overwrite** Same as for NPAG.

The ERR run can complete in seconds for small populations with analytic solutions, or days for large populations with complex differential equations. At the end of a successful run, the results will be automatically parsed and saved to the output directory. Your default browser will launch with a summary of the run that includes the estimates of the assay error coefficients for each output equation in the model.

## Simulator Runs

The Pmetrics simulator is a powerful Monte Carlo engine that is smoothly integrated with Pmetrics inputs and outputs. Unlike NPAG and IT2B, it is run from within R. No batch file is created or terminal window opened. However, the actual simulator is a Fortran executable compiled and run in an OS shell. It is documented with an example within R. You can access this by using the `help(SIMrun)` or `?SIMrun` commands from R.

In order to complete a simulator run you must include a data .csv file and a model file in the working directory. The structure of these files is identical to those used by NPAG and IT2B. The data .csv contains the template dosing and observation history as well as any covariates. Observation values (the OUT column) for EVID=0 events can be any number; they will be replaced with the simulated values. However, **do not use -99**, as this will simulate a missing value, which might be useful if you are testing the effects of such values. A good choice for the OUT value in the simulator template .csv file is -1.

You can have any number of subject records within a data .csv file, each with its own covariates if applicable. Each subject will cause the simulator to run one time, generating as many simulated profiles as you specify from each template subject. This is controlled from the `SIMrun()` command with the `include` and `nsim` arguments. The first specifies which subjects in the data .csv file will serve as templates for simulation. The second specifies how many profiles are to be generated from each included subject.

Simulation from a non-parametric prior distribution (from NPAG) can be done in one of two ways. The first is simply to take the mean, standard deviation and covariance matrix of the distribution and perform a standard Monte Carlo simulation. The second way is what we call semi-parametric, and was devised by Goutelle et al.[1] In this method, the non-parametric “support points” in the population model, each a vector of one value for each parameter in the model and the associated probability of that set of parameter values, serve as the mean of one multi-variate normal distribution in a multi-modal, multi-variate joint distribution. The weight of each multi-variate distribution is equal to the probability of the point. The overall population covariance matrix is divided by the number of support points and applied to each distribution for sampling.

Limits may be specified for truncated parameter ranges to avoid extreme or inappropriately negative values. When you load simulator output with `SIMparse()`, it will include values for the total number of simulated profiles needed to generate `nsim` profiles within the specified limits, as well as the means and standard deviations of the simulated parameters to check for simulator accuracy.

Output from the simulator will be controlled by further arguments to `SIMrun()`. If `makecsv` is not missing, a .csv file with the simulated profiles will be created with the name as specified by `makecsv`; otherwise there will be no .csv file created. If `outname` is not missing, the simulated values and parameters will be saved in a .txt file whose name is that specified by `outname`; otherwise the filename will be “simout”. In either case, integers 1 to `nsub` will be appended to `outname` or “simout”, e.g. “simout1.txt”, “simout2.txt”.

Output files from the simulator can be read into R using the `SIMparse()` command (see documentation in R). There is a plot method (`plot.PMsim`) for objects created by `SIMparse()`.

Here we detail the arguments to the `SIMrun()` function.

```
SIMrun(poppar, limits = NULL, model = "model.txt", data = "data.csv",
       split = F, include, exclude, nsim = 1000, predInt = 0, covariate,
       usePost = F, seed = -17, ode = -4, obsNoise, doseTimeNoise = rep(0,
        4), doseNoise = rep(0, 4), obsTimeNoise = rep(0, 4), makecsv, outname,
       clean = T, silent = F, nocheck = F)
```

1. **poppar** Either an object of class PMfinal or a list containing three items in this order, but of any name: vector of weights, vector of mean parameter values, and a covariance matrix. By far the easiest is to use the PMfinal object. However, the list option is available, for example, to simulate with values obtained from the literature or other sources. For the list option, if only one distribution is to be specified, the weights vector should be of length 1 and contain a 1. If multiple distributions are to be sampled, the weights vector should be of length equal to the number of distributions and its values should sum to 1, e.g. `c(0.25,0.05,0.7)`. The means matrix may be a vector for a single distribution, or a matrix with `length(weights)` rows and number of columns equal to the number of parameters, `npar`. The covariance matrix will be divided by `length(weights)` and applied to each distribution.

*Examples:* `SIMrun(poppar=final.1, ...)`,

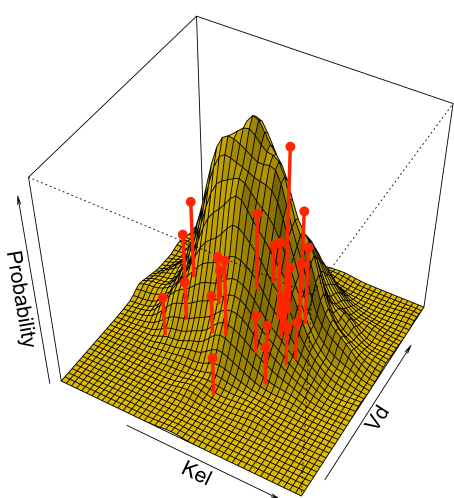
`SIMrun(poppar=list(weight=1, mean=c(2, 4), covar=diag(2, 2)), ...)`, `SIMrun(poppar=list(weight=c(0.75, 0.25), mean=matrix(c(2, 4, 1, 2), nrow=2, byrow=T), covar=diag(2, 2)), ...)`. The first example will use the population parameter distribution from Run 1 (NPAG or IT2B). The second example will use a distribution for two parameters with mean values of 2 and 4, and a covariance matrix with variances of 2 (the diagonals) and covariances of 0 (the off-diagonals). The third example will simulate from a bimodal, bivariate distribution. The first distribution, with weight 0.75, has means of 2 and 4 for the parameters. The second distribution, with weight 0.25, has means of 1 and 2. The overall covariance is the same as in example 2, but will be divided evenly across the two distributions. Example 3 is similar to what occurs with a PMfinal poppar object when `split` #5, below, is true.

2. **limits** If limits are specified, each simulated parameter set that contains any parameter value outside of the limits will be ignored and another set will be generated. This will result in a truncated distribution. See `SIMparse` for details on the full and truncated distributions returned by the simulator. Four options exist for limits. 1) The default `NULL` indicates that no limits are to be applied to simulated parameters. 2) The second option is to set limits to `NA`. This will use the parameter limits on the primary parameters that are specified in the model file. 3) The third option is a numeric vector of length 1 or 2, e.g. `3` or `c(0.5,4)`, which specifies what to multiply the columns of the limits in the model file. If length 1, then the lower limits will be the same as in the model file, and the upper limits will be multiplied by value specified. If length 2, then the lower and upper limits will be multiplied by the specified values. If this option is used, `poppar` must be a PMfinal object. 4) The fourth option for limits is a fully customized matrix of limits for simulated values for each parameter which will

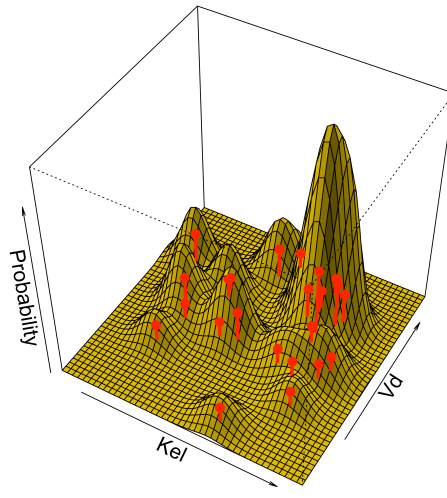
overwrite any limits in the model file. If specified, it should be a data.frame or matrix with number of rows equal to the number of random parameters and 2 columns, corresponding to the minimum and maximum values. For example, a `final$ab` object, or a directly coded matrix, e.g. `matrix(c(0.5,0.5,0.01,100), nrow=3,ncol=2,byrow=T)` for 3 parameters with limits of [0.5], [0.5] and [0.01,100], respectively. It is possible to convert a parameter to fixed by omitting the second limit. Means and covariances of the total number of simulated sets will be returned to verify the simulation, but only those sets within the specified limits will be used to generate output(s) and the means and covariances of the retained sets may (and likely will be) different than those specified by poppar.

