# Beautiful graphs with npde 3.0September 2020

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npde website: www.npde.biostat.fr

# 1 Running npde

```
Describe: - viral load data example - 20%
   Viral load data with 20% censored data
   [1] "computenpde.log<-function(npdeObject) {"
   _____
   Distribution of npde :
       nb of obs: 300
           mean = -0.03127 (SE= 0.053)
        variance= 0.8358 (SE= 0.068)
        skewness= 0.04592
        kurtosis= -0.3856
   _____
   Statistical tests
    t-test : 0.554
Fisher variance test : 0.0355 *
    SW test of normality
                          : 0.499
   Global adjusted p-value
                         : 0.107
   Signif. codes: '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1
   Warfarin PK data
       npde for the model without covariates
   _____
   Distribution of npde :
       nb of obs: 247
           mean= 0.03419 (SE= 0.06)
        variance= 0.8753 (SE= 0.079 )
        skewness= -0.1149
        kurtosis= -0.0497
   _____
   Statistical tests
    t-test
                        : 0.566
                       : 0.157
    Fisher variance test
    SW test of normality
                          : 0.371
   Global adjusted p-value
                         : 0.471
   Signif. codes: '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1
   _____
       npde for the model with covariates
   _____
   Distribution of npde :
       nb of obs: 247
           mean= 0.02928 (SE= 0.059 )
        variance= 0.8549 (SE= 0.077)
        skewness= -0.07211
        kurtosis= -0.4172
```

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Statistical tests

t-test : 0.619

Fisher variance test : 0.096 .

