Package 'npde'

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Title Normalised Prediction Distribution Errors for Nonlinear Mixed-Effect Models

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Description

Provides routines to compute normalised prediction distribution errors, a metric designed to evaluate non-linear mixed effect models such as those used in pharmacokinetics and pharmacodynamics. Documentation about 'npde' is provided by a comprehensive user guide on the github repository (https://github.com/ecomets/npde30/blob/main/userguide_npde_3.0.pdf), and references concerning the methods include the papers by Brendel et al. (2006, <doi:10.1007/s11095-006-9067-5>; 2010, <doi:10.1007/s10928-009-9143-

7>), Comets et al. (2008, <doi:10.1016/j.cmpb.2007.12.002>; 2010, http://journal-sfds.fr/article/view/45), and Nguyen et al. (2012, <doi:10.1007/s10928-012-9264-2>). See 'citation(``npde")' for details.

URL https://github.com/ecomets/npde30/blob/main/userguide_npde_3.0.pdf

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LazyLoad yes

LazyData yes

Depends gridExtra, ggplot2, grid

Imports methods, scales, mclust, rlang

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Collate 'NpdeSimData.R' 'NpdeData.R' 'aaa_generics.R'

'NpdeData-methods.R' 'NpdeRes.R' 'NpdeRes-methods.R'

'NpdeObject.R' 'NpdeObject-methods.R' 'compute_distribution.R'

'compute_npde.R' 'compute_pd.R' 'compute_ploq.R' 'mainNpde.R'

'npde.R' 'npdeControl.R' 'plotNpde-auxDistPlot.R'

'plotNpde-auxScatter.R' 'plotNpde-auxScatterPlot.R'

'plotNpde-binningPI.R' 'plotNpde-covplot.R'

'plotNpde-distributionPlot.R' 'plotNpde-methods.R'

'plotNpde-plotFunctions.R' 'plotNpde-scatterplot.R'

NeedsCompilation no

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Description

Routines to compute normalised prediction distribution errors, a metric designed to evaluate non-linear mixed effect models such as those used in pharmacokinetics and pharmacodynamics

Author(s)

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References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentr\'e. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

PDF documentation for npde 3.0: $https://github.com/ecomets/npde 30/blob/main/userguide_npde_3.0.pdf$

```
data(theopp)
data(simtheopp)

# Calling autonpde with dataframes

x<-autonpde(theopp,simtheopp,ix="Time",iy="Conc",iid="ID",boolsave=FALSE)
print(x)</pre>
```

4 autonpde

autonpde	Compute normalised prediction distribution errors	
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Description

These functions compute normalised prediction distribution errors (npde) and prediction discrepancies (pd). npde asks the user the name and structure of the files containing the data, using pdemenu, while autonpde takes these variables and others as arguments.

Usage

```
autonpde(namobs, namsim, iid, ix, iy, imdv = 0, icens = 0,
icov = 0, iipred = 0, boolsave = TRUE, namsav = "output", type.graph = "eps",
verbose = FALSE, calc.npde=TRUE, calc.npd=TRUE, decorr.method = "cholesky",
  cens.method = "cdf", units = list(x="",y=""), detect=FALSE, ties=TRUE, header=TRUE)
npde()
```

Arguments

namobs	name of the file containing the observed data, or a dataframe containing the observed data (in both cases, the column containing the various data required for the computation of the pde can be set using the arguments iid,ix and iy below)
namsim	name of the file containing the simulated data, or a dataframe containing the simulated data (the program will assume that subject ID are in column 1 and simulated Y in column 3, see User Guide)
iid	name/number of the column in the observed data containing the patient ID; if missing, the program will attempt to detect a column named id
ix	name/number of the column in the observed data containing the independent variable (X) ; if missing, the program will attempt to detect a column named X
iy	name/number of the column in the observed data containing the dependent variable (Y); if missing, the program will attempt to detect a column with the response
imdv	name/number of the column containing information about missing data (MDV), defaults to 0 (column not present)
icens	name/number of the column containing information about censored data (cens), defaults to 0 (column not present)
icov	name/number of the column(s) containing covariate information defaults to 0 (no covariates)
iipred	name/number of the column(s) with individual predictions (ipred), defaults to 0 (individual predictions not available)

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boolsave	a boolean (TRUE if graphs and results are to be saved to a file, FALSE otherwise), defaults to TRUE
namsav	name of the files to which results are to be saved (defaults to "output", which will produce a file called output.eps (if the default format of postscript is kept, see type.graph) for the graphs and a file called output.npde for the numerical results (see value)
type.graph	type of graph (one of "eps", "jpeg", "png", "pdf"), defaults to postscript ("eps")
verbose	a boolean (TRUE if messages are to be printed as each subject is processed, FALSE otherwise), defaults to FALSE
calc.npde	a boolean (TRUE if npde are to be computed, FALSE otherwise), defaults to TRUE
calc.npd	a boolean (TRUE if npd are to be computed, FALSE otherwise), defaults to TRUE
decorr.method	a character string indicating the method used to decorrelate observed and simulated data in the computation of npde (see npde.decorr.method) defaults to cholesky
cens.method	a character string indicating the method used to handle censored data (see $npde.cens.method$) defaults to cdf
units	a list with components x, y and cov (optional), specifying the units respectively for the predictor (x), the response (y), and the covariates (a vector of length equal to the number of covariates). Units will default to (-) if not given.
detect	a boolean controlling whether automatic recognition of columns in the dataset is on, defaults to FALSE
ties	a boolean (if FALSE, the distributions of pd and npde are smoothed by jittering the values so that there are no ties), defaults to TRUE
header	a boolean (TRUE if input files have headers, FALSE otherwise), defaults to TRUE

Details

Both functions compute the normalised prediction distribution errors (and/or prediction discrepancies) in the same way. npde is an interactive function whereas autonpde takes all required input as arguments.

Diagnostic graphs are produced for npd, and npde are used in the tests as their distribution takes into account the correlation between repeated observations.

When the computation of npde fails because of numerical problems, error messages are printed out, then pd are computed instead and graphs of pd are plotted so that the user may evaluate why the computation failed.

The function also prints out the characteristics of the distribution of the npde (mean, variance, skewness and kurtosis) as well as the results of the statistical tests applied to npde. In addition, if boolsave is TRUE, two files are created:

results file the numerical results are saved in a file with extension .npde (the name of which is given by the user). The file contains the components id, xobs, ypred, npde, pd stored in columns

graph file the graphs are saved to a file with the same name as the results file, and with extension depending on the format.

