Package 'qpNCA'

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Type Package

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calc.ctmax

Calculate Cmax and Tmax

Description

Calculates Cmax and Tmax from raw data for each PK curve defined using by.

Usage

```
calc.ctmax(x, by = character(0), timevar = "time", depvar = "dv")
```

Arguments

x data.frame

by column names in x indicating grouping variables

timevar variable name containing the actual sampling time after dose

depvar variable name containing the dependent variable (e.g., concentration)

Details

Input dataset can contain all uncorrected data, including LOQ; estimate first occurence of maximum concentration for each PK curve; if all concentrations are NA, sets Cmax and Tmax also to NA.

Value

A dataset with estimates for the Cmax (maximum concentration) and Tmax (time of first occurence of cmax) parameters: one observation per subject

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Examples

```
example(est.thalf)
ctmax <- x %>% calc.ctmax(by = 'subject')
ctmax %>% head
```

calc.par

Calculate NCA Parameters

Description

Calculates PK parameters for which half-life is not needed in the calculation for each PK curve defined using by.

Usage

```
calc.par(
    X,
    by = character(0),
    tau = NA,
    tstart = NA,
    tend = NA,
    teval = NA,
    route = "EV",
    method = 1
)
```

Arguments

X	contains all data after time/concentration deviation corrections obtained from correct.time and correct.conc
by	column names in x indicating grouping variables
tau	dosing interval (for multiple dosing); NA (default) for if single dose; x\$tau overrides
tstart	start time of partial AUC (start>0); NA (default) if not requested; x\$tstart overrides
tend	end time of partial AUC; NA (default) if not requested; x\$tend overrides
teval	user selected AUC interval; NA (default) if not requested; x\$teval overrides
route	route of drug administration ("EV", "IVB", "IVI"); x\$route overrides
method	method for trapezoidal rule

- 1: linear up linear down
- 2: linear up logarithmic down
- 3: linear before first Tmax, logarithmic after first Tmax

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Value

A dataset with estimates for the following parameters, one observation per subject:

Parameter	Description
t0.ok	flags if t=0 concentration could be corrected/imputes. If not, no AUCs starting at t=0 are calculated
tlast.ok	flags if there is at least one measurable concentration. If not, no AUClast can be calculated
tlast	time of last sample with measurable concentration
clast.obs	observed concentration at tlast
aucall	auc calculated over all observations, including values below LOQ (which are set to 0)
auclast	auc calculated using all observations up to and including the last measurable concentration (clast.obs at tlast
aumcall	aumc calculated over all observations, including values below LOQ (which are set to 0)
aumclast	aumc calculated using all observations up to and including the last measurable concentration (clast.obs at tla
tau	the dosing interval (if specified)
calc.tau	flags if AUCtau could be calculated
auctau	auc calculated over the dosing interval, only calculated if tau is specified
aumctau	aumc calculated over the dosing interval, only calculated if tau is specified
teval	user selected AUC interval starting at t=0 (if specified)
calc.teval	flags if AUCteval could be calculated
aucxx	auc calculated from t=0 up to/including teval, only calculated if teval is specified (xx is substituted by teval)
calc.part	flags if AUCpart could be calculated
tstart	start time of partial AUC (if specified)
tend	end time of partial AUC (if specified)
aucx_y	partial auc from time=x up to/including time=y, where x>0, only calculated if tstart and tend are specified
c0	back-extrapolated concentration at t=0 for IV bolus administration
area.back.extr	area back-extrapolated to 0

Examples

```
example(correct.conc)
par <- x %>% calc.par(by = 'subject')
par %>% head
```

calc.par.th Calculate Lambda_z Parameters

Description

Calculates PK parameters that need lambda_z.

