

# Package ‘ncappc’

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**Title** NCA Calculations and Population Model Diagnosis

**Version** 1.0.0

**Description** A flexible tool that can perform  
(i) traditional non-compartmental analysis (NCA) and  
(ii) Simulation-based posterior predictive checks for population  
pharmacokinetic (PK) and/or pharmacodynamic (PKPD) models using  
NCA metrics. The methods are described in Acharya et al. (2016)  
<[doi:10.1016/j.cmpb.2016.01.013](https://doi.org/10.1016/j.cmpb.2016.01.013)>.

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2.0.0), grDevices, stats, bookdown

**Suggests** testthat

**License** GPL-3

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<https://uupharmacometrics.github.io/ncappc/>

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---

calc.stat	<i>Calculates a set of statistics for a given array of numbers.</i>
-----------	---

---

### Description

**calc.stat** calculates a set of statistics for a given array of numbers.

### Usage

```
calc.stat(x)
```

### Arguments

x                    a numeric array

### Details

**calc.stat** calculates a set of statistics for a given array of numbers. The calculated statistics are

- Ntot = length of the array
- Nunique = Number of unique elements
- Min = Minimum value of the array
- Max = Maximum value of the array
- Mean = Mean value of the array
- Median = Median value of the array

- SD = Standard deviation value of the array
- SE = Standard error value of the array
- CVp = Percent coefficient of variation of the array
- CI95u = Upper limit of the 95% confidence interval of the array
- cI95l = Lower limit of the 95% confidence interval of the array
- gMean = Geometric mean value of the array
- gCVp = Geometric percent coefficient of variation of the array

**Value**

An array of calculated statistics of a given set of numbers

---

dv.plot

*Plots drug plasma concentration vs time data*

---

**Description**

**dv.plot** plots DV vs Time data.

**Usage**

```
dv.plot(  
  df,  
  xvar = "Time",  
  yvar = "Conc",  
  obsLog = FALSE,  
  myXlab = "Time",  
  myYlab = "Concentration",  
  color = NULL,  
  group = NULL,  
  guide = TRUE,  
  onlyLin = FALSE,  
  onlyLog = FALSE,  
  XYlog = FALSE,  
  STRATY = ".",  
  STRATX = ".",  
  myYBr = waiver(),  
  myXBr = waiver(),  
  myYBrLog = waiver(),  
  myXBrLog = waiver(),  
  myYlim = NULL,  
  myXlim = NULL,  
  myYlimLog = NULL,  
  myXlimLog = NULL,  
  title = NULL  
)
```

**Arguments**

df	A data frame to be used for the plot
xvar	is the independent variable, default is " <b>TIME</b> "
yvar	is the dependent variable, default is " <b>DV</b> "
obsLog	is a logical variable (TRUE, FALSE). If TRUE, concentration in observed data is assumed to be in logarithmic scale. Default is FALSE
myXlab	is the x-axis label, default is " <b>Time</b> "
myYlab	is the y-axis label, defaults is " <b>Concentration</b> "
color	is the column name of the color stratification variable, e.g. "DOSEF". Default is <b>NULL</b>
group	is the column name of the variable used to group data, default is " <b>ID</b> "
guide	if <b>TRUE</b> , show guide, default is <b>TRUE</b>
onlyLin	if <b>TRUE</b> , presents only the linear version of the plot, default is <b>FALSE</b>
onlyLog	if <b>TRUE</b> , presents only the log version of the plot, default is <b>FALSE</b>
XYlog	if <b>TRUE</b> , both X and Y axes of the log version of the plot is shown on the logarithmic scale; if <b>FALSE</b> , only the Y-axis is shown on the logarithmic scale. Default is <b>FALSE</b> .
STRATY	is the row stratification variable, default is "."
STRATX	is the column stratification variable, default is "."
myYBr	are the breaks for the Y-axis for the linear plot
myXBr	are the breaks for the X-axis for the linear plot
myYBrLog	are the breaks for the Y-axis for the log plot
myXBrLog	are the breaks for the X-axis for the log plot
myYlim	sets Y-axis limits for the linear plot
myXlim	sets X-axis limits for the linear plot
myYlimLog	sets the Y-axis limit for the log plot
myXlimLog	sets the X-axis limit for the log plot
title	The title of the plot.

**Details**

**dv.plot** plots DV vs Time data

**Value**

returns a graphical object created by arrangeGrob function

---

 est.nca

*Estimate individual NCA metrics.*


---

## Description

Estimates a comprehensive set of NCA metrics for a given individual using concentration vs. time data.