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Value

An object of class NpdeObject

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

PDF documentation for npde 3.0: https://github.com/ecomets/npde30/blob/main/userguide_npde_3.0.pdf

See Also

```
npde.graphs, gof.test
```

Examples

```
data(theopp)
data(simtheopp)

# Calling autonpde with dataframes
x<-autonpde(theopp,simtheopp,1,3,4,boolsave=FALSE)
x
head(x["results"]["res"])
plot(x)</pre>
```

dist.pred.sim

Compute distribution of pd/npde using simulations

Description

This function is used to build the distribution of pd/npde using the simulations under the model. The default is to build only the distribution of pd, and to sample from N(0,1) when building the distribution of npde under the null hypothesis.

Usage

```
dist.pred.sim(npdeObject,nsamp, ...)
```

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Arguments

npdeObject an object returned by a call to npde or autonpde

nsamp number of datasets (defaults to 100 or to the number of replications if it is

smaller)

... additional arguments. Currently only the value of calc.npd and calc.npde may

be passed on, and will override their corresponding value in the "options" slot

of npdeObject

Value

an object of class NpdeObject; the ["results"] slot will contain pd and/or npde for a sample of the simulated datasets (depending on whether calc.npd/calc.npde are set), stored in pd.sim and/or npde.sim

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

See Also

npde, autonpde

```
data(theopp)
data(simtheopp)
x<-autonpde(theopp, simtheopp,1,3,4,boolsave=FALSE)
# Use random samples from N(0,1) to obtain a prediction interval on the empirical cdf of the npde
plot(x,plot.type="ecdf",bands=TRUE,approx.pi=TRUE)
# defaults to computing the pd and npde for 100 simulated datasets
# (in the theophylline example, this uses all the simulated datasets)
x<-dist.pred.sim(x)
# Use the npde from the simulated datasets to obtain a prediction interval on the empirical cdf
plot(x,plot.type="ecdf",bands=TRUE,approx.pi=FALSE)</pre>
```

8 gof.test

gof.test

Goodness-of-fit tests for npde

Description

Performs test on the selected variable (which=one of npde, pd or npd) or on a numeric vector

Usage

```
gof.test(object, parametric = TRUE, ...)
printgoftest(object, which = "npde", ...)
```

Arguments

object an object (currently has methods for types numeric, NpdeRes and NpdeObject)

parametric a boolean. If TRUE (default), parametric tests are performed

additional arguments passed on to the function; special arguments are na.action, which controls how to handle NAs in the results (na.action), verbose (if

which controls how to handle NAs in the results (na.action), verbose (if FALSE, suppresses printing of the results) and covsplit which requests the tests to be performed split by categories or quantiles of the data. If covsplit is TRUE, continuous covariates will be split in 3 categories (<Q1, Q1-Q3, >Q3) (see details in the PDF documentation), but this behaviour can be overriden by passing the argument ncat=XXX where XXX is the number of categories to di-

vide the continuous covariates in.

which character string giving (used by printgoftest)

Details

If object is an NpdeObject and an argument covsplit=TRUE is given in ..., in addition to the global descriptive statistics and tests, tests will be performed for each covariate in which.cov. This argument can be set in ...; barring an explicit specification, the component which.cov of the prefs slot for a NpdeObject object will be used. The default value is which.cov="all", which produces tests for each covariate in the dataset. Two additional dataframes will then be present:

cov.stat descriptive statistics and test p-values split by covariate and by categories

cov.p.value p-values split by covariate; for each covariate, two tests are performed: the first test is a correlation test for continuous covariates and a Chi-square test for categorical covariates; the second test is defined using the p-values of the global tests split by each category, and appling a Bonferroni correction to obtain an overall p-value (see PDF documentation for details)

The p.value elements is a named vector with four components:

p.mean p-value for the mean test (Wilcoxon test if parametric=FALSE, Student test if parametric=TRUE)

p.var p-value for the variance test (parametric=FALSE, Fisher test if parametric=TRUE)

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p.dist p-value for the distribution test (Shapiro-test for normality (npd, npde)/Kolmogorove-Smirnov test for uniformity)

p.global p-value for the global test (combination of the mean, variance and distribution tests with a Bonferroni correction)

The p-values are adjusted using a Bonferroni correction: the raw p-values of the 3 individual tests are multiplied by 3, and the p-value for the global test is equal to the minimum of the adjusted p-values.

Value

A list with the following elements:

```
mean mean
se.mean standard error of the mean
var variance
se.var standard error on variance
kurtosis kurtosis (see kurtosis)
skewness skewness (see skewness)
p.value p-values for several tests (see below)
```

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

K. Brendel, E. Comets, C. Laffont, and F. Mentre. Evaluation of different tests based on observations for external model evaluation of population analyses. *Journal of Pharmacokinetics and Pharmacodynamics*, 37:49–65, 2010.

See Also

```
kurtosis, skewness
```

```
data(theopp)
data(simtheopp)
#' # Calling autonpde with dataframes
x<-autonpde(theopp,simtheopp,1,3,4,boolsave=FALSE)
gof.test(x)</pre>
```

10 kurtosis

kurtosis

Kurtosis

Description

Computes the kurtosis.

Usage

kurtosis(x)

Arguments

Х

a numeric vector containing the values whose kurtosis is to be computed. NA values are removed in the computation.

Details

If N = length(x), then the kurtosis of x is defined as:

$$Nsum_i(x_i - mean(x))^4 (sum_i(x_i - mean(x))^2)^{(-2)}$$

3

Value

The kurtosis of x.

References

G. Snedecor, W. Cochran. Statistical Methods, Wiley-Blackwell, 1989

```
x <- rnorm(100)
kurtosis(x)</pre>
```

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npde.cens.method

Method used to handle censored data

Description

Specifies the method used to handle censored data (data below the limit of quantification LOQ

Details

Several methods are available to handle censored data.

omit pd and npde for censored data will be set to NA

cdf for an observation ycens_ij under the LOQ, a pd_ij will be imputed in the uniform distribution [0-pLOQ_ij] where pLOQ_ij is the probability that y_ij is below LOQ, according to the model; the predictive distribution will then be used to obtain a corresponding y*_ij. This is also performed for all simulated data, and the npde are then computed on the completed dataset containing the observed y_ij for the uncensored data and the y*_ij imputed for the censored data. This method is the default.

ipred an observation ycens_ij is replaced by the individual prediction according to the model (ipred, which must be present in the dataset). Simulated data are left untouched.

ppred an observation ycens_ij is replaced by the population prediction according to the model. Simulated data are left untouched.

loq an observation ycens_ij is replaced by the value of the LOQ. Simulated data are left untouched.