SW test of normality : 0.368

Global adjusted p-value : 0.288
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Signif. codes: '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1
```

The npde package has built-in diagnostic plots which can be accessed for an object y returned by autonpde() (or npde() for the interactive execution) by the usual R command plot():

plot(y)

> plot(yvir20)

produces the plot shown in figure 1.

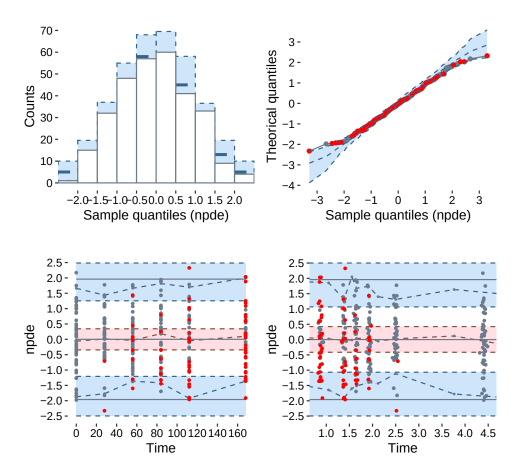


Figure 1: Default plots for the viral load data.

Eco TODO problems with the viral load data plots:

- not the same result when I execute interactively the viral load data (the p-value is NS interactively, but sometimes comes out significant)
- check the nubmer of simulations, maybe not enough ?

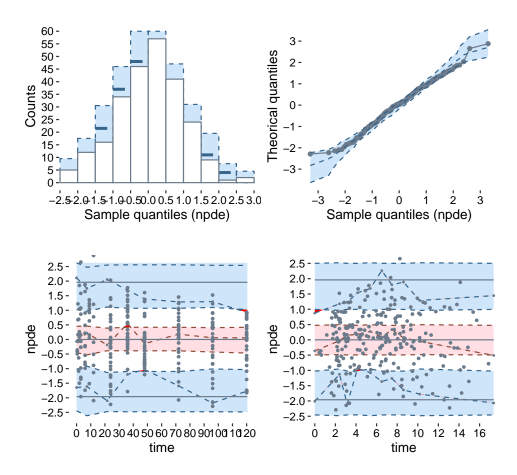


Figure 2: Default plots for the warfarin PK data, assuming a model without covariates.

# 2 Graphical options

### 2.1 Types of plots

Each of the default diagnostic plots, as well as a number of additional plots not shown by default, can also be produced on its own, using the argument plot.type="type". Table ?? lists the plots that can be created in this way. Apart from VPC and the plot of the probability of an observation being under the LOQ, all plots can be obtained for npde, npdor pd(this is controlled by the which="" argument, defaulting to which="npde").

Plot type	Description
data	Plots the observed data in the dataset
hist	Histogram of the npde
qqplot	QQ-plot of the npdeversus its theoretical distribution
ecdf	Empirical distribution function of the npde
x.scatter	Scatterplot of the npde versus the predictor X
pred.scatter	Scatterplot of the npde versus the population predicted values
cov.scatter	Scatterplot of the npde versus covariates
vpc	Plots a Visual Predictive Check (VPC)
loq	Plots the probability for an observation to be BQL, versus the
	predictor X

Table 1: Plot types available in the npde library. QQ-plots, histograms, cumulative cdf, and scatter plots can be produced for npde, pd or npd.

Table 2: tab:plottypes

To produce only the QQ-plot as a standalone graph, we request a "qqplot":

The plots can also be produced by the other metrics computed by npde(), for instance we might want to consider the histogram of pdwith the following command:

# > plot(yvir20, plot.type="qqplot")

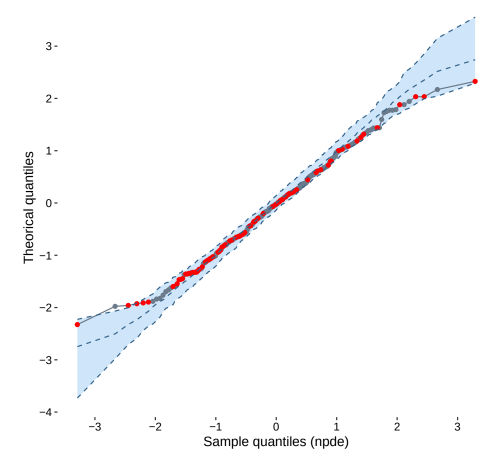


Figure 3: QQ-plot for the viral load data.

> plot(yvir20, plot.type="hist",which="pd")

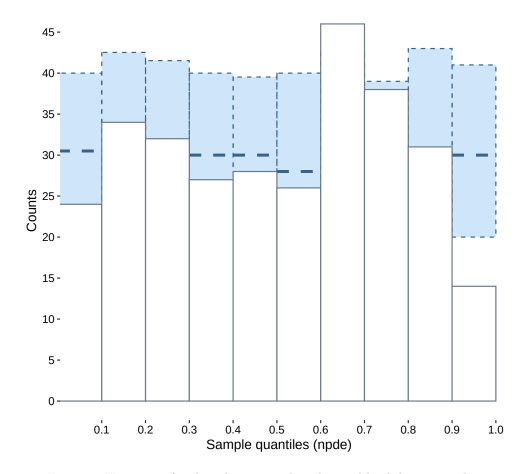


Figure 4: Histogram for the pd, computed in the viral load data example.

# 2.2 Changing graphical parameters

TODO: show different features in group, eg colours/symbols, titles/axes, boxes/grids Decide on following options: verbose

Argument	Description	Default value
verbose	Output is produced for some plots (most notably	FALSE
	when binning is used, this prints out the boundaries	
	of the binning intervals) if TRUE	
main	Title	depends on plot
sub	Subtitle	empty
size.main	Size of the main title 14	
size.sub	Size of the title for covariate	12
xlab	Label for the X-axis	depends on plot
ylab	Label for the Y-axis	depends on plot
size.xlab	Size of the label for the X-axis	12
size.ylab	Size of the label for the Y-axis	12
breaks.x	Number of tick marks on the X-axis	10
breaks.y	Number of tick marks on the Y-axis	10
size.x.text	Size of tick marks and tick labels on the X-axis	10
size.y.text	Size of tick marks and tick labels on the Y-axis	10
xlim	Range of values on the X-axis	empty, adjusts to the data
ylim	Range of values on the Y-axis	empty, adjusts to the data
xaxt	A character whether to plot the X axis. Specifying	"y"
	"n" suppresses plotting of the axis	
yaxt	A character whether to plot the Y axis. Specifying	"y"
	"n" suppresses plotting of the axis	
xlog	Scale for the X-axis (TRUE: logarithmic scale)	FALSE
ylog	Scale for the Y-axis (TRUE: logarithmic scale)	FALSE

Table 3: Graphical parameters that can be passed on the plot function: titles and axes.

Argument	Description	Default value
col	Main colour for observed data (applied to lines and	"slategray4"
	symbols pertaining to observations if no other option	
	is given to supersede this value)	
lty	Line type for observed data	1
lwd	Line width for observed data	0.5
pch	Symbol used to plot observed data	20
alpha	Transparencyfor observed data	1
size	Symbol size to plot observed data	1
fill	Colour used to fill area elements related to observed	"white
	data (such as histogram bars)	

Table 4: Graphical parameters that can be passed on the plot function: colours and symbols.

#### 2.3 Saving to file

#### 2.3.1 Using ggsave from ggplot

#### 2.3.2 Handling transparency with postscript files

Postcript doesn't handle the transparency used in the npde plots made with the ggplot2 library. Using ggsave(), the prediction intervals will likely be missing from the output. A workaround is to output the files to PDF format. Another is to use the following code

TODO

#### 2.4 Arranging individual plots

# 3 Covariate plots

#### 3.1 Plots of npde versus covariates

#### 3.2 Stratified plots

The different diagnostic plots produced in section 2.1 can be stratified using the covsplit=TRUE option. By default, the plots will be stratified for each covariate separately. The plots produced depend on the nature of the covariate, following [?]:

- for categorical covariates, a plot is created for each category of the covariate
- for continuous covariates, three plots

# 4 Reference profile