## Usage

```
est.nca(
  time,
  conc,
  backExtrp = FALSE,
  negConcExcl = FALSE,
  doseType = "ns",
  adminType = "extravascular",
  doseAmt = NULL,
  method = "linearup-logdown",
  AUCTimeRange = NULL,
  LambdaTimeRange = NULL,
  LambdaExclude = NULL,
  doseTime = doseTime,
  Tau = NULL,
  TI = NULL,
  simFile = NULL,
  dset = "obs",
  onlyNCA = FALSE,
  extrapolate = FALSE,
  sparse_compute = FALSE,
  force_extrapolate = FALSE,
  ...
)
```

## Arguments

time	Numeric array for time
conc	Numeric array for concentration
backExtrp	If back-extrapolation is needed for AUC (TRUE or FALSE) ( <b>FALSE</b> )
negConcExcl	Exclude -ve conc ( <b>FALSE</b> )
doseType	Steady-state (ss) or non-steady-state (ns) dose (" <b>ns</b> ")
adminType	Route of administration (iv-bolus,iv-infusion,extravascular) (" <b>extravascular</b> ")
doseAmt	Dose amounts (" <b>NULL</b> ")

method	Method to estimate AUC. The "linear" method applies the linear trapezoidal rule to estimate the area under the curve. The "log" method applies the logarithmic trapezoidal rule to estimate the area under the curve. The "linearup-logdown" method applies the linear trapezoidal rule to estimate the area under the curve for the ascending part of the curve and the logarithmic trapezoidal rule to estimate the area under the curve for the descending part of the curve.
AUCTimeRange	User-defined window of time used to estimate AUC ("NULL")
LambdaTimeRange	User-defined window of time to estimate elimination rate-constant ("NULL")
LambdaExclude	User-defined excluded observation time points for estimation of elimination rate-constant ("NULL")
doseTime	Dose time prior to the first observation for steady-state data (NULL)
Tau	Dosing interval for steady-state data ("NULL")
TI	Infusion duration ("NULL")
simFile	Name of the simulated concentration-time data if present ("NULL")
dset	Type, i.e., observed or simulated concentration-time data set ("obs" or "sim") ("obs")
onlyNCA	If TRUE only NCA is performed and ppc part is ignored although simFile is not NULL. Default is FALSE
extrapolate	Should the function extrapolate from the last observation to infinity?
sparse_compute	Should NCA metrics be computed even with only one sample?
force_extrapolate	Extrapolate AUC_inf with sparse data. Sparse data is defined as fewer than 3 points after Cmax for non-bolus input. In that case, Cmax is included in the extrapolation and only one extra point is required for extrapolation.
...	Arguments passed from other functions. Not used.

## Details

**est.nca** estimates a comprehensive set of NCA metrics using the concentration-time profile of an individual. NCA metrics are estimated according to traditional PK calculations. The names of the various NCA metrics estimated in this package are assigned mainly following the names used in WinNonlin. This package accepts any of the three different types of drug administration, (i) iv-bolus, (ii) iv-infusion and (iii) extravascular; **ncappc** also can accept both non-steady state and steady-state data. The NCA metrics that are estimated and reported by **ncappc** are listed below.

- **C0** is the initial concentration at the dosing time. It is the observed concentration at the dosing time, if available. Otherwise it is approximated using the following rules.
- **Cmax, Tmax and Cmax\_D** are the value and the time of maximum observed concentration, respectively. If the maximum concentration is not unique, the first maximum is used. For steady state data, The maximum value between the dosing intervals is considered. Cmax\_D is the dose normalized maximum observed concentration.
- **Clast and Tlast** are the last measurable positive concentration and the corresponding time, respectively.

- **AUClast** is the area under the concentration vs. time curve from the first observed to last measurable concentration.
- **AUMClast** is the first moment of the concentration vs. time curve from the first observed to last measurable concentration.
- **MRTlast** is the mean residence time from the first observed to last measurable concentration.
- **No\_points\_Lambda\_z** is the number of observed data points used to determine the best fitting regression line in the elimination phase.
- **AUC\_pBack\_Ext\_obs** is the percentage of AUCINF\_obs that is contributed by the back extrapolation to estimate C0.
- **AUC\_pBack\_Ext\_pred** is the percentage of AUCINF\_pred that is contributed by the back extrapolation to estimate C0.
- **AUClower\_upper** is the AUC under the concentration-time profile within the user-specified window of time provided as the "AUCTimeRange" argument. In case of empty "AUCTimeRange" argument, AUClower\_upper is the same as AUClast.
- **Rsqr, Rsqr\_adjusted and Corr\_XY** are regression coefficient of the regression line used to estimate the elimination rate constant, the adjusted value of Rsqr and the square root of Rsqr, respectively.
- **Lambda\_z** is the elimination rate constant estimated from the regression line representing the terminal phase of the concentration-time data.
- **Lambda\_lower and Lambda\_upper** are the lower and upper limit of the time values from the concentration-time profile used to estimate Lambda\_z, respectively, in case the "Lambda-TimeRange" is used to specify the time range.
- **HL\_Lambda\_z** is terminal half-life of the drug.
- **AUCINF\_obs and AUCINF\_obs\_D** are AUC estimated from the first sampled data extrapolated to infinity and its dose normalized version, respectively. The extrapolation in the terminal phase is based on the last observed concentration Clast\_obs.
- **AUC\_pExtrap\_obs** is the percentage of the AUCINF\_obs that is contributed by the extrapolation from the last sampling time to infinity.
- **AUMCINF\_obs** is AUMC estimated from the first sampled data extrapolated to infinity. The extrapolation in the terminal phase is based on the last observed concentration.
- **AUMC\_pExtrap\_obs** is the percentage of the AUMCINF\_obs that is contributed by the extrapolation from the last sampling time to infinity.
- **Vz\_obs** is the volume of distribution estimated based on total AUC
- **Cl\_obs** is total body clearance.
- **AUCINF\_pred and AUCINF\_pred\_D** are AUC from the first sampled data extrapolated to infinity and its dose normalized version, respectively. The extrapolation in the terminal phase is based on the last predicted concentration obtained from the regression line used to estimate Lambda\_z (Clast\_pred).
- **AUC\_pExtrap\_pred** is the percentage of the AUCINF\_pred that is contributed by the extrapolation from the last sampling time to infinity.
- **AUMCINF\_pred** is AUMC estimated from the first sampled data extrapolated to infinity. The extrapolation in the terminal phase is based on the last predicted concentration obtained from the regression line used to estimate Lambda\_z (Clast\_pred).