Value

This is not a function and does not have a return value, this is a statistical method.

More details can be found in the PDF documentation.

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

```
data(warfarin)
data(simwarfarinCov)
wcov<-autonpde(namobs=warfarin,namsim=simwarfarinCov, iid=1,ix=2,iy=4,icov=c(3,6:8),
namsav="warfCov", units=list(x="hr",y="mg/L", covariates=c("mg","kg","-","yr")))
# Diagnostic plots for warfarin with a covariate model
plot(wcov)
# Covariate plots</pre>
```

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```
xwt.scatt<-plot(wcov, plot.type="x.scatter", covsplit=TRUE, which.cov="wt")
xwt.qqplot<-plot(wcov, plot.type="qqplot", covsplit=TRUE, which.cov="wt")
xwt.box<-plot(wcov, plot.type="covariates", which.cov="wt")
xsex.scatt<-plot(wcov, plot.type="x.scatter", covsplit=TRUE, which.cov="sex")
xsex.qqplot<-plot(wcov, plot.type="qqplot", covsplit=TRUE, which.cov="sex")
xsex.box<-plot(wcov, plot.type="covariates", which.cov="sex")

# Transforming the reference profile for npd, compared to a VPC plot
plot.tnpd<-plot(wcov, plot.type="x.scatter", ref.prof=list(id=2),
main="tnpd with reference profile ID=2")
plot.vpc<-plot(wcov, plot.type="vpc", main="VPC")
grid.arrange(grobs=list(plot.tnpd, plot.vpc), nrow=1, ncol=2)</pre>
```

npde.decorr.method

Decorrelation methods in npde

Description

Specifies the method used to decorrelate observed and simulated data

Arguments

x a square matrix

cholesky decorrelation is performed through the Cholesky decomposition (default)

inverse decorrelation is performed by inverting Vi through the eigen function **polar** the singular-value decomposition (svd) is used

@return This is not a function and does not have a return value, this is a statistical method.

Details

More details can be found in the PDF documentation.

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

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npde.graphs

Save the graphs for a NpdeObject object to a file

Description

Save the graphs to a file on disk

Usage

```
npde.graphs(object, ...)
```

Arguments

```
object a NpdeObject object
... optional arguments to replace options in object
```

Details

The following options can be changed by passing the appropriate arguments: namsav (string giving the root name of the files, an extension depending on the type of graph will be added), namgr (string giving the full name of the file), type.graph (one of "eps", "pdf", "jpeg", "png")

Value

No return value, called for side effects

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F.Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

npde.plot.covariate

Covariate diagnostic plots

Description

Boxplot of the selected variable versus categories of covariates

Usage

```
npde.plot.covariate(npdeObject, which.y="npd", ...)
```

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Arguments

npdeObject an object returned by a call to npde or autonpde

which.y a string specifying the variable on the Y-axis (one of "yobs", "npde", "pd",

"npd")

... additional arguments to be passed on to the function, to control which metric

(npde, pd, npd) is used or to override graphical parameters (see the PDF docu-

ment for details, as well as set.plotoptions and npdeControl)

Details

For a categorical covariate, boxplots are produced for each category. Continuous covariates are split into quantile (by default, first quartile (<Q1), interquartile range (Q1-Q3) and upper quartile (>Q3), but the number of categories can be set by using the neat argument).

For each category, the median according to simulations under the model is shown (it can be suppressed by using the argument bands=FALSE)..

Value

a ggplot object or a list of ggplot objects (grobs)

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, F. Mentre F. Evaluation of different tests based on observations for external model evaluation of population analyses. *Journal of Pharmacokinetics and Pharmacodynamics*, 37:49-65, 2010.

See Also

npde, autonpde, set.plotoptions, npdeControl

npde.plot.data

Plot a NpdeData object

Description

Produces a spaghetti plot of the data

Usage

```
npde.plot.data(npdeObject, ...)
```

npde.plot.default

Arguments

npdeObject an object returned by a call to npde or autonpde

additional arguments to be passed on to the function, to control which metric

(npde, pd, npd) is used or to override graphical parameters (see the PDF docu-

ment for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

npde.plot.default

Diagnostic plots

Description

The default diagnostic plots produced after a call to npde or autonpde include a histogram of the distribution, a QQ-plot compared to the theoretical distribution, and scatterplots versus the independent variable and versus the population predictions from the model

Usage

```
npde.plot.default(npdeObject, ...)
```

Arguments

npdeObject an object returned by a call to npde or autonpde

... additional arguments to be passed on to the function, to control which metric

(npde, pd, npd) is used or to override graphical parameters (see the PDF docu-

ment for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

npde.plot.dist

npde.plot.dist Distribution plots of pd/n	ıpde
---	------

Description

Produces a plot of the cdistribution of a metric compared to their theoretical distribution. Three types of distribution plots are available: a histogram, a QQ-plot, or the empirical cdf.

Usage

```
npde.plot.dist(npdeObject, which="npd", dist.type="qqplot", ...)
```

Arguments

npdeObject	an object returned by a call to npde or autonpde
which	a string determining which metric to plot (one of "npde", "pd" or "npd"), defaults to "npd"
dist.type	string, one of "ecdf" (empirical cumulative density function), "hist" (histogram) or "qqplot" (QQ-plot of the empirical distribution versus the theoretical quantiles) to determine which type of plot (default is "qqplot")
	additional arguments to be passed on to the function, to control which metric (npde, pd, npd) is used or to override graphical parameters (see the PDF document for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

See Also

```
npde, autonpde, set.plotoptions
```

npde.plot.loq

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Plot of the probability that the observations are below the LOQ

Description

Plots the probability that the observations are below the LOQ along with the model predicted interval

Usage

```
npde.plot.loq(npdeObject,xaxis="x",nsim=200,...)
```

Arguments

npdeObject	an object returned by a call to npde or autonpde
xaxis	a string character, one of "x" (to plot $P(Y < LOQ)$) versus the value of the independent predictor) or "ypred" (versus the value of the population predictions). Defaults to "x"
nsim	number of simulations to be used for the computation of the prediction interval
	additional arguments to be passed on to the function, to control which metric (npde, pd, npd) is used or to override graphical parameters (see the PDF document for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

See Also