Usage

```
calc.par.th(
  x,
  by = character(0),
```

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```
th = th,
covariates = NA,
dose = "dose",
factor = 1,
reg = "SD",
ss = "N",
route = "EV"
)
```

Arguments

X	result parameter dataset from calc.par
by	column names in x indicating grouping variables
th	result dataset from est.thalf
covariates	covariates dataset (containing at least dose for CL calculation); defaults to unique combinations of by and dose evaluated on x; can be character name of csv file or local object
dose	variable containing the dose amount; default 'dose' set to 1 if not in names(x)
factor	conversion factor for CL and V calculation (e.g. dose in mg, conc in ng/mL, factor=1000); x $\$$ factor overrides
reg	regimen, "sd" or "md"; x\$reg overrides
SS	is steady state reached (y/n); x\$ss overrides
route	of drug administration ("EV", "IVB", "IVI"); x\$route overrides

Value

A dataset containing all parameters calculated in est.thalf and calc.par with estimates for the following parameters added, one observation per subject:

Parameter	Description
clast.pred	predicted concentration at tlast
aucinf.obs	aucinf based on observed concentration at tlast
aucinf.pred	aucinf based on predicted concentration at tlast
aumcinf.obs	area under the first moment curve extrapolated to infinity, based on observed concentration at tlast
aumcinf.pred	area under the first moment curve extrapolated to infinity, based on predicted concentration at tlast
cl.obs, cl.f.obs	clearance based on aucinf.obs, at steady state based on auctau
cl.pred, cl.f.pred	clearance based on aucinf.pred
cl.ss, cl.f.ss	clearance at steady state, based on auctau
mrt.obs	mean residence time based on aumcinf.obs and aucinf.obs
mrt.pred	mean residence time based on aumcinf.pred and aucinf.pred
vz.obs, vz.f.obs	distribution volume based on cl.f.obs, at steady state based on auctau
vz.pred, vz.f.pred	distribution based on cl.pred/cl.f.pred
vss.obs	steady-state volume based on cl.obs and mrt.obs
vss.pred	steady-state volume based on cl.pred and mrt.pred
pctextr.pred	percentage of AUC extrapolated to infinity, based on aucinf.pred
pctextr.obs	percentage of AUC extrapolated to infinity, based on aucinf.obs
pctback.pred	percentage of AUC extrapolated back to 0, based on aucinf.pred

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pctback.obs

percentage of AUC extrapolated back to 0, based on aucinf.obs

Note: ctmax must be merged separately as those were calculated from uncorrected data.

Examples

```
example(calc.par) # creates par
# notice x includes (optional) loqrule, includeCmax, reg, method, route, ss
covs <- Theoph %>%
    select(subject = Subject, Wt, dose = Dose) %>%
    unique %>%
    mutate(dose = dose * Wt, subject=as.numeric(as.character(subject))) # see ?Theoph
y <- x %>% select(subject, reg, ss, loqrule) %>% unique
y %<>% mutate(factor = 1)
par %<>% left_join(y, by = 'subject')
par %<% calc.par.th(by = 'subject', th = th, covariates = covs)
par %<% left_join(ctmax, ., by = 'subject')
par %>% head
par %>% data.frame %>% head(2)
```

check.input

Check qpNCA function arguments for validity

Description

Checks whether all function arguments are valid and entered column names are present in input data

See qpNCA for description of the arguments

Usage

```
check.input(
    x,
    by = NA,
    nomtimevar = NA,
    timevar = NA,
    depvar = NA,
    loqvar = NA,
    loqvar = NA,
    loqvar = NA,
    loqvar = NA,
    includeCmax = NA,
    exclvar = NA,
    plotdir = NA,
    timelab = NA,
    deplab = NA,
```

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```
tau = NA,
tstart = NA,
tend = NA,
teval = NA,
covariates = NA,
dose = NA,
factor = NA,
reg = NA,
rs = NA,
route = NA,
method = NA
```