- **AUMC\_pExtrap\_pred** is the percentage of the AUMCINF\_pred that is contributed by the extrapolation from the last sampling time to infinity.
- **Vz\_pred** is the volume of distribution estimated based on AUCINF\_pred.
- **Cl\_pred** is the total body clearance estimated based on AUCINF\_pred.
- **MRTINF\_obs** is the mean residence time from the first sampled time extrapolated to infinity based on the last observed concentration (Clast\_obs).
- **MRTINF\_pred** is the mean residence time from the first sampled time extrapolated to infinity based on the last predicted concentration obtained from the regression line used to estimate Lambda\_z (Clast\_pred).
- **Tau** is the dosing interval for steady-state data.
- **Cmin and Tmin** are the minimum concentration between 0 and Tau and the corresponding time, respectively.
- **Cavg** is the average concentration between 0 and Tau for steady-state data.
- **AUCtau and AUMCtau** are AUC and AUMC between 0 and Tau for steady-state data.
- **Clss** is an estimate of the total body clearance for steady-state data.
- **Vss\_obs and Vss\_pred** are estimated volume of distribution at steady-state based on Clast\_obs and Clast\_pred, respectively.
- **p\_Fluctuation** is the percentage of the fluctuation of the concentration between 0 and Tau for steady-state data.
- **Accumulation\_Index** is  $1/(1 - e^{-(\lambda_z * \tau)})$

### Value

An array of estimated NCA metrics

---

histobs.plot

*Plots histogram of selected set of NCA metrics.*

---

### Description

**histobs.plot** plots histogram of selected set of NCA metrics (e.g. AUClast, AUCINF\_obs, Cmax and Tmax).

### Usage

```
histobs.plot(
  plotData,
  figlbl = NULL,
  param = c("AUClast", "AUCINF_obs", "Cmax", "Tmax"),
  cunit = NULL,
  tunit = NULL,
  spread = "npi"
)
```

**Arguments**

plotData	A data frame with the estimated NCA metrics
figlbl	Figure label based on dose identifier and/or population stratifier ( <b>NULL</b> )
param	A character array of the NCA metrics. The allowed NCA metrics for this histograms are "AUClast", "AUClower_upper", "AUCINF_obs", "AUCINF_pred", "AUMClast", "Cmax", "Tmax" and "HL_Lambda_z". ( <b>c("AUClast", "AUCINF_obs", "Cmax", "Tmax")</b> )
cunit	Unit for concentration (default is <b>NULL</b> )
tunit	Unit for time (default is <b>NULL</b> )
spread	Measure of the spread of simulated data (ppi (95% parametric prediction interval) or np_i (95% nonparametric prediction interval)) ( <b>"npi"</b> )

**Details**

**histobs.plot** plots histogram of selected set of NCA metrics. The allowed NCA metrics for this histograms are "AUClast", "AUClower\_upper", "AUCINF\_obs", "AUCINF\_pred", "AUMClast", "Cmax", "Tmax" and "HL\_Lambda\_z". By default, this function produces histogram of AUClast, AUCINF\_obs, Cmax and Tmax.

**Value**

returns a graphical object created by arrangeGrob function

---

histpop.plot	<i>Plots population histogram of the NCA metrics selected for model diagnosis.</i>
--------------	--

---

**Description**

**histpop.plot** plots population histogram of the NCA metrics selected for model diagnosis (e.g. AUClast, AUCINF\_obs, Cmax and Tmax).

**Usage**

```
histpop.plot(
  obsdata = outData,
  simdata = smedianData,
  figlbl = NULL,
  param = c("AUClast", "Cmax"),
  cunit = NULL,
  tunit = NULL,
  spread = "npi"
)
```

**Arguments**

obsdata	Data frame with the values of the NCA metrics estimated from the observed data
simdata	Data frame with the values of the NCA metrics estimated from the simulated data
figlbl	Figure label based on dose identifier and/or population stratifier ( <b>NULL</b> )
param	A character array of the NCA metrics. The allowed NCA metrics for this histograms are "AUClast", "AUClower_upper", "AUCINF_obs", "AUCINF_pred", "AUMClast", "Cmax", "Tmax" and "HL_Lambda_z". ( <b>c("AUClast", "Cmax")</b> )
cunit	Unit for concentration (default is <b>NULL</b> )
tunit	Unit for time (default is <b>NULL</b> )
spread	Measure of the spread of simulated data (ppi (95% parametric prediction interval) or npci (95% nonparametric prediction interval)) ( <b>"npi"</b> )

**Details**

**histpop.plot** plots histogram of the NCA metrics selected for the model diagnosis and compares with the corresponding metrics estimated from the observed data. The allowed NCA metrics for this histograms are "AUClast", "AUClower\_upper", "AUCINF\_obs", "AUCINF\_pred", "AUMClast", "Cmax", "Tmax" and "HL\_Lambda\_z". By default, this function produces histogram of AUClast and Cmax.