```
npde, autonpde, set.plotoptions
```

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npde.plot.npde

Plots for pd and npde

Description

Plots for pd and npde

Usage

```
npde.plot.pd(npdeObject, ...)
npde.plot.npde(npdeObject, ...)
npde.plot.npd(npdeObject, ...)
```

Arguments

npdeObject

an object returned by a call to npde or autonpde

additional arguments to be passed on to the function, to control which metric (npde, pd, npd) is used or to override graphical parameters (see the PDF document for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

npde.plot.scatterplot ########### Scatterplots - VPC, scatterplots versus indepen-plots and VPC

Description

Produces a scatterplot. Different types of scatterplots can be produced, with associated prediction bands (see details).

Usage

```
npde.plot.scatterplot(npdeObject, which.x="x", which.y="npd", ref.prof=NULL, ...)
```

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Arguments

npdeObject	an object returned by a call to npde or autonpde
which.x	a string specifying the variable on the X-axis (one of "x", "pred", "cov")
which.y	a string specifying the variable on the Y-axis (one of "yobs", "npde", "pd", "npd"), defaults to "npd"
ref.prof	either a character string (one of "covariate" or "all"), or a named list specifying the characteristics of the reference profile (see details)
	additional arguments to be passed on to the function, to control which metric (npde, pd, npd) is used or to override graphical parameters (see the PDF document for details, as well as set.plotoptions and npdeControl)

Details

VPC: obtained using which.x="x", which.y="yobs"

Scatterplots of npde/pd/npd can be obtained versus "x" (independent variable) or "pred" (population predictions from the model)

Scatterplots of npde/pd/npd/observations can be obtained versus covariates by setting the which.x argument to "cov" and selecting the appropriate which.y. The function will use the covariates in the which.cov element of the prefs slot. This can be overriden to cycle over all the covariates in the dataset by supplying the argument which.cov="all" in the call to the function.

Reference profile: a reference profile can be added to scatterplots of npd and npde versus the independent variable (see Comets et al. 2013)

If ref.prof="all" (and covsplit is FALSE), the reference plot will be computed over all subjects using the mean and SD of all simulated data in each bin (see documentation).

If ref.prof="covariate" and an additional argument covsplit is given (covsplit=TRUE), the reference plot will be adjusted for each covariate category over all the covariates in the which.cov element of the prefs slot (see npdeControl for details on the prefs slot of the npdeObject).

If ref.prof is given as a named list (eg list(ID=c(1,5)) or list(sex=0, dose=c(50,100)), where names should refer to columns in the data file (eg ID should be a column in the data)), the reference profile will be obtained by combining (in the first example above, the reference profile will be obtained using the simulated data for subjects 1 and 5, while in the second example it will be computed using the subjects with sex=0 given doses 50 or 100).

Value

a ggplot object or a list of ggplot objects (grobs)

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

20 npde.plot.select

E. Comets, T.H.T. Nguyen, and F. Mentré F. Additional features and graphs in the new npde library for R. 22th meeting of the Population Approach Group in Europe, Glasgow, United Kingdom, 2013.

See Also

npde, autonpde, set.plotoptions, npdeControl

 ${\tt npde.plot.select} \qquad \qquad \textit{Select plot for a NpdeObject object}$

Description

Select plot for a NpdeObject object

Usage

npde.plot.select(npdeObject,data=FALSE,ecdf=FALSE,qqplot=FALSE, histogram=FALSE, x.scatter=FALSE,pred.scatter=FALSE,x.box=FALSE,pred.box=FALSE, cov.scatter=FALSE, cov.x.scatter=FALSE, cov.pred.scatter=FALSE,cov.x.box=FALSE,cov.pred.box=FALSE, cov.ecdf=FALSE, cov.hist=FALSE, cov.qqplot=FALSE, vpc=FALSE,...)

Arguments

npdeObject	an object returned by a call to npde or autonpde				
data	boolean, whether to produce a plot of the data				
ecdf	boolean, whether to produce a distribution plot of the empirical distribution function				
qqplot	boolean, whether to produce a QQ-plot of the empirical distribution function				
histogram	boolean, whether to produce a histogram of the metric				
x.scatter	boolean, whether to produce a scatterplot of the metric as a function of X				
pred.scatter	boolean, whether to produce a scatterplot of the metric as a function of predictions				
x.box	boolean, whether to produce whisker plots of the metric as a function of X				
pred.box	boolean, whether to produce whisker plots of the metric as a function of predictions				
cov.scatter	boolean, whether to produce a scatterplot of the metric as a function of covariate(s) $ \\$				
cov.x.scatter	boolean, whether to produce a scatterplot of the metric as a function of X, split by covariate(s)				
cov.pred.scatter					
	boolean, whether to produce a scatterplot of the metric as a function of predictions, split by covariate(s)				
cov.x.box	boolean, whether to produce whisker plots of the metric as a function of X, split by covariate(s)				

npde.plot.splitcov 21

cov.pred.box	boolean, whether to produce whisker plots of the metric as a function of predictions, split by covariate(s)
cov.ecdf	boolean, whether to produce a distribution plot of the empirical distribution function, split by covariate(s)
cov.hist	boolean, whether to produce a distribution plot of the empirical distribution function, split by covariate(s)
cov.qqplot	boolean, whether to produce a distribution plot of the empirical distribution function, split by covariate(s)
vpc	boolean, whether to produce a VPC
	additional arguments to be passed on to the function, to control which metric (npde, pd, npd) is used or to override graphical parameters (see the PDF document for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

See Also

npde, autonpde, set.plotoptions

npde.plot.splitcov	Plots split by covariate for a NpdeObject object	

Description

Plots split by covariate for a NpdeObject object (equivalent to using covsplit=TRUE with the appropriate plot.type)

Usage

```
npde.plot.splitcov(npdeObject, which.plot="x", ...)
```

22 npde.save

Arguments

npdeObject an object returned by a call to npde or autonpde

which.plot one of "x" (scatterplots of the metric versus X), "pred" (scatterplots of the metric

versus predictions), "ecdf" (empirical distribution function), "hist" (histogram),

"qqplot"

... additional arguments to be passed on to the function, to control which metric

(npde, pd, npd) is used or to override graphical parameters (see the PDF docu-

ment for details, as well as set.plotoptions)

Value

a ggplot object or a list of ggplot objects (grobs)

npde.save

Save the results contained in a NpdeObject object to a file

Description

Save the results to a table on disk

Usage