*Examples:* `SIMrun(...,limits=NULL)`, `SIMrun(...,limits=NA)`, `SIMrun(...,limits=3)`, `SIMrun(...,limits=c(0,5)`, `SIMrun(...,limits=matrix(c(0,1,0.5,1000), nrow=2,byrow=T))`. The first example has no limits. The second example uses the limits on the primary parameters in the model file. The third example uses the lower limits on the primary parameters in the model file, and multiplies the upper limits of each primary parameter in the model file by 3. The fourth example multiplies the lower limits of the primary parameter values in the model file by 0, and the upper limits are multiplied by 5. The fifth example sets a custom matrix of limits, with ranges of 0 to 1 for the first parameter, and 0.5 to 1000 for the second.

3. **model** Name of a suitable model file template in the working directory. The default is “model.txt”. This file will be converted to a fortran model file. If it is detected to already be a fortran file, then the simulation will proceed without any further file conversion. This is the same model file used for NPAG and IT2B runs.
4. **data** Either a PMmatrix object previously loaded with PMreadMatrix or character vector with the filename of a Pmetrics data file that contains template regimens and observation times. This file is the same format as for NPAG and IT2B runs, and the value for the OUT column (EVID=0 rows) in this file can be coded as any number(s) other than -99. We typically use -1. The number(s) will be replaced in the simulator output with the simulated values.
5. **split** Boolean operator controlling whether to split an NPAG PMfinal object into one distribution per support point, with means equal to the vector of parameter values for that point, and covariance equal to the population covariance divided by the number of support points. See the figures below to better understand what happens when **split** is true or false. The red points are the non-parametric population distribution, while the gold wireframe overlays are smoothed probability functions from which samples are drawn in simulations.



**split=F**, unimodal, multivariate sampling



**split=T**, multimodal, multivariate sampling

6. **include** A vector of subject IDs in the data file to iterate through, with each subject serving as the source of an independent simulation. If missing, all subjects in the datafile will be used. This is analogous to the same argument in `NPrun()` and `ITrun()`.

7. **exclude** A vector of subject IDs to exclude in the simulation, e.g. `c(4,6:14,16:20)` If a `makecsv` filename is supplied, ID numbers will be of the form `nsub.nsim`, e.g. 1.001 through 1.1 for the first subject, 2.001 through 2.1 for the second subject, etc. if 1000 simulations are made from each subject. This is analogous to the same argument in `NPrun()` and `ITrun()`.
8. **nsim** The number of simulated profiles to create, per subject. Default is 1000.
9. **predInt** The interval in fractional hours for simulated predicted outputs at times other than those specified in the template data. The default is 0, which means there will be simulated outputs only at times specified in the data file. Values of `predInt > 0` result in simulated outputs at the specified value of `predInt`, e.g. every 15 minutes for `predInt = 0.25` from time 0 up to the maximal time in the template file, per subject if `nsub > 1`. You may also specify `predInt` as a vector of 3 values, e.g. `c(1,4,1)`, similar to the R command `seq`, where the first value is the start time, the second is the stop time, and the third is the step value. Outputs for times specified in the template file will also be simulated. To simulate outputs only at the output times in the template data (i.e. `EVID=0` events), use `predInt=0`, which is the default. Note that the maximum number of predictions total is 594, so the interval must be sufficiently large to accommodate this for a given number of output equations and total time to simulate over. If `predInt` is set so that this cap is exceeded, predictions will be truncated.

*Examples:* `SIMrun(...)`, `SIMrun(..., predInt=1)`, `SIMrun(..., predInt=c(1,12,0.5))`. The first example is the same as `predInt=0` (the default) which will simulate observations only at the times specified in the template date file (i.e. the times of all `EVID=0` events). The second example simulates observations hourly starting at 1 hour, plus any specified by `EVID=0` event times in the data file, and continuing until the maximum time in the template data file. If the number of simulated observations exceeds 594, they will be truncated to this value. The third example simulates observations half-hourly from 1 to 12 hours, plus at any `EVID=0` event times in the data file.

10. **covariate** If you are using the results of an NPAG or IT2B run to simulate, i.e. a `PMfinal` object as `poppar`, then you can also simulate with covariates. This argument is a list with names “cov”, “mean”, “sd”, “limits”, and “fix”.
  - The first item is a `PMcov` object, such as that loaded with `PMload`. Pmetrics will use this object to calculate the correlation matrix between all covariates and Bayesian posterior parameter values.
  - The second item, “mean”, allows you to specify a different mean for one or more of the covariates. This argument is a named list, where each item in the list is the name of a covariate in your data that is to have a different mean. If this argument is missing then the mean covariate values in the population will be used for simulation. The same applies to any covariates that are not named in the “mean” list.
  - The third item in the covariate argument list is “sd”, and this functions just as the “mean” object does - allowing you to specify different standard deviations for covariates in the simulation. If “sd” is missing, then the standard deviations of the covariates in the population are used.
  - The fourth item in the covariate argument list is “limits”, and this functions a bit differently than the limits argument for parameters does (#2 above). Because the desired limits of a covariate, e.g. weight, are generally known, unlike for population parameters, this argument is specified in the same way as for `mean` and `sd`, that is, a named list with the limits for each covariate. If this list item is missing altogether, no limits will be applied to simulated covariates. If it is supplied, then covariates which are not fixed and not included in the list will have the same limits as in the original population. If you want to simulate some covariates with limits and some without, specify the latter with very large ranges.
  - The fifth and final item in the covariate argument list is “fix”. This is simply a character vector of the names of the covariates that you wish not to simulate, i.e. fix to the values in the template data file.

Whether you use the means and standard deviations in the population or specify your own, the covariance matrix in `poppar` will be augmented by the covariate covariances. The parameter plus covariate means and this augmented covariance matrix will be used for simulations. In effect, all covariates are moved into the `#Primary` block of the model file to become parameters that are simulated. In fact, a copy of your model file is made with a “`c_`” prepended to the model name (e.g. “`model.txt`” -> “`c_model.txt`”). Likewise, the data file is copied to a new file with “`c_`” prepended to the data file name (e.g. “`data.csv`” -> “`c_data.csv`”).

*Examples:* assuming that the covariates `wt` (weight) and `ccr` (creatinine clearance) are in your data and you are simulating from run 1. `SIMrun(..., covariate=list(cov=cov.1))`. This will simulate all

parameters and covariates that were present in run 1, using the covariate means and covariances that existed in the population. No limits will be applied. `SIMrun(...,covariate=list(cov=cov.1,mean=list(wt=100)))`. Note the list within a list. This simulation will be the same as the first example, except that the mean weight will now be 100, retaining the same standard deviation as in the original population. `SIMrun(...,covariate=list(cov=cov.1, mean=list(wt=100), sd=list(wt=30)))`. This will now also replace the standard deviation of weight in the population with 30. `SIMrun(...,covariate=list(cov=cov.1,limits=list(wt=c(10, 90), age=c(-1,1000)))`. This will use the existing means and covariances of the covariates in the population, but will limit simulated values for weight to be a minimum of 10 and maximum of 90. Age will effectively have no limits. Other covariates will have limits equal to the limits in the original population. `SIMrun(...,covariate=list(cov=cov.1,fix=c("wt", "age")))` This will simulate with existing means, standard deviations, and limits for all covariates except weight and age, which will be fixed to the values in the template data file used for simulations.