**Value**

returns a graphical object created by arrangeGrob function

---

nca.check.obs	<i>Check observed data</i>
---------------	----------------------------

---

**Description**

**nca.check.obs** Checks observed data for compatibility with ncappc and processes the data with various filtering criteria.

**Usage**

```
nca.check.obs(
  obsData,
  idNmObs = "ID",
  timeNmObs = "TIME",
  concNmObs = "DV",
  doseType = "ns",
  doseTime = NULL,
  Tau = NULL,
  filterNm = NULL,
  filterExcl = NULL,
```

```

    str1Nm = NULL,
    str1 = NULL,
    str2Nm = NULL,
    str2 = NULL,
    str3Nm = NULL,
    str3 = NULL,
    AUTimeRange = NULL,
    LambdaTimeRange = NULL,
    adminType = "extravascular",
    TI = NULL,
    doseAmtNm = NULL,
    dateColNm = NULL,
    dateFormat = NULL,
    timeFormat = "number",
    concUnit = NULL,
    timeUnit = NULL,
    doseUnit = NULL,
    blqNm = NULL,
    blqExcl = 1,
    evid = TRUE,
    evidIncl = 0,
    mdv = FALSE
)

```

### Arguments

obsData	Observed concentration-time data.
idNmObs	Column name for ID in observed data. Default is " <b>ID</b> ".
timeNmObs	Column name for time in observed data. Default is " <b>TIME</b> ".
concNmObs	Column name for concentration in observed data. Default is " <b>DV</b> ".
doseType	Steady-state (ss) or non-steady-state (ns) dose. Default is " <b>ns</b> ".
doseTime	Dose time prior to the first observation for steady-state data. Default is NULL.
Tau	Dosing interval for steady-state data. Default is NULL.
filterNm	Column name to filter data. Default is NULL.
filterExcl	Row exclusion criteria based on the column defined by filterNm. This can be numeric value or logical condition (e.g. c(1, 2, "<20", ">=100", "!=100")). Default is NULL.
str1Nm	Column name for 1st level population stratifier. Default is NULL.
str1	Stratification ID of the members within 1st level stratification (e.g. c(1,2)). Default is NULL.
str2Nm	Column name for 2nd level population stratifier. Default is NULL.
str2	Stratification ID of the members within 2nd level stratification (e.g. c(1,2)). Default is NULL.
str3Nm	Column name for 3rd level population stratifier. Default is NULL.

str3	Stratification ID of the members within 3rd level stratification (e.g c(1,2)). Default is NULL
AUCtimeRange	User-defined window of time used to estimate AUC. Default is NULL
LambdaTimeRange	User-defined window of time to estimate elimination rate-constant. This argument lets the user to choose a specific window of time to be used to estimate the elimination rate constant (Lambda) in the elimination phase. The accepted format for the input to this argument is a numeric array of two elements; c(14, 24) will estimate the Lambda using the data within the time units 14 to 24. Default is NULL
adminType	Route of administration. Allowed options are iv-bolus, iv-infusion or extravascular. Default is " <b>extravascular</b> "
TI	Infusion duration. If TI is a single numeric value, TI is the same for all individuals. If TI is the name of a column with numeric data present in the data set, TI is set to the unique value of the column for a given individual. Default is NULL
doseAmtNm	Column name to specify dose amount. Default is NULL
dateColNm	column name for date if used (e.g. "Date", "DATE"). Default is NULL
dateFormat	date format (D-M-Y, D/M/Y or any other combination of D,M,Y). Default is NULL
timeFormat	time format (number, H:M, H:M:S). Default is " <b>number</b> "
concUnit	Unit of concentration (e.g. "ng/mL"). Default is NULL
timeUnit	Unit of time (e.g. "h"). Default is NULL
doseUnit	Unit of dose amount (e.g. "ng"). Default is NULL
blqNm	Name of BLQ column if used to exclude data. Default is NULL
blqExcl	Excluded BLQ value; either a numeric value or a logical condition (e.g. 1 or ">=1" or c(1,">3)"). Used only if the blqNm is not NULL. Default is " <b>1</b> "
evid	If TRUE EVID is used to filter data. Default is TRUE
evidIncl	Included values in EVID. Default is " <b>0</b> "
mdv	If TRUE MDV is used to include data when MDV=0. Default is FALSE

### Details

**nca.check.obs** Checks observed data for compatibility with ncappc.

### Value

A list of objects

---

nca.check.sim	<i>Check simulated data</i>
---------------	-----------------------------

---

## Description

**nca.check.sim** Checks simulated data for compatibility with ncappc and processes the data with various filtering criteria.