```
npde.save(object, ...)
```

Arguments

object a NpdeObject object

... optional arguments to replace options in object

Details

The following options can be changed by passing the appropriate arguments: namsav (string giving the root name of the files, an extension .npde will be added), nameres (string giving the full name of the file)

Value

No return value, called for side effects

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F.Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

npdeControl 23

npdeControl	Set options for an NpdeObject

Description

Set, replace and check options for an NpdeObject

match.

Usage

```
npdeControl(boolsave = TRUE, namsav = "output", type.graph = "eps",
verbose = FALSE, calc.npde = TRUE, calc.npd = TRUE, decorr.method = "cholesky",
cens.method = "omit", ties = TRUE, sample = FALSE)

check.control.options(opt)

replace.control.options(opt,...)

check.control.options(opt, ...)
```

Arguments

boolsave	whether to save the results (a file containing the numerical results and a file with the graphs)
namsav	the root name of the files to save to (the file with the results will be named ROOTNAME.npde and the graphs will be saved to ROOTNAME.format where format is given by the type.graph argument)
type.graph	type of graph to save to (one of "eps", "pdf", "jpeg", "png")
verbose	a boolean; if TRUE, a message is printed as the computation of the npde begins for each new subject
calc.npde	a boolean; TRUE to compute npde
calc.npd	a boolean; TRUE to compute npd
decorr.method	the method used to decorrelate simulated and observed data (see npde.decorr.method)
cens.method	the method used to handle censored data (see npde.cens.method)
ties	if FALSE, a smoothing will be applied to prediction discrepancies to avoid ties
sample	if TRUE, the test on the pd will be performed after randomly sampling only pd per subject
opt	a list of control options to be checked
• • •	named parameters to be changed. The names will be compared to the names of the control variables and changed, with warnings issued for names that do not

24 npdeData

Value

A list of settings for the computation of pd/npde

npdeData	Creates a NpdeData object	

Description

This function is used to create a NpdeData object, representing a longitudinal data structure, and fill it with data from a dataframe or a file on disk

Usage

```
\label{lem:npdeData} $$ name.data,header=TRUE,sep="",na.strings=c(".","NA"),name.group, \\ name.predictor, name.response, name.covariates,name.cens,name.miss,name.ipred, \\ units=list(x="",y="",covariates=c()), detect=TRUE,verbose=FALSE) \\
```

Arguments

name of the file containing the observed data, or a dataframe containing the observed data
boolean indicating whether the file has a header (mandatory if detect is TRUE)
field separator (for files on disk)
strings to be considered as indicating NA
name/number of the column in the observed data containing the patient ID (if missing and detect is TRUE, columns named id, subject or sujet (regardless of case) will be assumed to contain this information)
name/number of the column in the observed data containing the independent variable X (if missing and detect is TRUE, columns named xobs, time, dose, x, temps, tim (regardless of case) will be assumed to contain this information)
name/number of the column in the observed data containing the dependent variable Y (if missing and detect is TRUE, columns named yobs, response, resp, conc, concentration (regardless of case) will be assumed to contain this information)
S
name/number of the column(s) containing covariate information (optional)
name/number of the column containing information about censored data (cens) (if missing and detect is TRUE, column with a name containing cens (regardless of case) will be assumed to contain this information)
name/number of the column containing information about missing data (MDV) (if missing and detect is TRUE, column called mdv or miss (regardless of case) will be assumed to contain this information)

NpdeData-class 25

name.ipred	name/number of the column(s) with individual predictions (ipred) (if missing and detect is TRUE, column with a name containing ipred (regardless of case) will be assumed to contain this information)
units	a list with components x, y and cov (optional), specifying the units respectively for the predictor (x), the response (y), and the covariates (a vector of length equal to the number of covariates). Units will default to (-) if not given.
detect	a boolean controlling whether automatic recognition of columns in the dataset is on, defaults to TRUE
verbose	whether to print warning messages, defaults to FALSE (set to TRUE to check how data is being handled)

Value

an object of class NpdeData

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F. Mentr\'e. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

See Also

```
npde, autonpde
```

Examples

```
data(theopp)
x<-npdeData(theopp) # Automatic detection
print(x)
x<-npdeData(theopp,name.group="ID",name.predictor="Time",name.response="Conc",
name.covariates=c("Wt"),units=list(x="hr",y="mg/L",covariates="kg")) # Explicit
print(x)</pre>
```

NpdeData-class

Class "NpdeData" representing the structure of the longitudinal data

Description

A longitudinal data structure

26 NpdeObject-class

Objects from the Class

NpdeData objects are typically created by a call to npdeData contain the following slots:

name.group character string giving the name of the grouping term (ID)

name.predictor character string giving the name of the predictor (X)

name.response character string giving the name of the response (Y)

name.cens character string giving the name of the censoring indicator

name.mdv character string giving the name of the missing data indicator

name.covariates vector of character string giving the name(s) of the covariates

name.ipred character string giving the name of the individual predictions

units (optional) a list with the units for X, Y, and covariates

data a dataframe containing the data

N number of subjects

ntot.obs total number of non-missing observations

nind.obs vector of size N giving the number of non-missing observations for each subject

ind index of non-missing observations

icens index of censored observations (non-missing)

not.miss a vector of boolean indicating for each observation whether it is missing (FALSE) or available (TRUE)

loq the censoring value

Methods

show(npde.data): Prints a short summary of object npde.data

qqplot.npde(npde.data): QQ-plot for NpdeData object (TODO: change for NpdeObject in final package)

Examples

```
methods(class="NpdeData")
showClass("NpdeData")
```

NpdeObject-class

Class "NpdeObject"

Description

An object of class NpdeObject

npdeSimData 27

Objects from the Class

NpdeObject objects are typically created by calls to npde or autonpde. They contain the following slots:

data an object of class NpdeData, containing the observed data sim.data an object of class NpdeSimData, containing the simulated data results an object of class NpdeRes, containing the results options a list of options

prefs a list of graphical preferences for the plots

Methods