Arguments

x data.frame

by column names in x indicating grouping variables

nomtimevar variable name containing the nominal sampling time after dose timevar variable name containing the actual sampling time after dose

depvar variable name containing the dependent variable (e.g., concentration)

bloqvar variable name containing the BLOQ flag (0: no, 1: yes)

loqvar variable name containing the LOQ value

logrule rule number to be applied to the LOQ values in the curve; x\$logrule overrides if

provided

includeCmax in half-life estimation? (y/n); x\$includeCmax overrides if pro-

vided

exclvar variable name indicating points to be excluded in half-life estimation (these

should have exclvar = 1)

plotdir directory where regression plots (.PNG) will be saved; NA gives default loca-

tion, NULL skips regression plots

timelab label for time axis in regression plots

deplab label for dependent variable axis in regression plots

tau dosing interval (for multiple dosing); NA (default) if single dose; x\$tau over-

rides

tstart start time of partial AUC (start>0); NA (default) if not requested; x\$tstart over-

rides

tend end time of partial AUC; NA (default) if not requested; x\$tend overrides

teval user selected AUC interval (starting at t=0); NA (default) if not requested; x\$teval

overrides

covariates covariates dataset; Must contain the dose variable

dose variable containing the dose amount

factor conversion factor for CL and V calculation (e.g. dose in mg, conc in ng/mL,

factor=1000); x\$factor overrides if provided

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reg	regimen, "SD" or "MD"; x\$reg overrides if provided
SS	is steady state reached (y/n); x\$ss overrides if provided
route	route of drug administration ("EV", "IVB", "IVI"); x\$route overrides if provided

method for trapezoidal rule; x\$method overrides if provided

Value

Check results

method

correct.conc

Correct Missing Concentration

Description

Corrects missing concentration at critical time points (e.g, predose, TAU, start and end of user selected AUC interval).

Usage

```
correct.conc(
    x,
    by = character(0),
    nomtimevar = "ntad",
    tau = NA,
    tstart = NA,
    tend = NA,
    teval = NA,
    th = NA,
    reg = "SD",
    ss = "N",
    route = "EV",
    method = 1
)
```

Arguments

X	input dataset name (after Time Deviation Correction Rules have been applied by correct.time)
by	column names in x indicating grouping variables
nomtimevar	variable name containing the nominal sampling time after dose
tau	dosing interval (for multiple dosing); NA (default) for if single dose; x tau overrides
tstart	start time of partial AUC (start>0); NA (default) if not requested; x\$tstart overrides
tend	end time of partial AUC; NA (default) if not requested; x\$tend overrides

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teval	user selected AUC interval; NA (default) if not requested; x\$teval overrides
th	lamdba_z information for each curve; like output of est.thalf
reg	regimen, "sd" or "md"; x\$reg overrides
SS	is steady state reached (y/n); x\$ss overrides
route	route of drug administration ("EV","IVB","IVI"); x\$route overrides
method	method for trapezoidal rule; x\$method overrides
	• 1: linear up - linear down
	• 2: linear up - logarithmic down
	• 3: linear before first Tmax, logarithmic after first Tmax

Details

- Use interpolation if there is a measurable concentration BEFORE and AFTER the missing concentration
- Use extrapolation if there is NO measurable concentration AFTER the missing concentration
- Set missing concentration at predose to 0 (SD, non-endogenous) or value at t=TAU (steady state only)
- Set missing concentration at t=TAU to value at t=0 (steady state only)

The following Concentration Deviation Correction Rules will be applied to critical time points (t=0, tau, tstart, tend, teval), if needed:

Rule	Regimen	Description
SDC-1	sd	Set concentration to 0 (only non-endogenous compounds)
SDC-2	sd	impute missing concentration by interpolation
SDC-3	sd	impute missing concentration by extrapolation
SDC-4	sd (IVB)	impute missing concentration by back-extrapolation
MDC-1	md	impute missing concentration by existing conc at t=0 or t=tau (only if steady state has been reached)
MDC-2	md	impute missing concentration by interpolation
MDC-3	md	impute missing concentration by extrapolation
MDC-4	md (IVB)	impute missing concentration by back-extrapolation

Value

a dataset with missing concentrations imputed. The following variables are added:

Variable	Description
crule.nr	correction rule number
crule.txt	text explaining what was altered
applies.to.conc	lists all critical time points to which the concentration correction rule applies

Examples

```
example(correct.time)
```

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```
x %<>% mutate(ss = 'N', route = 'EV')
# route redefined for completeness
x %<>% correct.conc(by = 'subject') # ignoring th
x %>% head
```

correct.log

Impute Concentrations Below the Limit of Quantitation

Description

Imputes LOQ values according to the chosen LOQ substitution rule.

Usage