## Usage

```
nca.check.sim(
  simData,
  idNmSim = "ID",
  timeNmSim = "TIME",
  concNmSim = "DV",
  filterNm = NULL,
  filterExcl = NULL,
  str1Nm = NULL,
  str1 = NULL,
  str2Nm = NULL,
  str2 = NULL,
  str3Nm = NULL,
  str3 = NULL,
  adminType = "extravascular",
  TI = NULL,
  doseAmtNm = NULL,
  blqNm = NULL,
  blqExcl = 1,
  evid = TRUE,
  evidIncl = 0,
  mdv = FALSE
)
```

## Arguments

simData	Simulated concentration-time data.
idNmSim	Column name for ID in simulated data. Default is " <b>ID</b> "
timeNmSim	Column name for time in simulated data. Default is " <b>TIME</b> "
concNmSim	Column name for concentration in simulated data. Default is " <b>DV</b> "
filterNm	Column name to filter data. Default is NULL
filterExcl	Row exclusion criteria based on the column defined by filterNm. This can be numeric value or logical condition (e.g. c(1, 2, "<20", ">=100", "!=100")). Default is NULL
str1Nm	Column name for 1st level population stratifier. Default is NULL

str1	Stratification ID of the members within 1st level stratification (e.g c(1,2)). Default is NULL
str2Nm	Column name for 2nd level population stratifier. Default is NULL
str2	Stratification ID of the members within 2nd level stratification (e.g c(1,2)). Default is NULL
str3Nm	Column name for 3rd level population stratifier. Default is NULL
str3	Stratification ID of the members within 3rd level stratification (e.g c(1,2)). Default is NULL
adminType	Route of administration. Allowed options are iv-bolus, iv-infusion or extravascular. Default is " <b>extravascular</b> "
TI	Infusion duration. If TI is a single numeric value, TI is the same for all individuals. If TI is the name of a column with numeric data present in the data set, TI is set to the unique value of the column for a given individual. Default is NULL
doseAmtNm	Column name to specify dose amount. Default is NULL
blqNm	Name of BLQ column if used to exclude data. Default is NULL
blqExcl	Excluded BLQ value; either a numeric value or a logical condition (e.g. 1 or ">=1" or c(1,">3")). Used only if the blqNm is not NULL. Default is " <b>1</b> "
evid	If TRUE EVID is used to filter data. Default is TRUE
evidIncl	Included values in EVID. Default is " <b>0</b> "
mdv	If TRUE MDV is used to include data when MDV=0. Default is FALSE

### Details

**nca.check.sim** Checks simulated data for compatibility with ncappc.

### Value

A list of objects

---

nca.deviation.plot	<i>Plot individual deviation of NCA metrics estimated from observed and simulated data</i>
--------------------	--

---

### Description

**nca.deviation.plot** plots individual deviation of selected NCA metrics estimated from observed and simulated data.

**Usage**

```
nca.deviation.plot(
  plotdata,
  xvar = NULL,
  devcol = NULL,
  figlbl = NULL,
  spread = "npi",
  cunit = NULL,
  tunit = NULL
)
```

**Arguments**

plotdata	A data frame containing the scaled deviation values of the NCA metrics
xvar	x-variable against which the deviation data will be plotted ( <b>NULL</b> )
devcol	column names/numbers of the data frame containing deviation data ( <b>NULL</b> )
figlbl	Figure label based on dose identifier and/or population stratifier ( <b>NULL</b> )
spread	Measure of the spread of simulated data (ppi (95% parametric prediction interval) or npi (95% nonparametric prediction interval)) (" <b>npi</b> ")
cunit	Unit for concentration (default is <b>NULL</b> )
tunit	Unit for time (default is <b>NULL</b> )

**Details**

**nca.deviation.plot** plots individual deviation of selected NCA metrics estimated from observed and simulated data. This function requires three mandatory arguments, (i) deviation data, (ii) independent variable and (iii) dependent variables. The deviation of the NCA metrics values estimated from the observed and simulated data are scaled by the "spread" of the simulated metrics values. The "speed" of the simulated data is measured either by the standard deviation or the 95

**Value**

returns the data frame with the NPDE values based on the input data.

---

nca.ind.data

---

*Prepare individual PK data*


---

**Description**

**nca.ind.data** Extracts time-conc data for a given individual.

**Usage**

```
nca.ind.data(
  pkData,
  ID,
  dvLog = FALSE,
  dataType = "obs",
  idNm = "ID",
  timeNm = "TIME",
  concNm = "DV",
  adminType = "extravascular",
  TI = NULL,
  dateColNm = NULL,
  dateFormat = NULL,
  timeFormat = "number"
)
```

**Arguments**

pkData	PK concentration-time data.
ID	ID number of the individual.
dvLog	If TRUE concentration is in logarithmic scale. Default is FALSE
dataType	Indicates if the data is observed ("obs") or simulated ("sim"). Since the simulated data is assumed to be obtained from NONMEM output, DATE and clock time (dateColNm, dateFormat, timeFormat) are not used for time data. Default is <b>"obs"</b> .
idNm	Column name for ID in PK data. Default is <b>"ID"</b>
timeNm	Column name for time in PK data. Default is <b>"TIME"</b>
concNm	Column name for concentration in PK data. Default is <b>"DV"</b>
adminType	Route of administration. Allowed options are iv-bolus, iv-infusion or extravascular. Default is <b>"extravascular"</b>
TI	Infusion duration. If TI is a single numeric value, TI is the same for all individuals. If TI is the name of a column with numeric data present in the data set, TI is set to the unique value of the column for a given individual. Default is NULL
dateColNm	column name for date if used (e.g. "Date", "DATE"). Default is NULL
dateFormat	date format (D-M-Y, D/M/Y or any other combination of D,M,Y). Default is NULL
timeFormat	time format (number, H:M, H:M:S). Default is <b>"number"</b>

**Details**

**nca.ind.data** Extracts time-conc data for a given individual.

**Value**

A list of objects with time-conc data and individual infusion duration for iv-infusion data

---

nca.npde	<i>Calculates individual normalized prediction distribution errors (NPDE) from PDE data.</i>
----------	--

---

### Description

**nca.npde** calculates individual normalized prediction distribution errors (NPDE) of selected NCA metrics from the PDE data.