```
print(x): Prints a summary of object
show(x): Prints a short summary of object
showall(x): Prints a detailed summary of object
plot(x): Diagnostic and other plots. More details can be found in plot.NpdeObject
summary(x): Returns a summary of object x in list format
gof.test(x, parametric=TRUE, ...): Returns goodness-of-fit tests
set.plotoptions(x): Sets options for graphs (internal method used in plots)
```

See Also

```
npde, autonpde, NpdeData, NpdeSimData, NpdeRes, gof.test
```

Examples

```
methods(class="NpdeObject")
showClass("NpdeObject")
```

npdeSimData

Creates a NpdeSimData object

Description

This function is used to create a NpdeSimData object containing the simulated data corresponding to an NpdeData object

Usage

```
npdeSimData(npde.data, name.simdata, header=TRUE, verbose=FALSE)
```

28 NpdeSimData-class

Arguments

npde.data a NpdeData object

name simdata name of the file containing the simulated data, or a dataframe containing it boolean indicating whether the file has a header (mandatory if detect is TRUE) werbose whether to print warning messages, defaults to FALSE (set to TRUE to check

how data is being handled)

Value

an object of class NpdeSimData

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

See Also

NpdeData, npde, autonpde

NpdeSimData-class Class "NpdeSimData" representing the structure of the longitudinal

data

Description

A longitudinal data structure, with simulated data

Objects from the Class

NpdeSimData objects are created by associating an NpdeData object with matching simulated data, and they contain the following slots.

nrep number of replications)

datsim a dataframe containing the simulated data, with columns: idsim (subject id), irsim (replication index), xsim (simulated x), ysim (simulated response). After a call to npde or autonpde, an additional column ydsim (decorrelated replicated data) will be added.

Methods

print(npde.simdata): Prints a summary of object npde.simdatashow(npde.simdata): Prints a short summary of object npde.simdatashowall(npde.simdata): Prints a detailed summary of object npde.simdata

See Also

npde, autonpde

plot.NpdeData 29

Examples

```
showClass("NpdeSimData")
```

plot.NpdeData

Plots a NpdeData object

Description

Plots the data in a NpdeData object

Usage

```
## S3 method for class 'NpdeData' plot(x, y, ...)
```

Arguments

x a NpdeData object

y unused, here for compatibility with the base plot function

... additional graphical parameters to be passed on to the plot

Details

The default plot is a spaghetti plot of all the data, with a line joining the observations for each subject. If censored data is present, it is shown with a different symbol and colour.

Value

currently does not return anything, use plot(x, plot.type="data") on the npdeObject x (TODO; a ggplot object)

References

K. Brendel, E. Comets, C. Laffont, C. Laveille, and F.Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. *Pharmaceutical Research*, 23:2036–49, 2006.

See Also

```
set.plotoptions
```

30 plot.NpdeObject

Examples

```
data(theopp)
x<-npdeData(theopp,name.group="ID",name.predictor="Time",name.response="Conc",
name.covariates=c("Wt"),units=list(x="hr",y="mg/L",covariates="kg"))
plot(x)
```

plot.NpdeObject

Plots a NpdeObject object

Description

Plots the data and diagnostic plots in a NpdeObject object

Usage

```
## S3 method for class 'NpdeObject'
plot(x, y, ...)
```

Arguments

a NpdeObject object Χ

unused, here for compatibility with the base plot function У

additional graphical parameters, which when given will supersede graphical

preferences stored in the object

Details

The default plots are represented as a 2x2 array with distribution plots on the top row (histogram and QQ-plot), and scatterplots of npde versus independent variable and population predictions on the bottom row. The graph is plotted in a graphic device window, unless the result is stored in an object (eg myplot<-plot(x)) which can then be printed (eg using print(myplot)).

@references K. Brendel, E. Comets, C. Laffont, C. Laveille, and F.Mentre. Metrics for external model evaluation with an application to the population pharmacokinetics of gliclazide. Pharmaceutical Research, 23:2036-49, 2006.

Value

```
a ggplot object or a list of ggplot objects (grobs)
```

See Also

```
set.plotoptions
```

print.NpdeData 31

Examples

```
data(theopp)
data(simtheopp)

x<-autonpde(theopp,simtheopp,iid="ID",ix="Time", iy="Conc", boolsave=FALSE)
plot(x)</pre>
```

print.NpdeData

Prints objects from the npde package

Description

prints objects of classes NpdeData, NpdeSimData, NpdeRes and NpdeObject

Usage

```
## S3 method for class 'NpdeData'
print(x, nlines = 10, ...)

## S3 method for class 'NpdeRes'
print(x, nlines = 10, ...)

## S3 method for class 'NpdeObject'
print(x, nlines = 10, ...)
```

Arguments

```
    x an object of class NpdeData, NpdeSimData, NpdeRes or NpdeObject
    nlines number of lines from the dataset to print
    ... Additional arguments (ignored)
```

Value

None

32 set.plotoptions

set.plotoptions

Set graphical preferences

Description

This function is used to set options for graphs

Usage

```
set.plotoptions(object)
## Default S3 method:
set.plotoptions(object)
## S3 method for class 'NpdeData'
set.plotoptions(object)
## S3 method for class 'NpdeObject'
set.plotoptions(object)
```

Arguments

object

an object of class NpdeData or NpdeObject

Details

See documentation for a list of available options.

Value

a list of options for graphs

Author(s)

Emmanuelle Comets <emmanuelle.comets@bichat.inserm.fr>

See Also

```
npde, autonpde
```

show 33

show

Displays npde objects

Description

Prints the structure of objects from the package

Usage

```
## S4 method for signature 'NpdeData'
show(object)

## S4 method for signature 'NpdeSimData'
show(object)

## S4 method for signature 'NpdeRes'
show(object)

## S4 method for signature 'NpdeObject'
show(object)
```

Arguments

object

an object from the npde package (NpdeData, NpdeRes, NpdeObject)

showall

Contents of an object

Description

Prints the contents of an object

Usage