```
correct.loq(
    x,
    by = character(0),
    nomtimevar = "ntad",
    timevar = "time",
    depvar = "dv",
    bloqvar = "bloq",
    loqvar = "loq",
    loqrule = 1
)
```

Arguments

input dataset name contains all uncorrected data, including LOQ Х by column names in x indicating grouping variables nomtimevar variable name containing the nominal sampling time after dose variable name containing the actual sampling time after dose timevar variable name containing the dependent variable (e.g., concentration) depvar bloqvar variable name containing the BLOQ flag (0: no, 1: yes) loqvar variable name containing the LOQ value loqrule rule number to be applied to the LOQ values in the curve. x\$loqrule overrides if provided

- 1: 0 before first measurable concentration (FMC); NA after FMC
- 2: 0 before FMC; 0 after FMC
- 3: 0 before FMC; 0.5xLOQ for first consecutive LOQ after FMC, NA for other LOQ
- 4: 0 before FMC; 0.5xLOQ for first consecutive LOQ after FMC, 0 for other LOQ

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Details

Imputations will be applied to the original depvar (no new concentration variable will be created).

Value

A dataset with imputed BLOQ concentrations using the chosen imputation rule

Examples

```
library(magrittr)
library(dplyr)
library(qpNCA)
x <- Theoph
ntad <- c(0,0.25,0.5,1,2,4,5,7,9,12,24)
for(i in 1:nrow(x)){
  time <- x$Time[[i]]</pre>
  delta <- abs(ntad - time)</pre>
  best <- min(delta)</pre>
  index <- match(best, delta)</pre>
  nom <- ntad[[index]]</pre>
  x$ntad[[i]] \leftarrow nom
rm(list = c('time', 'delta', 'best', 'index', 'nom', 'i', 'ntad'))
x %<>% rename(time = Time, dv = conc)
x %<>% mutate(bloq = ifelse(dv==0,1,0), loq = 0.01, tad = time, loqrule = 1,
               subject=as.numeric(Subject), ntad=as.numeric(ntad))
x %>% head
x %<>% correct.loq('subject')
x %>% head
```

correct.time

Correct Concentrations for Time Deviations

Description

Corrects concentrations at critical, but deviating time points (e.g, predose, TAU, start and end of user selected AUC interval), and adds missing records at these critical time points.

Usage

```
correct.time(
    x,
    by = character(0),
    nomtimevar = "ntad",
    timevar = "time",
    depvar = "dv",
    tau = NA,
```

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```
tstart = NA,
tend = NA,
teval = NA,
th = NA,
reg = "SD",
method = 1
)
```

Arguments

input dataset name (after LOQ values have been imputed by correct.log) Х column names in x indicating grouping variables by variable name containing the nominal sampling time after dose nomtimevar variable name containing the actual sampling time after dose timevar variable name containing the dependent variable (e.g., concentration) depvar dosing interval (for multiple dosing); NA (default) if single dose; x\$tau overtau rides tstart start time of partial AUC (start>0); NA (default) if not requested; x\$tstart overend time of partial AUC; NA (default) if not requested; x\$tend overrides tend teval user selected AUC interval, starting at t=0; NA (default) if not requested; x\$teval overrides th lamdba_z information for each curve; like output of est.thalf regimen, "sd" or "md"; x\$reg overrides reg method method for trapezoidal rule; x\$method overrides if provided • 1: linear up - linear down • 2: linear up - logarithmic down

Details

 Records with missing NOMINAL time will be removed and this must be corrected before the function is called

• 3: linear before first Tmax, logarithmic after first Tmax

- If a record at the critical time point is missing, add it and set time to nominal time and set dv conc to NA
- Use interpolation if there is a measurable concentration AFTER the nominal time point (i.e. sample is taken too late)
- Use extrapolation if there is NO measurable concentration AFTER the nominal time point (i.e. sample is taken too early)
- Set deviating time at predose after single dose to 0
- Original time and conc will be kept in original variables.