11. **usePost** Boolean argument. Only applicable when **poppar** is an NPAG PMfinal object. If so, and **usePost** is TRUE, the posterior for each subject (modified by include or exclude) in poppar will be used to simulate rather than the population prior. The number of subjects in the template data file must be the same. Normally one would use the same data file as used to make the model final parameter distribution in **poppar**, but if different templates are desired, the number must be equivalent to the number of included subjects from whom the posteriors are obtained.
12. **seed** The seed for the random number generator. For `nsub > 1`, should be a vector of length equal to `nsub`. Shorter vectors will be recycled as necessary. Default is -17. Examples: `SIMrun(...,seed=c(2,-3))`. This will run the simulator for the two subjects in the data file, with 2 as the seed for the first subject, and -3 for the second subject. The value of the seed is that it will generate the same random parameter sets each time the seed is specified. This allows random, but reproducible results for comparing simulations where other factors have changed, such as the dosing regimen used in the template data file. Setting the seed to fewer than the number of subjects (e.g. 1 seed for 3 subjects) will result in each subject having the same set of simulated parameter values because Pmetrics recycles the seed as needed. This may or may not be desirable.
13. **ode** Ordinary Differential Equation solver log tolerance or stiffness. Default is -4, i.e. 0.0001. Higher values will result in faster simulations, but simulated concentrations may not be as accurate. This is the same parameter as for NPAG and IT2B runs.
14. **obsNoise** This is the noise added to simulated observations. If present will override any other values in the data file or model file. Should be a vector of length 4 times the number of output equations, e.g. `c(0.1,0.1,0,0)` for one output and `c(0.1,0.1,0,0,0.01,0.2,-0.001,0)` for two output equations. Each number is a coefficient that allows calculation of the standard deviation of the normal distribution around zero from which the noise is generated, such that  $SD = C0 + C1*sim + C2*sim^2 + C3*sim^3$ , where "sim" is the simulated value. This is directly analogous to the assay error polynomial discussed in the section on the [model file](#) format. If you do not specify **obsNoise**, all coefficients will be set to 0 for all output equations, i.e. no noise. If you specify **obsNoise = NA**, values in the data file will be used (similar to **limits** above). If these values in the data file are missing, values in the model file will be used. Examples: `SIMrun(...)`, `SIMrun(...,obsNoise=NA)`, `SIMrun(...,obsNoise=c(0,0.1,0,0,1,0.15,0,0))`. The first example sets all the observation noise terms to 0. The second example uses the error terms in the model file. The third example sets the noise terms for the first output equation to 0, 0.1, 0, 0 and for the second output equation to 1, 0.15, 0, 0.
15. **doseTimeNoise** A vector of dose time error polynomial coefficients, analogous to **obsNoise** above. The default is 0 for all coefficients. This is applied to all dose times, regardless of drug, so can only be of length 4, e.g `c(0.1,0.1,0,0)`.
16. **doseNoise** A vector of dose amount error polynomial coefficients, analogous to **obsNoise** above. The default is 0 for all coefficients. This is applied to all dose amounts, regardless of drug, so can only be of length 4, e.g `c(0.1,0.1,0,0)`.
17. **obsTimeNoise** A vector of observation timing error polynomial coefficients. The default is 0 for all coefficients. This is applied to all observation times, regardless of output number, so can only be of length 4, e.g `c(0.1,0.1,0,0)`.
18. **makecsv** A character vector for the name of the single .csv file to be made for all simulated "subjects". If missing, no files will be made. Examples: `SIMrun(...,makecsv="newsubj.csv")`. This example will

make a .csv file in Pmetrics format called “newsubj.csv” containing the simulated subjects. If there were 3 subjects in the template data file, and nsim=10, then newsubj.csv will contain  $3*10=30$  simulated subject records.

19. **outname** The name for the simulator output file(s) without an extension. Numbers 1 to nsub will be appended to the files. If missing, will default to “simout”. *Examples:* SIMrun(..., outname="sim2\_"). This will make the following output files for a template with 3 subjects: sim2\_1.txt, sim2\_2.txt, sim2\_3.txt. These are the files that are parsed by SIMparse().
20. **clean** This is a rarely used Boolean parameter to specify whether temporary files made in the course of the simulation run should be deleted. It is primarily used for debugging. Defaults to True.
21. **silent** Boolean operator controlling whether a model summary report is given. Default is False.
22. **nocheck** Suppress the automatic checking of the data file with PMcheck(). Default is False.

To get the results of a simulator run back into R, you need to use the `SIMparse()` command, which is detailed below.

```
SIMparse(file, include, exclude, combine = F, silent = F, parallel)
```

Note that `SIMparse()` returns the parsed output of a simulator run as a **PMsim** object, so use the command like this: `simdata <- SIMparse(...)` so that `simdata` will contain the results. The arguments to `SIMparse` follow.

1. **file** An output file or files of the simulator in the current working directory, or the full pathname to the file. To load and combine multiple outputs, specify files separated by commas or using wild cards. For file specification “?” will be matched by just a single numeral or character; “\*” will be matched by any number of consecutive alphanumeric characters. *Examples:* `simdata <- SIMparse(file="simout1.txt, simout2.txt,simout3.txt")`, `simdata <- SIMparse(file="simout?.txt")` and `simdata <- SIMparse(file="sim*.txt")`. All three examples will find the files `simout1.txt`, `simout2.txt`, and `simout3.txt` in the working directory. The second example would also find `simout4.txt`, etc. The third example would also find `sim_1.txt` if that existed.
2. **include** A vector of files to include in the parsing. *Example:* `simdata <- SIMparse("simout?.txt", include=c(1, 4))`. If the wildcard match returned four files: `simout1.txt`, `simout2.txt`, `simout3.txt` and `simout4.txt`, and you wished to only parse the first and fourth files, this would accomplish that filtering.
3. **exclude** A vector of files to exclude in the parsing. *Example:* `simdata <- SIMparse("simout?.txt", include=c(1, 4))`. If the wildcard match returned four files: `simout1.txt`, `simout2.txt`, `simout3.txt` and `simout4.txt`, and you wished to only parse the second and third files, this would accomplish that filtering. Be careful using **include** and **exclude** at the same time; you might end up with no files!
4. **combine** Boolean parameter, default False, which specifies whether you wish to combine the parsed files into a single **PMsim** object. This can be useful for making visual predictive checks, for example. If `combine=F`, and multiple files are parsed, then the return object will be a list of **PMsim** objects, which can be plotted or otherwise accessed using standard list referencing, e.g. `plot(simlist[[1]])`, `plot(simlist[[2]])`, etc. In this case, each element of the list corresponds to a single simulation template. *Examples:* `simdata <- SIMparse("simout?.txt", combine=F)`, `simdata <- SIMparse("simout?.txt", combine=T)`. If your template data file had four subjects, and you did not combine the results as in the first example, `simdata` would be a list of length 4 (`simdata[[1]]`, `simdata[[2]]`, `simdata[[3]]`, `simdata[[4]]`), each containing a **PMsim** simulation output object corresponding to a subject in the template data file. In the second example, all 4 simulation outputs would be combined into a single **PMsim** object. You can combine any simulator output files, even with differing numbers of simulated parameter sets. The number of outputs and times of observations also may differ, although combining these may lead to strange plots since not all profiles have the same observations.
5. **silent** Suppress messages about parsing. Default is False.

6. **parallel** Runs in parallel mode. Defaults to true if multiple files are to be parsed, otherwise false. Can be overridden by specifying True or False

## Plotting

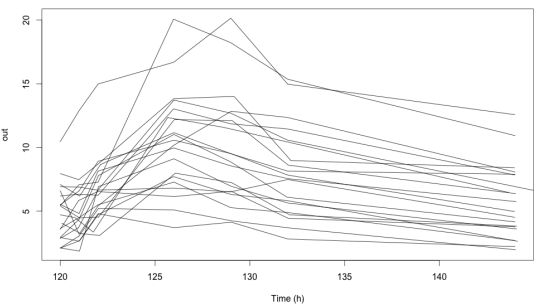
There are numerous plotting methods included in Pmetrics to generate standardized, but customizable graphical visualizations of Pmetrics data. Taking advantage of the class attribute in R, a single `plot()` command is used to access all of the appropriate plot methods for each Pmetrics object class.

To access the R help for these methods, you must query each method specifically to get details, for `?plot` will only give you the parent function.