```
showall(object)
## S3 method for class 'NpdeRes'
showall(object)
## S3 method for class 'NpdeObject'
showall(object)
```

Arguments

object

a NpdeData object

34 simtheopp

Value

No return value, shows the object

simtheopp	Simulated data for the computation of normalised prediction distribution errors in the theophylline dataset
	······································

Description

The simtheopp dataset contains 100 simulations using the design of dataset theopp. These simulations are used to compute npde. The control file used to perform the simulations can be found in the subdirectory 'doc' within the library npde.

Usage

simtheopp

Format

A data frame with 132000 rows and 3 variables This data frame contains the following columns:

ID an ordered factor with levels 1, ..., 12 identifying the subject on whom the observation was made. The ordering is first by simulation then by increasing time.

xsim time since drug administration when the sample was drawn (hr).

ysim simulated theophylline concentration (mg/L).

Details

See the opp for a description of the original dataset.

The simulated data was obtained using the software *NONMEM*. A one-compartment model was fit to the data. An exponential interindividual variability was assumed for the three parameters (absorption rate constant ka, volume of distribution V and clearance CL) and a combined additive and proportional residual error model was usd. The estimated parameters were then used to simulate 100 datasets with the same structure as the original dataset. Thus, for each observation in the original dataset, the simulated dataset contains 100 simulations under the model used for the estimation.

This dataset is provided so that users can figure out what type of data is needed for the computation of prediction distribution errors. More information can be found in the User Guide distributed along with this package, which contains a run-through of the theophylline example.

Source

Boeckmann, A. J., Sheiner, L. B. and Beal, S. L. (1994), *NONMEM Users Guide: Part V*, NON-MEM Project Group, University of California, San Francisco.

skewness 35

References

PDF documentation for npde 3.0: https://github.com/ecomets/npde30/blob/main/userguide_npde_3.0.pdf

See Also

theopp

Examples

```
data(simtheopp)
# Plotting the simulated data for subject 1 in the first simulation
plot(ysim[2:12]~xsim[2:12],data=simtheopp,xlab="Time after dose (hr)",
ylab="Theophylline concentration (mg/L)",type="l",
main="Example of simulated data for subject 1")
# Plotting a 90% prediction interval for the observations in theopp
# using the simulated data in simtheopp
# note : differences in doses between subjects are not taken into account
data(theopp)
xp1<-c(0,0.25,0.5,1,2,3.5,5,7,9,12,24)
xpl1 < -list(c(0,0.1),c(0.2,0.4),c(0.5,0.65),c(0.9,1.2),c(1.9,2.2),c(3.4,4),
c(4.9,5.2), c(6.9,7.2), c(8.8,9.4), c(11.5,12.2), c(23.7,24.7))
ypl<-cbind(xpl=xpl,binf=xpl,median=xpl,bsup=xpl)</pre>
for(i in 1:(length(xpl))) {
  vec<-simtheopp$ysim[simtheopp$xsim>=xpl1[[i]][1] &simtheopp$xsim<=xpl1[[i]][2]]</pre>
  ypl[i,2:4] < -quantile(vec,c(0.05,0.5,0.95))
}
plot(Conc~Time, data=theopp, xlab="Time after dose (hr)",
ylab="Theophylline concentration (mg/L)")
lines(ypl[,1],ypl[,3],lwd=2)
lines(ypl[,1],ypl[,2],lty=2)
lines(ypl[,1],ypl[,4],lty=2)
```

skewness

Skewness

Description

Computes the skewness.

Usage

```
skewness(x)
```

36 subset.NpdeData

Arguments

Х

a numeric vector containing the values whose skewness is to be computed. NA values are removed in the computation.

Details

If N = length(x), then the skewness of x is defined as

$$N^{-1} \operatorname{sd}(x)^{-3} \sum_{i} (x_i - \operatorname{mean}(x))^3.$$

Value

The skewness of x.

References

G. Snedecor, W. Cochran. Statistical Methods, Wiley-Blackwell, 1989

Examples

```
x <- rnorm(100)
skewness(x)</pre>
```

subset.NpdeData

Subsetting a NpdeData object

Description

Return subset of data from a NpdeData object

Usage

```
## S3 method for class 'NpdeData'
subset(x, subset, ...)
## S3 method for class 'NpdeObject'
subset(x, subset, ...)
```

Arguments

x A NpdeData object

subset logical expression indicating elements or rows to keep: missing values are taken

as false.

... Additional arguments (ignored)

Value

a NpdeData object with a subset of the original data

summary.NpdeData 37

summary.NpdeData

Summary of a NpdeData object

Description

Extracts elements from a NpdeData object

Usage

```
## S3 method for class 'NpdeData'
summary(object, print = TRUE, ...)
## S3 method for class 'NpdeRes'
summary(object, print = TRUE, ...)
## S3 method for class 'NpdeObject'
summary(object, ...)
```

Arguments

object A NpdeData object

print whether to print to data to stdev

... Additional arguments (ignored)

Value

A list with elements N (nb of subjects), data (dataframe containing the data), ntot.obs (total nb of observations), nind.obs (nb of observations per subject)

theopp

Pharmacokinetics of theophylline

Description

The theopp data frame has 132 rows and 5 columns of data from an experiment on the pharmacokinetics of theophylline.

Usage

theopp

38 theopp

Format

This data frame contains the following columns:

ID an ordered factor with levels 1, ..., 12 identifying the subject on whom the observation was made. The ordering is by Time at which the observation was made.

Dose dose of the ophylline administered or ally to the subject (mg/kg).

Time time since drug administration when the sample was drawn (hr).

Conc theophylline concentration in the sample (mg/L).

Wt weight of the subject (kg).

Details

Boeckmann, Sheiner and Beal (1994) report data from a study by Dr. Robert Upton of the kinetics of the anti-asthmatic drug theophylline. Twelve subjects were given oral doses of theophylline then serum concentrations were measured at 11 time points over the next 25 hours.

These data are analyzed in Davidian and Giltinan (1995) and Pinheiro and Bates (2000) using a two-compartment open pharmacokinetic model.

These data are also available in the library datasets under the name Theoph in a slightly modified format and including the data at time 0. Here, we use the file in the format provided in the *NONMEM* installation path (see the User Guide for that software for details).

Source

Boeckmann, A. J., Sheiner, L. B. and Beal, S. L. (1994), *NONMEM Users Guide: Part V*, NON-MEM Project Group, University of California, San Francisco.

Davidian, M. and Giltinan, D. M. (1995) *Nonlinear Models for Repeated Measurement Data*, Chapman & Hall (section 5.5, p. 145 and section 6.6, p. 176)

Pinheiro, J. C. and Bates, D. M. (2000) *Mixed-effects Models in S and S-PLUS*, Springer (Appendix A.29)

References

```
PDF documentation for npde 3.0: https://github.com/ecomets/npde30/blob/main/userguide_npde_3.0.pdf
```