The following Time Deviation Correction Rules will be applied to critical time points (t = 0, tau, tstart, tend, teval), if needed:

est.thalf

Rule	Regimen	Description	Applied to
SDT-1	sd	Set actual time to 0	t = 0
SDT-2	sd	Correct concentration at deviating time by interpolation	t = tau, tstart, tend, teval
SDT-3	sd	Correct concentration at deviating time by extrapolation	t = tau, tend, teval
MDT-1	md	If predose sample taken after dosing, set actual time to 0 and conc to NA	t = 0
MDT-2	md	Correct concentration at deviating time by interpolation (too late)	t = tau, tstart, tend, teval
MDT-3	md	Correct concentration at deviating time by extrapolation (too early)	t = 0, tau, tend, teval
MDT-3a	md	Set actual time to zero if concentration is BLOQ (too early)	t = 0

Value

a dataset with time deviation corrections applied (timevar and depvar adapted). The following variables are added:

Variable	Description
create.nr	is a missing record created?
create.txt	explanation of what is created
trule.nr	correction rule number
trule.txt	text explaining what was altered
applies.to.time	lists all critical time points to which the time deviation rule applies
time.tau, conc.tau	time and conc, corrected for AUCtau calculation
time.teval, conc.teval	time and conc, corrected for AUCteval calculation (AUC0-teval)
time.part, conc.part	time and conc, corrected for partial AUC calculation (AUCstart-end, start>0)
time.lastall, conc.lastall	time and conc, corrected for AUClast and AUCall calculation
t0.flag, tau.flag, tstart.flag, tend.flag, teval.flag	flags for what timepoint the correction was needed

The following are preserved if present in x: tau, tstart, tend, teval, reg, ss, route, method.

Examples

```
example(calc.ctmax)
x %<>% mutate(reg = 'SD', method = 1, route = 'EV')
# route not used yet, but still preserved
x %<>% correct.time(by = 'subject', th = th)
x %>% head
```

est.thalf Calculate Lambda_z and Elimination Half-life

Description

Calculates lambda_z and thalf for each PK curve identified using by.

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Usage

```
est.thalf(
   x,
   by = character(0),
   timevar = "time",
   depvar = "dv",
   includeCmax = "Y",
   exclvar = NA
)
```

Arguments

x a dataset

by column names in x indicating grouping variables

timevar variable name containing the actual sampling time after dose

depvar variable name containing the dependent variable (e.g., concentration)

includeCmax include results of regression including Cmax in selection? (y/n); x\$includeCmax

overrides if provided

exclvar a variable name containing information about points to be excluded (these should

have exclvar = 1)

Details

The function starts with the last three sample points and performs log-linear regression on it. It then adds one sampling point at a time (including and ending at tmax) and performs the regression again. The results of the regression with the highest adjusted R-squared are returned.

Visual outliers can be excluded from the regression analysis.

Value

a dataset with estimates for each regression analysis in one observation. The following parameters are available.

- no.points number of data points used in the regression analysis
- intercept estimated intercept
- lambda_z -1*estimated slope
- r.squared square of the correlation coefficient
- adj.r.squared adjusted square of the correlation coefficient
- thalf elimination half-life
- start_th time of first sample included in the thalf estimation
- end_th time of last sample included in the thalf estimation
- includeCmax include results of regression including Cmax in selection? (y/n)
- points_excluded are time points excluded from the half-life estimation? (y/n)

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Examples

```
example(correct.loq)
x %<>% mutate(includeCmax = 'Y')
th <- x %>% est.thalf(by='subject',exclvar=)
th %>% head
```

filenamefun

Create File Name for Regression Plots

Description

Creates file name for regression plots (*.png) from by-variables in plot_reg function

Usage

```
filenamefun(x, by)
```

Arguments

x data.frame

by column names in x indicating grouping variables

Value

character

interpol

Interpolate Concentrations

Description

Interpolates concentrations. Used by correct.xx functions to interpolate concentrations. Uses linear interpolation unless method is 2 (log down), c1 > c2, and both concentrations are non-zero.