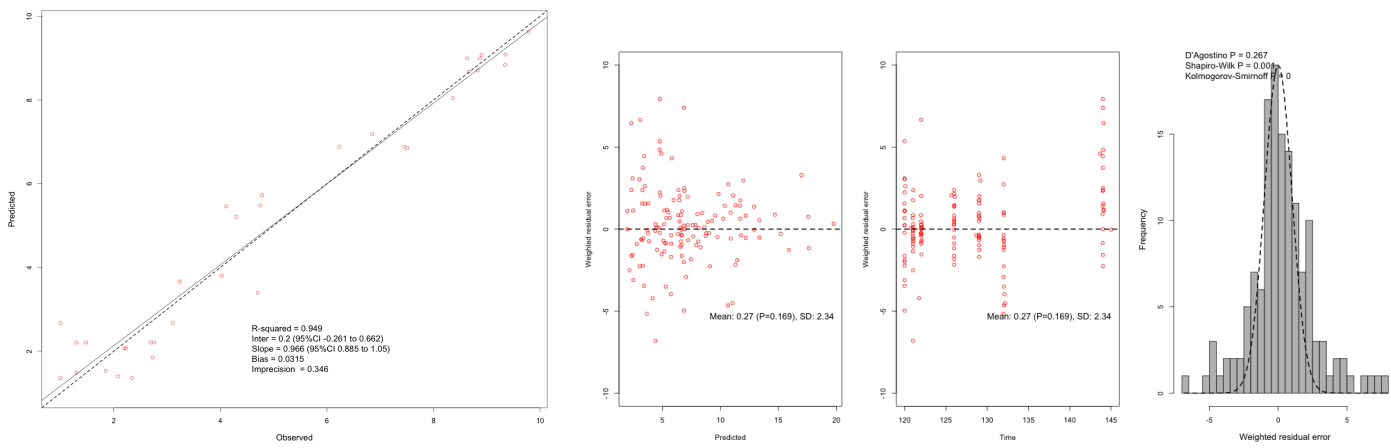
Object Classes	Creating functions	R help	Description
PMop	<code>PMload()</code> , <code>makeOP()</code>	<code>?plot.PMop</code>	Plot population or individual Bayesian posterior predicted data vs. observed. Optionally, you can generate residual plots.
PMfinal	<code>PMload()</code> , <code>makeFinal()</code>	<code>?plot.PMfinal</code>	Plot marginal final cycle parameter value distributions. Specifying a formula of the form $y \sim x$ will generate a bivariate plot. For NPAG only, a formula of the form $prob \sim x$ will plot the Bayesian posterior parameter $x$ distributions for included subjects.
PMcycle	<code>PMload()</code> , <code>makeCycle()</code>	<code>?plot.PMcycle</code>	Plots a panel with the following windows: -2 times the log-likelihood at each cycle, gamma/lambda at each cycle; Akaike Information Criterion at each cycle and Bayesian (Schwartz) Information Criterion at each cycle, the mean parameter values at each cycle (normalized to starting values); the normalized standard deviation of the population distribution for each parameter at each cycle; and the normalized median parameter values at each cycle. The default is to omit the first 10% of cycles as a burn-in from the plots.
PMcov	<code>makeCov()</code>	<code>?plot.PMcov</code>	Plots the relationship between any two columns of a PMcov object.

Object Classes	Creating functions	R help	Description
PMmatrix	PMreadMatrix()	?plot.PMmatrix	Plots raw time-observation data from a data .csv file read by the PMreadMatrix() command, with a variety of options, including joining observations with line segments, including doses, overlaying plots for all subjects or separating them, including individual posterior predictions ( <b>post</b> objects as described above), color coding according to groups and more.
PMsim	SIMParse()	?plot.PMsim	Plots simulated time-concentration profiles overlaid as individual curves or summarized by customizable quantiles (e.g. 5th, 25th, 50th, 75th and 95th percentiles). Inclusion of observations in a population can be used to return a visual and numerical predictive check.
PMnpde	makeNPDE()	?plot.PMnpde	<p>Plots an NPDE qqnorm, NPDE histogram, NPDE vs. time, NPDE vs. prediction and others to visualize results of simulation based internal model diagnostics accessed with the makeNPDE() command.</p> <p>More documentation is available at <a href="http://www.biostat.fr/NPDE/index.php">http://www.biostat.fr/NPDE/index.php</a>.</p>
PMpta	makePTA()	?plot.PMpta	Plots superimposed curves corresponding to each dose, with target (e.g. MIC) on the x-axis and proportion of the simulated time-concentration profiles for the dose with a target statistic (e.g. %time > MIC) above a user-defined success threshold