### Usage

```
nca.npde(pdedata, pdecol)
```

### Arguments

pdedata	A data frame containing the prediction distribution errors (PDE) of NCA metrics
pdecol	The range of column numbers in the data frame containing the PDE values, which will be used to calculate the corresponding NPDE

### Details

**nca.npde** calculates individual normalized prediction distribution errors (NPDE) of selected NCA metrics from PDE data. The deviation of each estimated NCA metrics is scaled by the "spread" of the simulated values. By default, this function calculates the NPDE values of all columns of the input data frame.

### Value

returns the data frame with the NPDE values based on the input data.

---

nca.npde.plot	<i>Plots population histogram of the NCA metrics selected for model diagnosis.</i>
---------------	--

---

### Description

**nca.npde.plot** plots individual NPDE values and histogram of the NPDE values within a population group

## Usage

```
nca.npde.plot(  
  plotdata,  
  xvar = NULL,  
  npdecol = NULL,  
  figlbl = NULL,  
  cunit = NULL,  
  tunit = NULL  
)
```

## Arguments

plotdata	Data frame with the values of the NPDE values of each individual for the NCA metrics
xvar	Name of the independent variable column against which NPDE values will be plotted
npdecol	Column names or column numbers of containing the NPDE values
figlbl	Figure label based on dose identifier and/or population stratifier ( <b>NULL</b> )
cunit	Unit for concentration (default is <b>NULL</b> )
tunit	Unit for time (default is <b>NULL</b> )

## Details

**nca.npde.plot** individual NPDE values and histogram of the NPDE values of NCA metrics within a population group.

## Value

returns a data frame with the mean and SD of population NPDE values of each NCA metric and two graphical objects created by arrangeGrob function for the individual and population histogram of the NPDE values

---

nca.pde.deviation.outlier

*Calculates individual prediction distribution errors (PDE) and scaled deviation of NCA metrics estimated from observed and simulated data. Identifies outlier to population PK model.*

---

## Description

**nca.pde.deviation.outlier** calculates individual prediction distribution errors (PDE) and scaled deviation of NCA metrics estimated from observed and simulated data. Identifies outlier to population PK model.

**Usage**

```
nca.pde.deviation.outlier(
  obsdata,
  simdata,
  idNm = "ID",
  id = NULL,
  spread = "npi",
  figlbl = NULL,
  calcparam = c("AUClast", "Cmax"),
  diagparam = c("AUClast", "Cmax"),
  cunit = NULL,
  tunit = NULL,
  noPlot = FALSE,
  onlyNCA = onlyNCA
)
```

**Arguments**

obsdata	A data frame containing the NCA metrics values estimated from the observed data
simdata	A data frame containing the NCA metrics values estimated from the simulated data
idNm	Column name for ID (" <b>ID</b> ")
id	ID of the individual whose data is being evaluated
spread	Measure of the spread of simulated data (ppi (95% parametric prediction interval) or npi (95% nonparametric prediction interval)) (" <b>npi</b> ")
figlbl	Figure label based on dose identifier and/or population stratifier, in addition to ID ( <b>NULL</b> )
calcparam	A character array of the NCA metrics used for calculations of PDE and deviation. The allowed NCA metrics for this histograms are "AUClast", "AUClower_upper", "AUCINF_obs", "AUCINF_pred", "AUMClast", "Cmax", "Tmax" and "HL_Lambda_z". ( <b>c("AUClast", "Cmax")</b> )
diagparam	A character array of the NCA metrics used for diagnostic test to detect outliers. The allowed NCA metrics for this histograms are "AUClast", "AUClower_upper", "AUCINF_obs", "AUCINF_pred", "AUMClast", "Cmax", "Tmax" and "HL_Lambda_z". ( <b>c("AUClast", "Cmax")</b> )
cunit	Unit for concentration (default is <b>NULL</b> )
tunit	Unit for time (default is <b>NULL</b> )
noPlot	Perform only NCA calculations without any plot generation ( <b>TRUE</b> , <b>FALSE</b> ) ( <b>FALSE</b> )
onlyNCA	If <b>TRUE</b> only NCA is performed and ppc part is ignored although simFile is not <b>NULL</b> . Default is <b>FALSE</b>

## Details

**nca.pde.deviation.outlier** calculates individual prediction distribution errors (PDE) and scaled deviation of NCA metrics estimated from observed and simulated data. The deviation of each estimated NCA metrics is scaled by the "spread" of the simulated values. The "spread" is measured either by the 95% parametric prediction interval or 95% non-parametric prediction interval. Any individual yielding an absolute value of the scaled deviation for any of the selected NCA metrics greater than 1, is assigned as an outlier to the corresponding population PK model. The allowed NCA metrics for this diagnostic tests are "AUClast", "AUClower\_upper", "AUCINF\_obs", "AUCINF\_pred", "AUMClast", "Cmax", "Tmax" and "HL\_Lambda\_z". By default, this function uses AUClast and Cmax metrics for the comparison.