Usage

```
interpol(c1 = NA, c2 = NA, t1 = NA, t2 = NA, t3 = NA, method = 1)
```

lag_lead

Arguments

c1	concentration 1 lagconc
c2	concentration 2 leadconc
t1	time 1 tiem where conc should be calculated
t2	time 2 lagtime
t3	time 3 leadtime
method	calculation method (1, 2, or 3)

lag_lead

Estimate Lagging and Leading Times and Concentrations

Description

Estimates lagging and leading times and concentrations. Used by correct.xx functions to estimate lagging and leading timepoints and concentrations for each timepoint.

Usage

```
lag_lead(
    x,
    nomtimevar1 = NA,
    depvar1 = NA,
    timevar1 = NA,
    lagc = NA,
    lagt = NA,
    leadc = NA,
    leadt = NA,
    ...
)
```

Arguments

```
data.frame
Х
                  column name in x indicating nominal time after dose
nomtimevar1
                  column name in x indicating concentration
depvar1
timevar1
                  column name in x indicating actual time after dose
                  concentration at previous sampling time
lagc
lagt
                  previous sampling time
leadc
                  concentration at next sampling time
                  next sampling time
leadt
                  ignored
. . .
```

Value

data.frame

plot_reg

plot_reg	Plot Regression Curves	

Description

Plots regression curves for each set of records defined using by. A log-linear plot will be made for each curve.

Usage

```
plot_reg(
    x,
    by = character(0),
    th = NA,
    bloqvar = "bloq",
    timevar = "tad",
    depvar = "dv",
    timelab = "timevar",
    deplab = "depvar",
    exclvar = NA,
    plotdir = NA,
    ...
)
```

Arguments

X	input dataset name
by	column names in x indicating grouping variables
th	file name of file with half-life estimation information for each curve
bloqvar	variable name containing the BLOQ flag (0: no, 1: yes)
timevar	variable name containing the actual sampling time after dose
depvar	variable name containing the dependent variable (e.g., concentration)
timelab	X-axis label (default: "timevar")
deplab	Y-axis label (default: "depvar")
exclvar	variable name containing information about points to be excluded (these should have exclvar = 1)
plotdir	directory where individual plot files will be saved
	ignored

Details

If elimination half-life was estimated for that curve, the following will be indicated in the plot:

• Cmax (Yellow, even if no half-life was estimated)

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- points used in regression and resulting regression line (green)
- points excluded from regression (red crossed)
- estimate of elimination half-life and adjusted R-squared

Input dataset:

- uncorrected dataset, used for half-life estimation
- dataset containing results of the half-life estimation

Value

(invisible) plotdir. If the attribute 'plotdir' is empty, plots will be generated in standard output, otherwise plots will be saved as PNG file in the designated directory.

Examples

```
example(est.thalf)
x %>% filter(dv > 0) %>% plot_reg(by = 'subject', th = th)
```

qpNCA

Perform Non-compartmental Analysis

Description

Consecutively executes the following NCA steps:

- correct.log impute LOQ values
- est.thalf calculate lambda_z and half-life
- plot_reg plot each regression curve
- calc.ctmax calculate Cmax and Tmax
- correct.time correct time deviations at critical time points
- correct.conc impute missing concentrations at critical time points
- tab.corr tabulate data alterations
- calc.par calculates parameters not dependent on lambda_z
- calc.par.th calculates parameters dependent on lambda_z

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Usage

```
qpNCA(
 Х,
  by = character(0),
  nomtimevar = "ntad",
  timevar = "time",
  depvar = "dv",
 bloqvar = "bloq",
  loqvar = "loq",
  loqrule = 1,
  includeCmax = "Y",
  exclvar = NA,
 plotdir = NA,
  timelab = "timevar",
  deplab = "depvar",
  tau = NA,
  tstart = NA,
  tend = NA,
  teval = NA,
  covariates = NA,
  dose = "dose",
  factor = 1,
  reg = "SD",
  ss = "N",
 route = "EV",
 method = 1
)
```