## Examples of Pmetrics plots

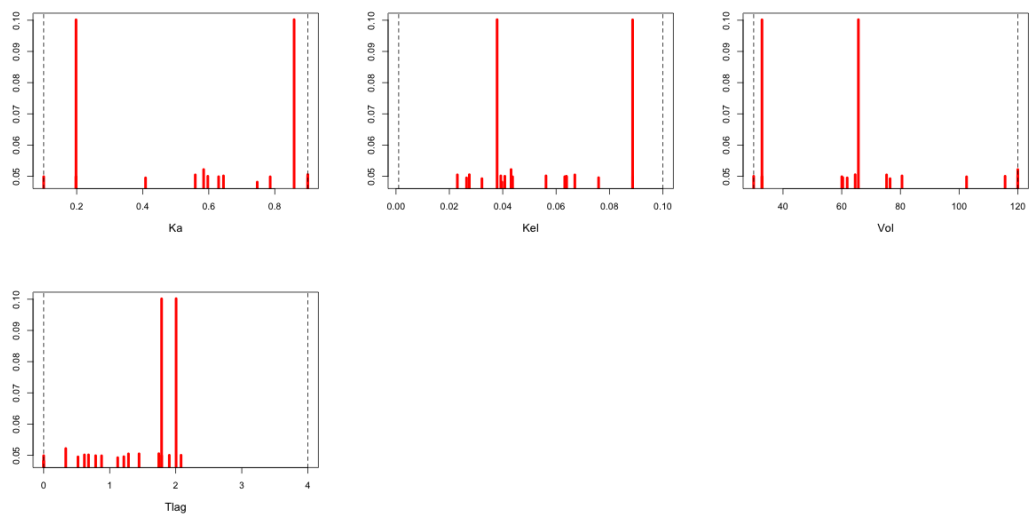


**plot(PMmatrix object)**

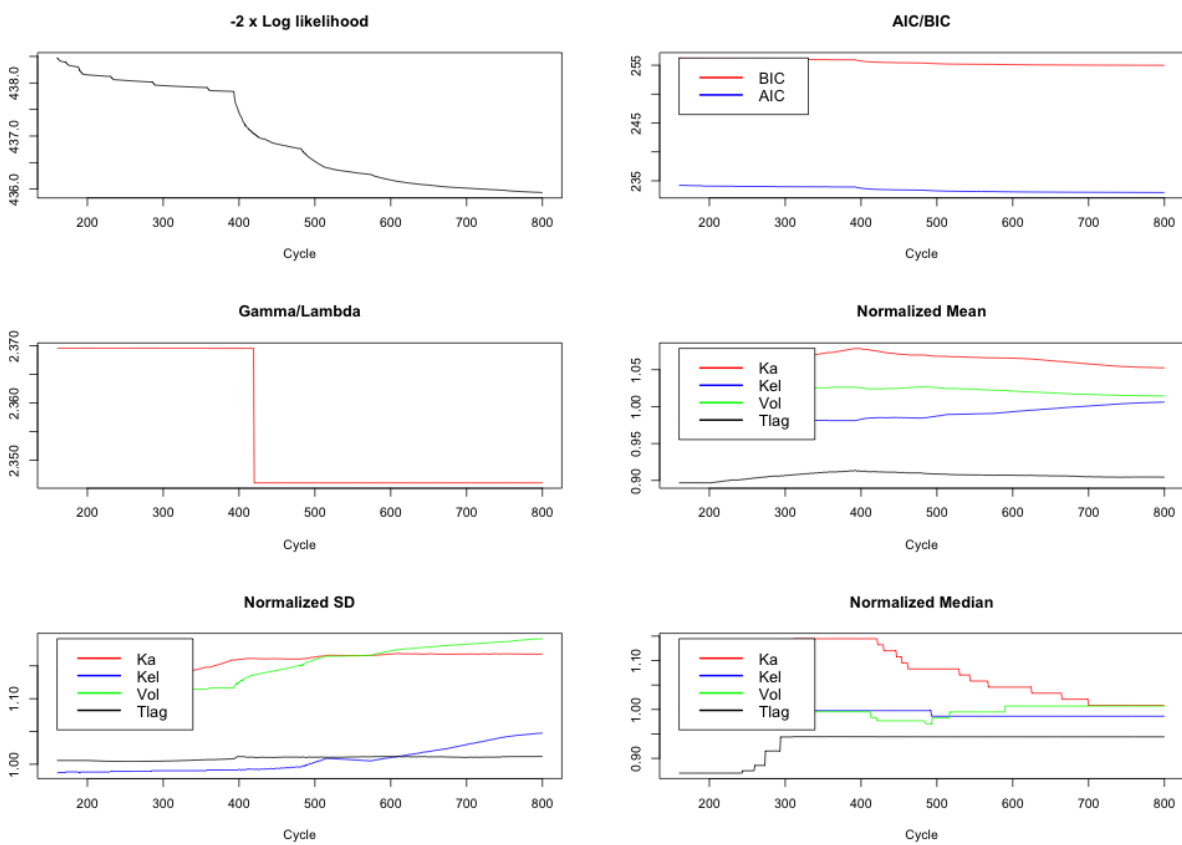


**plot(PMop object)**

**plot(PMop object, resid=T)**

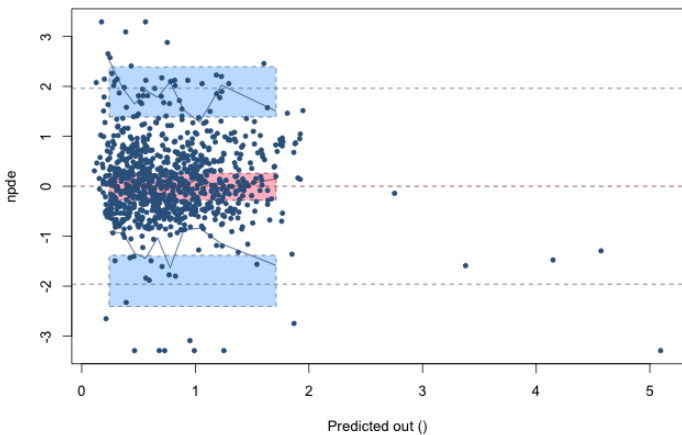
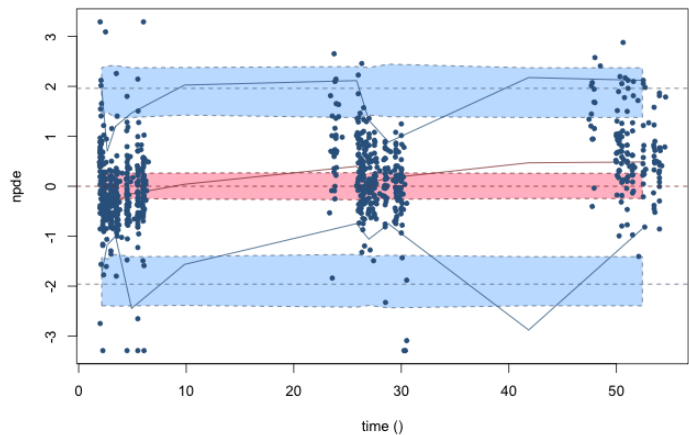
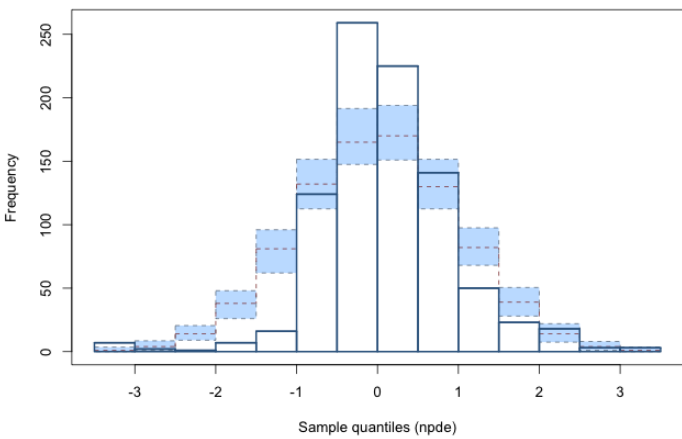
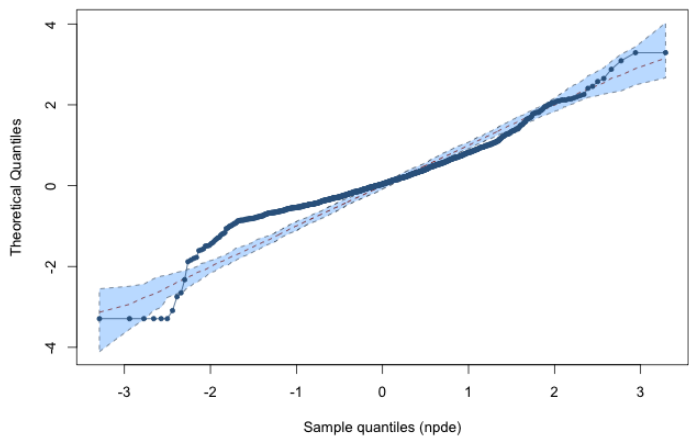


**plot(PMfinal object)**

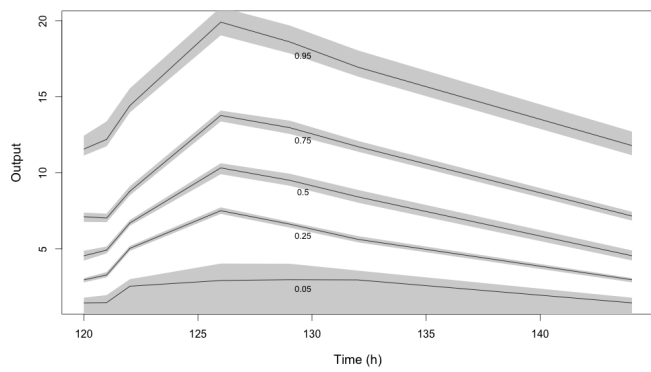


**plot(PMcycle object)**

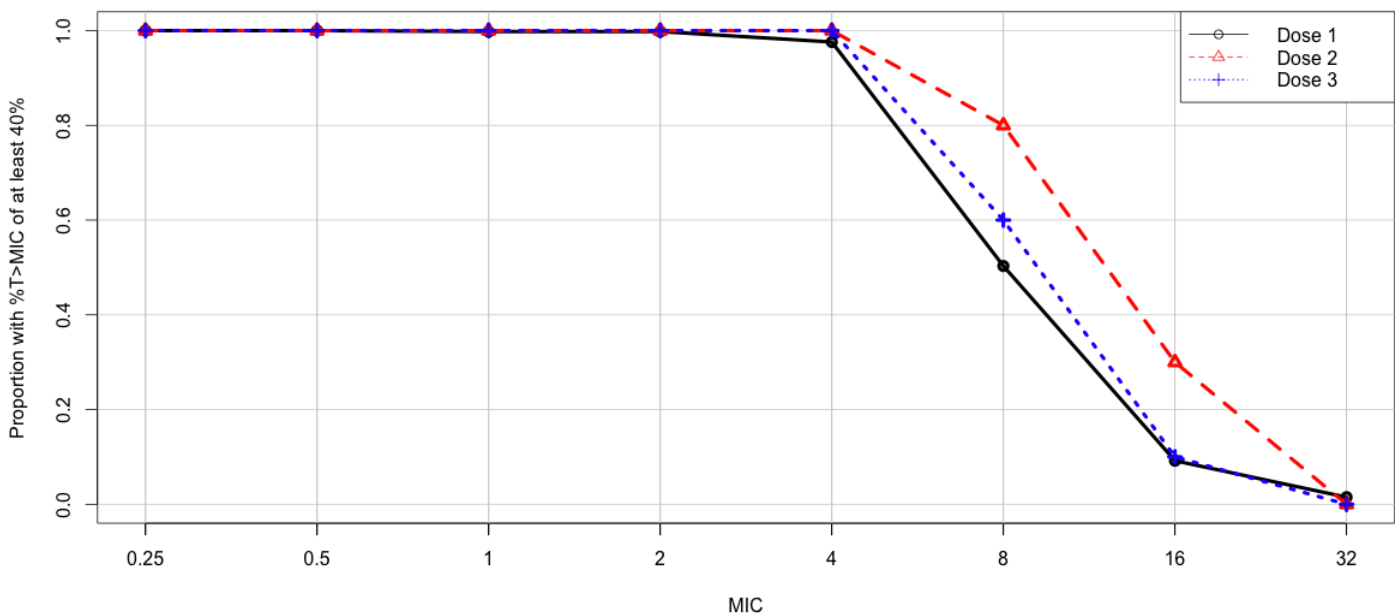
Q-Q plot versus  $N(0,1)$  for npde



**plot(PMnpde)**



**plot(PMsim object)**



**plot(PMpta object)**

## Probability of Target Attainment

PTA analyses and plots are a powerful application of simulation in Pmetrics. Simulated output profiles, e.g time-concentration profiles, are automatically compared to a target to generate a pharmacodynamic index (PDI). This index can be one of several, such as the proportion of a dosing interval that the concentration remains above the target, the ratio of area under the time-concentration curve to target, ratio of maximum or minimum concentration to target, or ratio of concentration at a specific time to target.