## Value

returns the observed data frame with added distance and simulation mean of the nCA metrics, and a data frame with the PDE values of the NCA metrics. If the individual is identified as an outlier for the PK model, histograms of the diagnostic NCA metrics are produced and a graphical object created by arrangeGrob function is returned.

---

nca.read.sim

*Check observed data*

---

## Description

**nca.read.sim** Reads NONMEM simulation output file.

## Usage

```
nca.read.sim(simFile = "nca_simulation.1.npctab.dta", MDV.rm = FALSE)
```

## Arguments

simFile	NONMEM simulation output with the simulated concentration-time data from an internal data frame or an external table. Default is " <b>nca_simulation.1.npctab.dta</b> ".
MDV.rm	If TRUE MDV column is used to remove non-observation rows. Default is FALSE

## Details

**nca.read.sim** Reads NONMEM simulation output file.

## Value

A list of objects

---

ncappc	<i>Performs NCA calculations and population PK model diagnosis.</i>
--------	---

---

## Description

**ncappc** is a flexible tool, to

1. perform a traditional NCA
2. perform simulation-based posterior predictive checks for a population PK model using NCA metrics.

## Usage

```
ncappc(  
  obsFile = "nca_original.npctab.dta",  
  simFile = "nca_simulation.1.npctab.dta.zip",  
  str1Nm = NULL,  
  str1 = NULL,  
  str2Nm = NULL,  
  str2 = NULL,  
  str3Nm = NULL,  
  str3 = NULL,  
  concUnit = NULL,  
  timeUnit = NULL,  
  doseUnit = NULL,  
  obsLog = FALSE,  
  simLog = obsLog,  
  psnOut = TRUE,  
  idNmObs = "ID",  
  timeNmObs = "TIME",  
  concNmObs = "DV",  
  idNmSim = idNmObs,  
  timeNmSim = timeNmObs,  
  concNmSim = concNmObs,  
  onlyNCA = FALSE,  
  AUCTimeRange = NULL,  
  backExtrp = FALSE,  
  LambdaTimeRange = NULL,  
  LambdaExclude = NULL,  
  doseAmtNm = NULL,  
  adminType = "extravascular",  
  doseType = "ns",  
  doseTime = NULL,  
  Tau = NULL,  
  TI = NULL,  
  method = "linearup-logdown",  
  blqNm = NULL,  
)
```

```

blqExcl = 1,
evid = TRUE,
evidIncl = 0,
mdv = FALSE,
filterNm = NULL,
filterExcl = NULL,
negConcExcl = FALSE,
param = c("AUClast", "Cmax"),
timeFormat = "number",
dateColNm = NULL,
dateFormat = NULL,
spread = "npi",
tabCol = c("AUClast", "Cmax", "Tmax", "AUCINF_obs", "Vz_obs", "Cl_obs", "HL_Lambda_z"),
figFormat = "tiff",
noPlot = FALSE,
printOut = TRUE,
studyName = NULL,
new_data_method = TRUE,
overwrite_SIMDATA = NULL,
overwrite_sim_est_file = NULL,
outFileNm = NULL,
out_format = "html",
gg_theme = theme_bw(),
parallel = FALSE,
extrapolate = FALSE,
timing = FALSE,
...
)

```

## Arguments

obsFile	Observed concentration-time data from an internal data frame or an external table with comma, tab or space as separators.
simFile	NONMEM simulation output with the simulated concentration-time data from an internal data frame or an external table. NULL produces just the NCA output, a filename or data frame produces the NCA output as well as the PopPK diagnosis. If new_data_method=TRUE then this can be a compressed file as well.
str1Nm	Column name for 1st level population stratifier. Default is NULL
str1	Stratification ID of the members within 1st level stratification (e.g c(1,2)). Default is NULL
str2Nm	Column name for 2nd level population stratifier. Default is NULL
str2	Stratification ID of the members within 2nd level stratification (e.g c(1,2)). Default is NULL
str3Nm	Column name for 3rd level population stratifier. Default is NULL
str3	Stratification ID of the members within 3rd level stratification (e.g c(1,2)). Default is NULL