Arguments

X	input dataset name
by	column names in x indicating grouping variables
nomtimevar	variable name containing the nominal sampling time after dose
timevar	variable name containing the actual sampling time after dose
depvar	variable name containing the dependent variable (e.g., concentration)
bloqvar	variable name containing the BLOQ flag (0: no, 1: yes)
loqvar	variable name containing the LOQ value
loqrule	rule number to be applied to the LOQ values in the curve; x\$loqrule overrides if provided
includeCmax	include Cmax in half-life estimation? (y/n); x =includeCmax overrides if provided
exclvar	variable name indicating points to be excluded in half-life estimation (these should have $exclvar = 1$)
plotdir	directory where regression plots (.PNG) will be saved; NA gives default location, NULL skips regression plots

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timelab label for time axis in regression plots

deplab label for dependent variable axis in regression plots

tau dosing interval (for multiple dosing); NA (default) if single dose; x\$tau over-

rides

tstart start time of partial AUC (start>0); NA (default) if not requested; x\$tstart over-

rides

tend end time of partial AUC; NA (default) if not requested; x\$tend overrides

teval user selected AUC interval (starting at t=0); NA (default) if not requested; x\$teval

overrides

covariates covariates dataset; Must contain the dose variable

dose variable containing the dose amount

factor conversion factor for CL and V calculation (e.g. dose in mg, conc in ng/mL,

factor=1000); x\$factor overrides if provided

reg regimen, "SD" or "MD"; x\$reg overrides if provided ss is steady state reached (y/n); x\$ss overrides if provided

route route of drug administration ("EV","IVB", "IVI"); x\$route overrides if provided

method method for trapezoidal rule; x\$method overrides if provided

• 1: linear up - linear down

• 2: linear up - logarithmic down

• 3: linear before first Tmax, logarithmic after first Tmax

Value

(list)

- covariates covariates selected with the covariates argument
- half_life linear regression parameters
- ct_corr the time and concentration corrected dataset
- corrections descriptions of the corrections applied
- pkpar all estimated PK parameters
- plots generated plots

Examples

```
library(magrittr)
library(dplyr)
library(qpNCA)
x <- Theoph
ntad <- c(0,0.25,0.5,1,2,4,5,7,9,12,24)
for(i in 1:nrow(x)){
  time <- x$Time[[i]]
  delta <- abs(ntad - time)
  best <- min(delta)</pre>
```

tab.corr 21

tab.corr

Tabulate Corrections

Description

Tabulates what records were added, time deviations and concentration imputations were applied, for each subject.

Usage

```
tab.corr(x, by = character(0), nomtimevar = "time")
```

Arguments

x concentration dataset created by the correct.time and correct.conc functions, containing time and conc corrected data

by column names in x indicating grouping variables

nomtimevar column in x containing the nominal time after dose

Value

dataset with applied corrections (rule number and rule text) listed by by-variable(s) and nominal time

Examples

```
example(correct.conc)
corrtab <- x %>% tab.corr(by = 'subject')
corrtab %>% head
```

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titlefun

Create Title for Regression Plots

Description

Creates title for regression plots in plot_reg() using by-variables.

Usage

```
titlefun(x, by)
```

Arguments

x dataset containing concentration-time information of the current curve

by column names in x indicating grouping variables

Value

character

trap

Calculate Area Under the Curve Using Trapezoids

Description

Calculates AUC using the trapezoidal method. Assumes data represent a single profile. Despite choice of method, only linear interpolation is used for areas of intervals beginning or ending with y: 0.

Usage

```
trap(x = NA, y = NA, method = 1)
```

Arguments

x variable, i.e. time

y y variable, i.e. concentration

method method:

- 1: linear up linear down
- 2: linear up logarithmic down
- 3: linear before Tmax, logarithmic after Tmax

Value

```
area (length-one numeric)
```

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trapm

Calculate Area Under the Moment Curve Using Trapezoids

Description

Calculates AUMC using the trapezoidal method. Assumes data represent a single profile. Despite choice of method, only linear interpolation is used for areas of intervals beginning or ending with y: 0.

Usage

```
trapm(x = NA, y = NA, method = 1)
```

Arguments

x variable names of x coordinates y variable names of y coordinates

method method:

- 1: linear up linear down
- 2: linear up logarithmic down
- 3: linear before Tmax, logarithmic after Tmax

Value

area (length-one numeric)

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