Here are the details of the `makePTA()` function.

```
makePTA(simdata, simlabels, targets, target.type, success, outeq = 1,
       free.fraction = 1, start, end)
```

1. **simdata** A vector of simulator output filenames, e.g. `c("simout1.txt","simout2.txt")`, with wildcard support, e.g. `"simout*"` or `"simout?"`, or a list of PMsim objects made by [SIMparse](#) with suitable simulated regimens and observations, e.g. `simdata <- SIMparse("simout?.txt")`. The latter method is preferable so that you do not have to re-parse the simulator output each time you run `makePTA()`. The number and times of simulated observations does not have to be the same in all objects.
2. **simlabels** Optional character vector of labels for each simulation. Default is `c("Regimen 1", "Regimen 2", ...)`. These will be used in plots.
3. **targets** A vector of pharmacodynamic targets, such as Minimum Inhibitory Concentrations (MICs). This can also be a sampled distribution using `makePTATarget()`, which generates an object of class **PMpta.targ** that can be used in `makePTA()`. The single input to `makePTATarget()` is a data.frame or name of .csv file in working directory whose first two columns are targets and the number of samples for each target. An example can be seen for *Staphylococcus aureus* susceptibility to vancomycin at the EUCAST website at <http://mic.eucast.org/Eucast2/regShow.jsp?Id=1214> (see the bottom of the page). Examples:

`makePTA(..., targets=c(0.25, 0.5, 1, 2, 4, 8, 16, 32))` will test all profiles against the specific targets listed. On the other hand, `makePTA(..., targets=makePTATarget("staph.csv"))` will test each profile against one MIC drawn from a distribution of MICs for contained in the `staph.csv` file.

4. **target.type** A numeric or character vector, length 1.

- If numeric, must correspond to an observation time common to all PMsim objects in simdata, rounded to the nearest hour. In this case, the target statistic will be the ratio of observation at time `target.type` to target. This enables testing of a specific timed concentration (e.g. one hour after a dose or C1) which may be called a peak, but is not actually the maximum drug concentration. Be sure that the time in the simulated data is used, e.g. 122 after a dose given at 120.
- “time”, calculate the proportion of the time range specified by start and end time above target. The algorithm looks at each pair of concentrations within a simulate profile, and if both are below the target, the cumulative time is not incremented. If both are above, the cumulative time is incremented by the time interval between the pair. If one is above and the other below, the cumulative time is incremented by the fraction above the target, as estimated by linear regression between the paired concentrations.
- “auc”, calculate ratio of area under the curve within the start-end time range to target, using the trapezoidal approximation in `makeAUC()` to calculate AUC.
- “peak”, calculate ratio of peak concentration within the start-end time range to target
- “min”, calculate the ratio of minimum concentration within the start-end time range to target.

5. **success** A single value specifying the success statistic, e.g. 0.4 for proportion time (start to end) above target, or 100 for peak:target.

6. **outeq** An integer specifying the number of the simulated output equation to use. Default is 1.

7. **free.fraction** Proportion of free, active drug, from 0 to 1. Default is 1, i.e. 100% free drug or 0% protein binding.

8. **start** Specify the time to begin PTA calculations. Default is a vector with the first observation time for subjects in each element of simdata, e.g. dose regimen. If specified as a vector, values will be recycled as necessary.

9. **end** Specify the time to end PTA calculations so that PTA is calculated from start to end. Default for end is the maximum observation time for subjects in each element of simdata, e.g. dose regimen. If specified as a vector, values will be recycled as necessary. Subjects with insufficient data (fewer than 5 simulated observations) for a specified interval will trigger a warning. Ideally then, the simulated dataset should contain sufficient observations within the interval specified by start and end.

If a simulation exists, such that `simout1.txt`, ..., `simout4.txt` exist in the current working directory, and each contain a different simulated dosing regimen of 100 mg daily, 50 mg bid (twice daily), 200 mg daily, or 100 mg bid, all given for 5 days, then here are examples of different PTAs.

Example 1:

```
#parse the results into a list
simlist <- SIMparse("simout?.txt")

#make the PTA with discrete targets, and define success as proportion > 0.6 of
interval from 72 to 96 hours with concentration > target

pta.1 <- makePTA(simdata=simlist, targets=c(0.25, 0.5, 1, 2, 4, 8, 16),
target.type="time", success=0.6, start=72, end=96)
```

Example 2:

```
#parse the results into a list
simlist <- SIMparse("simout?.txt")
```

```
#make the PTA with sampled targets using a file with the distribution of
vancomycin MICs for staphylococcus aureus, and define success as an AUC:target ≥
400 from 72 to 96 hours

pta.1 <- makePTA(simdata=simlist, targets=makePTATarget("staph.csv"),
target.type="auc", success=400, start=72, end=96)
```

### Example 3:

```
#parse the results into a list
simlist <- SIMparse("simout?.txt")

#make the PTA with a target concentration of 4 mg/L 12 hours after (e.g. time=84)
a steady state dose given every 12 hours

pta.1 <- makePTA(simdata=simlist, targets=4, target.type=84, success=1)
```

The output of `makePTA()` is a list of class **PMpta**, which has 2 objects:

1. **results** A data frame with the following columns:

- *simnum* is the number of the simulation;
- *id* is the simulated profile number within each simulation;
- *target* is the specified target;
- *pdi* is the target pharmacodynamic index, e.g. time > target, auc:target, etc.

2. **outcome** A data frame summarizing the results with the following columns:

- *simnum* and *target* are as for results. If *targets* was specified via `makePTATarget()` to be a sampled distribution, then the *target* column will be missing from the outcome table.
- *prop.success* column has the proportion with a *pdi* > *success*, as specified in the function call.
- The *pdi.mean* and *pdi.sd* columns have the mean and standard deviation of the target pharmacodynamic index (e.g. proportion end-start above target, ratio of Cmax to target) for each simulation and target.

**PMpta** objects can be summarized with `summary(x)` and plotted with `plot(x)`. See `?summary.PMpta` and `?plot.PMpta` in R for additional help.

## Model Diagnostics

### Internal Validation

Several tools are available in Pmetrics to assist with model selection. The simplest methods are using `PMcompare()` and `plot.PMop()`, via the `plot()` command for a PMop object made by `makeOP()` or by using `PMload()` after a successful run. `PMstep()` is another option for covariate analysis. All these functions are carefully documented within R and accessible using the `?command` or `help(command)` syntax.

**Plotting observed vs. predicted objects (class: PMop).** When you have successfully completed an NPAG or IT2B run and loaded the results with `PMload()`, one of the objects will be a PMop object. *Example:* `PMload(1)` will include **op.1** in the list of loaded objects. You can also remake the PMop object, for example, if you want use the mean of parameter values for calculation of predictions rather than the default median: `op.1 <- makeOP(NPdata.1, "mean")`. In both cases, you can then simply `plot(op.1)`, which is a wrapper for `plot.PMop(op.1)`. See `?plot.PMop` for options to this plot.