```
data(theopp)
#Plotting the theophylline data
plot(Conc~Time,data=theopp,xlab="Time after dose (hr)",
ylab="Theophylline concentration (mg/L)")
```

virload 39

virload

Simulated HIV viral loads in HIV patients

Description

This is simulated data, based on real data obtained in a phase II clinical trial supported by the French Agency for AIDS Research, the COPHAR 3-ANRS 134 trial (Goujard et al., 2010). The original study included 35 patients, who received a once daily dose containing atazanavir (300 mg), ritonavir (100 mg), tenofovir disoproxil (245 mg) and emtricitabine (200 mg) during 24 weeks. Viral loads were measured 6 times over a treatment period of 24 weeks (day 0, 28, 56, 84, 112, 168).

Usage

virload

Format

This data frame contains the following columns:

ID an ordered factor with levels 1, ..., 50 identifying the subject on whom the observation was made. The ordering is by Time at which the observation was made.

Time time since the beginning of the study (days)

Log_VL logarithm (base 10) of the viral load (copies/L)

cens indicator variable (cens=1 for censored data, cens=0 for observed data)

ipred individual predictions

Details

The datasets were generated in a simulation study designed to evaluate the new method proposed to handle BQL data (Nguyen et al., 2011). Data was simulated using a simple bi-exponential HIV dynamic model describing the two-phase decline of viral load during anti-retroviral treatment.

The virload data frame has 300 rows and 4 columns of data. The dataset was then censored at two different LOQ levels (LOQ=20 or 50~copies/mL) to generate two datasets containing different proportions of BQL data, creating the data frames virload20 andvirload50 respectively

The file simvirload contains 500 simulations under the same model. A full version of the simulated data with 1000 simulations can be downloaded from the github for npde3.0: https://github.com/ecomets/npde30/blob/main/keep/data/simvirload.tab

Source

Goujard, C., Barrail-Train, A., Duval, X., Nembot, G., Panhard, X., Savic, R., Descamps, D., Vrijens, B., Taburet, A., Mentre, F., and the ANRS 134 study group (2010). Virological response to atazanavir, ritonavir and tenofovir/emtricitabine: relation to individual pharmacokinetic parameters and adherence measured by medication events monitoring system (MEMS) in naive HIV-infected patients (ANRS134 trial). *International AIDS Society 2010*, Abstr WEPE0094.

40 warfarin

References

Examples

```
data(virload)
str(virload)
data(virload50)
# Plotting the data
plot(Log_VL~Time,data=virload,xlab="Time (d)",ylab="Viral loads, base 10 log-scale (cp/mL)")
plot(Log_VL~Time,data=virload50,xlab="Time (d)",ylab="Viral loads, base 10 log-scale (cp/mL)")
```

warfarin

Pharmacokinetics of warfarin

Description

The warfarin data frame has 251 rows and 8 columns of data containing data on the pharmacokinetics of warfarin, an anticoagulant drug used in the prevention of thrombosis and thromboembolism.

Usage

warfarin

Format

This data frame contains the following columns:

age age of the subject (yr)

```
id an integer identifying the subject on whom the observation was made
time time since drug administration when the sample was drawn (hr)
amt total dose received by the subject (mg)
dv warfarin concentration in the sample (mg/L)
dvid observation type (1 for all observations)
wt weight of the subject (kg)
sex subject gender (0=female, 1=male)
```

warfarin 41

Details

The dataset is the PK part of a larger dataset including both warfarin concentrations and prothrombin complex activity (PCA), which measures the decreased coagulation activity resulting from the inhibition of vitamin K recycling, the mechanism of action of warfarin. It contains the concentrations measured in 32 healthy subjects after a single oral dose of warfarin sodium (1.5 mg/kg of body weight). The subjects in the study were sampled at different times over a period of up to 120 hours.

The data is distributed with the Monolix software as a demo for PK/PD modelling. The data has been slightly reformated for R, removing the line at time=0 and filling the amt column with the dose for each subject, following the output of simulx which was used to simulate data from two alternative models to fit this dataset.

Two datasets containing simulated data are associated with the warfarin data. For each dataset, 1000 simulations of the original data were performed for the computation of npde. The package contains only the simulated data simwarfarinCov because of size constraints. simwarfarinBase can be downloaded from the github for npde3.0: https://github.com/ecomets/npde30/blob/main/keep/data/simwarfarinBase.tab

simwarfarinBase the data in this dataset was simulated according to a base model without covariates: the PK model was a two-compartment model, with first-order absorption and a time-delay. Interindividual variability was modelled as log-normal distributions for parameters Tlag, ka, Cl and V1, and the error model was a combined error model. The parameters were estimated by Monolix.

simwarfarinCov the data in this dataset was simulated according to a model including several covariates: an age (centered on 30 yr) effect on Cl, a weight (centered on 70 kg) effect on Cl and V1, and a gender effect on V1. The covariate model was built in Monolix.

Source

O'Reilly (1968). Studies on coumarin anticoagulant drugs. Initiation of warfarin therapy without a loading dose. Circulation 1968, 38:169-177.

References

```
PDF documentation for npde 3.0: https://github.com/ecomets/npde 30/blob/main/userguide\_npde_3.0.pdf
```

```
data(warfarin)
#Plotting the warfarin PK data
plot(dv~time,data=warfarin,xlab="Time after dose (hr)",
ylab="Warfarin concentration (mg/L)")
```

 $\verb§[,NpdeSimData-method]{$ \textit{Get/set methods for NpdeData object}$}$

Description

Access slots of a NpdeData using the object["slot"] format

Usage

```
## S4 method for signature 'NpdeSimData'
x[i, j, drop]

## S4 method for signature 'NpdeData'
x[i, j, drop]

## S4 replacement method for signature 'NpdeData'
x[i, j] <- value

## S4 method for signature 'NpdeRes'
x[i, j, drop]

## S4 method for signature 'NpdeObject'
x[i, j, drop]</pre>
```

Arguments

X	object from which to extract element(s) or in which to replace element(s)
i, j	indices specifying elements to extract or replace. Indices are numeric or character vectors or empty (missing) or NULL
drop	For matrices and arrays. If TRUE the result is coerced to the lowest possible dimension (see the examples). This only works for extracting elements, not for the replacement. See drop for further details
value	typically an array-like R object of a similar class as x

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