One particularl option to `plot.PMop()`, `resid=T` will generate a residual plot instead of an observed vs. predicted plot. A residual plot consists of three panels: 1) weighted residuals (predicted - observed) vs. time; 2)

weighted residuals vs. predictions; 3) a histogram of residuals with a superimposed normal curve if the option `ref=T` is specified (the default). The mean of the weighted residuals (expected to be 0) is reported along with the probability that it is different from 0 by chance. Three tests of normality are reported for the residuals: D'Agostino[2], Shapiro-Wilk, and Kolmogorov-Smirnof. An example is shown in the [Plotting](#) section.

**Covariate analysis.** The `PMstep()` function will report a 2x2 matrix of P-values for the linear regression coefficients of each subject's covariates vs. Bayesian posterior parameter values from the `PMcov` object loaded with `PMload()`. *Example:* `PMstep(cov.1)`. Entries in the matrix are the multivariate P-values. A value of NA indicates that the variable was not retained in the final model. The default method is "both", but "forward" selection and "backward" elimination are possible. `PMstep()` uses the `step()` function in the `stats` package for R, which is a default package. It is possible to test non-linear relationships using capabilities of R and the `PMcov` object, for example with the `nls()` function for non-linear least squares analysis.

**Model comparison.** To compare models with `PMcompare()`, simply enter two or more PMetrics data objects, e.g. `PMcompare(NPdata.1, NPdata.2, NPdata.3)`. These should be of the `NPAG` or `IT2B` class, made either by using `PMload()` or `NPParse()`/`ITParse()`. Although it is possible to compare models of mixed classes, the validity of this is dubious. The return object will be a data frame with summaries of each model and key metrics such as log-likelihood, final-cycle Akaike Information and Bayesian Information Criteria, bias and imprecision of predictions relative to observations from the population prior distribution and individual posterior distributions, and a p-value for model.

The p-values are a comparison of the joint distribution of population parameter values for all models, using the first model in the list as the reference. Only parameters with common names to all models are included. Data objects supplied to `PMcompare()` should not be zero-cycle runs, which will generate an error message. By specifying the option `plot=T`, observed vs. predicted plots for all the models will be generated. The comparison is based on the nearest neighbors approach in the `MTSKNN` package. The option to generate residual plots of prediction errors, described next, can be specified with the additional switch `resid=T`, which is ignored if `plot=F`.

**Model bias and imprecision.** In both `PMcompare()` and in plots of `PMop` objects with `reg=T` (the default), PMetrics supplies two statistics: bias and imprecision. Bias is the mean weighted error of predicted - observed. Imprecision is the bias-adjusted, mean weighted squared error of predicted - observed. We adjust for bias because the mean squared error (MSE), calculated as  $(\text{pred} - \text{obs})^2$  for all predictions/observations, is equal to the sum of the variance of the predictions and the bias of the predictions, i.e.  $\text{Var}(\text{pred}) + \text{Bias}(\text{pred})$ . Therefore, the true variance or imprecision of the predictions is  $\text{MSE} - \text{Bias}$ .

For both calculations, the weighting is according to the calculated SD of the observation, using C0, C1, C2, and C3. The specific formulae are as follows.

- **Weighted prediction error, *wpe*:**  $(\text{pred}-\text{obs})/\text{SD}$  for each prediction/observation
- **Weighted squared prediction error, *wspe*:**  $(\text{pred}-\text{obs})^2/\text{SD}^2$  for each prediction/observation
- **Bias, mean weighted prediction error, *mwpe*:**  $\sum \text{wpe} / N$
- **Imprecision, bias-adjusted mean weighted squared prediction error, *bamwspe*:**  $\sum \text{wspe} / N - \text{mwpe}^2$

All these values are returned with `summary()` for `PMop` objects. Both `summary()` and `PMcompare()` will also provide the root mean squared error (RMSE) and %RMSE. RMSE is the square root of  $(\text{pred}-\text{obs})^2$ , i.e. an unweighted measure of imprecision used by many, but which we favor less given its failure to account for observation weights or adjust for bias. The %RMSE is the RMSE normalized to the total RMSE over all observations.

## Visual predictive checks and NPDE.

Two more complex and time-consuming options based on simulation are also available for internal model validation: the **normalized prediction distribution errors (NPDE)** method of Mentré and Escolano [3] and **visual predictive checks (VPC)**. Both of these can be computed from the same simulation.

For NPDE, the basic idea is that each subject in the population serves as a template for a simulation of 1000 further profiles using the population structural model and parameter values joint probability distribution, i.e. together the "population model". The simulated profiles are compared to the observed data, and the NPDE is generated. The command to generate a `PMnpde` list object is `makeNPDE()`, which is documented in R. Your current working

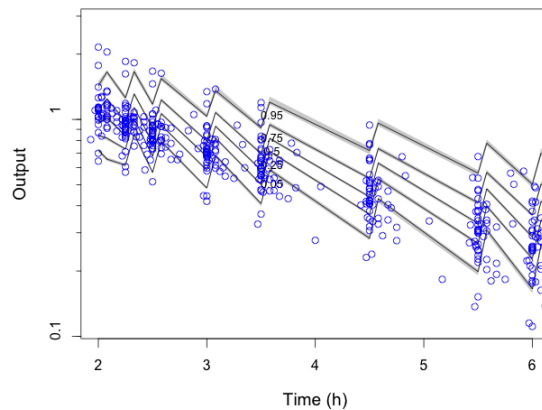
directory should be your runs folder, as the argument to `makeNPDE()` is the run (folder) number that you wish to analyze. Because of the extensive simulations involved, execution of this command can be slow if the population is large, the model complex, the time horizon long, and/or the number of observations to be simulated per profile is large. There are print and plot methods for `NpdeObjects`, which are contained within `PMnpde` objects (`print.NpdeObject()` and `plot.NpdeObject`) both of which are also documented within R and much more extensively online at <http://www.biostat.fr/NPDE/index.php>. An example of a NPDE plot is shown in the [Plotting](#) section.

Note that simulation from a population model can be a fickle thing, which may lead to errors when trying to execute this command. Parameter value distributions in linear space run the risk of simulating extreme or even inappropriately negative parameter values which can in turn lead to simulated observations far beyond anything corresponding to possible reality. Arguments to the Pmetrics simulator can be supplied in the `makeNPDE()` call, such as **split**, and **limits** to mitigate this problem.

In Pmetrics `makeNPDE()` automatically generates the simulations necessary for a **VPC**. VPCs are cumbersome when models include covariates or have heterogeneous dosing/sampling regimens among subjects in the population. It is nonetheless possible to obtain a VPC in two ways.

The first way to generate a VPC is to use the **PMsim** object saved after running `makeNPDE()`, which will be called **sim.x**, where **x** is the run number. If an observed vs. predicted `PMop` object made with `makeOP()` is passed to `plot.PMsim()` with the `obs` argument, the observed values will be overlaid upon simulated profiles if possible, and an NPC will be returned in addition to the plot. The NPC is simply a binomial test for the percentage of observations less than the quantiles specified by the `probs` argument (0.05, 0.25, 0.5, 0.75, 0.95 by default). It is up to the user to decide if the study population and model is homogeneous enough to justify a VPC.

*Example:* `vpc <- plot(sim.1, obs=op.1)`. Note that when we use the `obs` argument, the `plot` call will return a list of, in addition to a plot like the one below. Although the template subjects all received only one dose of drug in this interval, they were not sampled at the same times, so when the simulations were combined, it leads

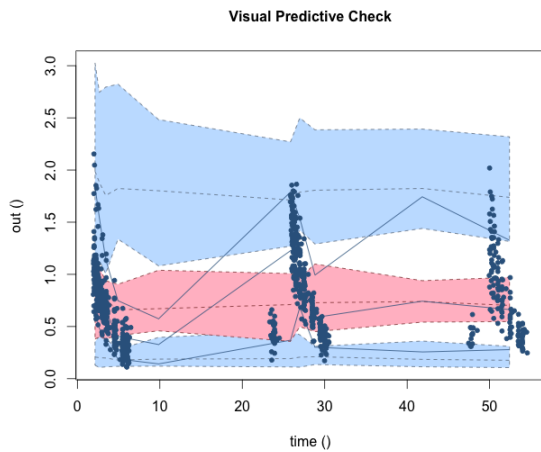


to the jagged quantiles. This can be improved (i.e. smooth the quantile curve) by setting the `binSize` argument when plotting, which controls the width of binning interval for simulated concentrations, in time units, e.g. hours. For example, a `binSize` of 0.5 will pull all simulated concentrations +/- 0.5 hours into the same time bin. The default is 0, i.e. no binning. The return object when **obs** is included is a list with the following items:

1. **npc** A data frame with three columns: `quantile`, `prop.less`, `pval`. `quantile` are those specified by the `prob` argument to the `plot` call; `prop.less` are the proportion of simulated observations at all times less than the quantile; `pval` is the P-value of the difference in the `prop.less` and `quantile` by the beta-binomial test.
2. **simsum** A data frame with the quantile concentration at each simulated time, with lower and upper confidence intervals
3. **obs** A data frame similar to an `PMop` object made by `makeOP()` with the addition of the quantile for each observation

So to get the NPC, simply type, from the example above, `vpc$npc`.

Visual predictive checks can also be made from a `PMnpde` object made with the `makeNPDE()` function with the following syntax: `plot(npde.1, plot.type="vpc")`. An example of this plot is shown below.



## External Validation

Should you wish to use your population model to test how well it predicts a second population that is separate from that used to build the model (i.e. externally validate your model) you may do that in Pmetrics. After completing an NPAG run, place the same model file (located in the `inputs` subdirectory of the NPAG run whose model you are validating) along with your new (validating) data file in your working directory. So there should be two files in your working directory:

- **model .txt file** This will be the same as for model building NPAG run, found in the `/inputs` subdirectory.
- **data .csv file** This will be a Pmetrics data input file containing the new subjects for validation.

Next, do the following steps to complete the validation NPAG run.

1. Load the model building run with `PMload(run_num1)` so that its `NPdata` object is in memory.
2. Initiate an NPAG run in Pmetrics as usual, but with an additional argument to specify the model density file which will serve as a non-uniform prior, e.g. `NPrun(model="mymodel.txt", data="validation.csv", prior=NPdata.1, cycles=0)`, where `NPdata.1` is an example of the object loaded in step 1, in this case with `PMload(1)`. Specifying 0 cycles will calculate a Bayesian posterior only for each subject in the validation data set.
3. Complete the NPAG run as usual.
4. Load the results with `PMload(run_num2)` and plot, etc. as usual.

## Pmetrics Outputs

At the end of a successful NPAG, IT2B or ERR run, your Run folder will have a new folder with the run number, which is sequentially designated from the previous run. Within this folder are 4 subfolders.

- **/etc** This folder contains files made during the run. Generally you do not need to look at these. They include the fortran code specifying the model (model.for) and the instructions (instr.inx) that NPAG or IT2B used.
- **/inputs** This folder contains the original data and model file used in the run.
- **/outputs** This folder contains the NPAGout.Rdata or IT2B.Rdata file (see Data Objects) which are created by the PMreport() function, as well as the NPAGreport.html or IT2Breport.html files, which you will see open automatically in your default browser at the end of a run. This html generates a tabbed page which will display properly as long as it is in the run /outputs folder where the supporting image files are also located. Within this folder are also .csv (comma separated values) tables of the population parameter value summaries (popparam.csv), covariance matrix (popcov.csv), correlation matrix (popcor.csv), and points (NPAG only, poppoints.csv). Images for default graphs are available as .png and .pdf files. Additional text files for NPAG (OUT0001, OUTT0001, DEN0001, ILOG0001, PRTB0001, NP\_RF0001.txt) contain combined cycle output/density, cycle output, density, convergence, posterior predictions, and R-friendly information, respectively. For IT2B, these text files (OUTF0001, OUFF0001, DENF0001, LAST0001, FROM0001, IT\_RF0001.txt) contain combined cycle output/density, cycle output, density, Bayesian posterior parameters, parameter ranges that can be passed to NPAG, and R-friendly information, respectively. Normally, you do not need to look at these files.

Finally, for outputs, there is an NPAGreport.tex or IT2Breport.tex file. For users familiar with LATEX, this file can be used to generate a pdf version of the html report page, useful for sharing with colleagues. You must have an installed LATEXengine. Our recommended LATEX engine for Mac users is [MacTex](#), which is very large (>1GB) but contains a complete installation. For Windows, [MikTex](#) is our recommended engine. Both are free.

Once you have a LATEX engine, it is easy to make a .pdf from the .tex file. Simply open it with Rstudio and click the “Make PDF” button that appears. The .pdf file will be in your /outputs folder.

- **/wrkcopy** This folder contains “working copy” files in the original format used by NPAG and IT2B, and which are still used “under the hood.” They are easy to read and can be used as a check if something goes wrong.

## Optimal Sampling

Pmetrics has two optimal sampling functions which can be used on the output of NPAG runs only (currently).

- **MMopt** or multiple model optimal sampling. This function uses the simulator, a template of candidate dosage regimens and sample times, and the PMfinal object from an NPAG run to generate the expected output from each support point in the model. The algorithm then finds the requested number of times (one to four) which minimize the chance of choosing the incorrect Bayesian posterior for a subject or group of subjects. This is also termed minimizing the maximum possible Bayesian risk of misclassification. In other words, for the least number of samples, how can you be the most certain that your Bayesian posterior parameter value distribution for an individual is the most likely to be “correct”? The algorithm finds the optimal times by locating the points of maximal separation of the output profiles of all the points in the model. The template data file may contain multiple regimens, in which case each regimen will be administered to each point in the model, generating *npoint x nregimen* output profiles, upon which the MMopt calculation will be performed.

There is an option for a weighted MMopt, currently only for AUC. This version of the optimization does not try to distinguish support points that generate the same AUC, regardless of other values, such as maximum or minimum output. This can be useful if the target for pharmacodynamic effect is an AUC rather than a specific concentration.

To use the function, see the help in Pmetrics under ?MMopt.

- **Dopt** This function will calculate d-optimal times from the output of an NPAG run. D-optimal sample times are chosen such that the smallest changes in parameter values result in the largest changes in output profile. Be sure that the working directory is the project /Runs folder, and then use Dopt(run), e.g. Dopt(2) to calculate the d-optimal times for that run. If you have no template data file in the /Runs folder, the first subject in the original run will be used as the template for dosage and starting sample times. However, if the number of observations in that subject exceeds 10, then 10 observation times will be randomly selected to serve as the starting times. If you include a template data file, you may specify any dosage regimen and candidate starting

times (up to 10 and at least as many as parameters) that you wish.

To use the function, see the help in Pmetrics under ?Dopt.

## Ancillary Functions

Pmetrics has some ancillary functions which can be helpful when you are involved in population PK modeling. These functions do not control the components of Pmetrics or process output.

- **qgrowth** This function will return a dataframe of height and weight for boys, girls or both for a given range of ages in months and body size percentile (e.g. the median). This can be useful for simulations in Pmetrics. The data for this function come from the 1977 National Center for Health Statistics Growth Chart Equations on the website of the United States [Centers for Disease Control](#).
- **ss.PK** Sample size calculations for PK studies are difficult as traditional power analysis is based on comparisons between 2 or more groups. One proposed method is to sample sufficient subjects to ensure that the width or precision of the standard error of the mean value for a given PK parameter is within a specified degree for a given probability. This function uses the formula for precision of the standard error of the mean of a distribution of PK parameter values to calculate either  $n$ , the sample size, or  $sd$ , the maximum standard deviation of a PK parameter value distribution for a given mean, desired degree of precision and confidence.

The formula is  $n = Z^2_{(1+ci)/2} * sd^2 / (\text{precision} * \text{mean})^2$ , where  $Z$  is the standard normal quantile,  $ci$  is the probability of the confidence interval (e.g. 0.95),  $sd$  is the standard deviation of the parameter values, *precision* is the desired width of the confidence interval as a proportion of the mean (e.g. 0.2 = 20%), and *mean* is the mean parameter value.

## References

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3. Mentré F, Escolano S. Prediction discrepancies for the evaluation of nonlinear mixed-effects models. *J Pharmacokinet Pharmacodyn* **2006**; 33:345–367.