concUnit	Unit of concentration (e.g. "ng/mL"). Default is NULL
timeUnit	Unit of time (e.g. "h"). Default is NULL
doseUnit	Unit of dose amount (e.g. "ng"). Default is NULL
obsLog	If TRUE concentration in observed data is in logarithmic scale. Default is FALSE
simLog	If TRUE concentration in simulated data is in logarithmic scale. Default is FALSE
psnOut	If TRUE observed data is an output from PsN or in NONMEM output format. Default is TRUE
idNmObs	Column name for ID in observed data. Default is " <b>ID</b> "
timeNmObs	Column name for time in observed data. Default is " <b>TIME</b> "
concNmObs	Column name for concentration in observed data. Default is " <b>DV</b> "
idNmSim	Column name for ID in simulated data. Default is " <b>ID</b> "
timeNmSim	Column name for time in simulated data. Default is " <b>TIME</b> "
concNmSim	Column name for concentration in simulated data. Default is " <b>DV</b> "
onlyNCA	If TRUE only NCA is performed and ppc part is ignored although simFile is not NULL. Default is FALSE
AUTimeRange	User-defined window of time used to estimate AUC. Default is NULL
backExtrp	If TRUE back-extrapolation is performed while estimating AUC. Default is FALSE
LambdaTimeRange	User-defined window of time to estimate elimination rate-constant. This argument lets the user to choose a specific window of time to be used to estimate the elimination rate constant (Lambda) in the elimination phase. The accepted format for the input to this argument is a numeric array of two elements; c(14, 24) will estimate the Lambda using the data within the time units 14 to 24. Default is NULL
LambdaExclude	User-defined excluded observation time points for estimation of Lambda. This can be numeric value or logical condition (e.g. c(1, 2, "<20", ">=100", "!=100")). Default is NULL
doseAmtNm	Column name to specify dose amount. Default is NULL
adminType	Route of administration. Allowed options are iv-bolus, iv-infusion or extravascular. Default is " <b>extravascular</b> "
doseType	Steady-state (ss) or non-steady-state (ns) dose. Default is " <b>ns</b> "
doseTime	Dose time prior to the first observation for steady-state data. Default is NULL
Tau	Dosing interval for steady-state data. Default is NULL
TI	Infusion duration. If TI is a single numeric value, TI is the same for all individuals. If TI is the name of a column with numeric data present in the data set, TI is set to the unique value of the column for a given individual. Default is NULL
method	Method to estimate AUC. linear method applies the linear trapezoidal rule to estimate the area under the curve. log method applies the logarithmic trapezoidal rule to estimate the area under the curve. linearup-logdown method applies the linear trapezoidal rule to estimate the area under the curve for the ascending part of the curve and the logarithmic trapezoidal rule to estimate the area under the curve for the descending part of the curve. Default is " <b>linearup-logdown</b> "

blqNm	Name of BLQ column if used to exclude data. Default is NULL
blqExcl	Excluded BLQ value; either a numeric value or a logical condition (e.g. 1 or ">=1" or c(1,">3")). Used only if the blqNm is not NULL. Default is <b>"1"</b>
evid	If TRUE EVID is used to filter data. Default is TRUE
evidIncl	Included values in EVID. Default is <b>"0"</b>
mdv	If TRUE MDV is used to include data when MDV=0. Default is FALSE
filterNm	Column name to filter data. Default is NULL
filterExcl	Row exclusion criteria based on the column defined by filterNm. This can be numeric value or logical condition (e.g. c(1, 2, "<20", ">=100", "!=100")). Default is NULL
negConcExcl	If TRUE negative concentrations are excluded. Default is FALSE
param	NCA parameters (AUClast, AUClower_upper, AUCINF_obs, AUCINF_pred, AUMClast, Cmax, Tmax, HL_Lambda_z). Default is <b>(c("AUClast", "Cmax"))</b>
timeFormat	time format (number, H:M, H:M:S). Default is <b>"number"</b>
dateColNm	column name for date if used (e.g. "Date", "DATE"). Default is NULL
dateFormat	date format (D-M-Y, D/M/Y or any other combination of D,M,Y). Default is NULL
spread	Measure of the spread of simulated data ("ppi" (95% parametric prediction interval) or "npi" (95% nonparametric prediction interval)). Default is <b>"npi"</b>
tabCol	Output columns to be printed in the report in addition to ID, dose and population strata information (list of NCA metrics in a string array). Default is <b>c("AUClast", "Cmax", "Tmax", "AUCINF_obs", "Vz_obs", "CI_obs", "HL_Lambda_z")</b>
figFormat	format of the produced figures (bmp, jpeg, tiff, png). Default is <b>"tiff"</b>
noPlot	If TRUE only NCA calculations are performed without any plot generation. Default is FALSE
printOut	If TRUE tabular and graphical outputs are saved on the disk. Default is TRUE
studyName	Name of the study to be added as a description in the report. Default is NULL
new_data_method	If TRUE a faster method of reading data is tested. Default is TRUE
overwrite_SIMDATA	If TRUE new information is created in the SIMDATA directory. If FALSE the information in the SIMDATA directory is used. If NULL a dialog will come up to ask the user what to do. Default is NULL
overwrite_sim_est_file	If TRUE The NCA metrics are created again based on the simulation data. If FALSE the information in the ncaSimEst file is used. If NULL a dialog will come up to ask the user what to do. Default is NULL
outFileNm	Additional tag to the name of the output html and pdf output file hyphenated to the standard ncappc report file name standard ncappc report file name. Default is NULL

out_format	What type of output format should the NCA report have? Pass "all" to render all formats defined within the rmarkdown file. Pass "first" to render the first format defined within the rmarkdown file. Pass "html" to render in HTML. Pass "pdf" to render in PDF.
gg_theme	Which ggplot theme should be used for the plots?
parallel	Should the nca computations for the simulated data be run in parallel? See <a href="#">start_parallel</a> for a description and additional arguments that can be added to this function and passed to <a href="#">start_parallel</a> .
extrapolate	Should the NCA calculations extrapolate from the last observation to infinity?
timing	Should timings of calculations be reported to the screen?
...	Additional arguments passed to other functions, including <a href="#">start_parallel</a> .

## Details

Non-compartmental analysis (NCA) calculates pharmacokinetic (PK) metrics related to the systemic exposure to a drug following administration, e.g. area under the concentration-time curve and peak concentration. **ncappc** performs a traditional NCA using the observed plasma concentration-time data. In the presence of simulated plasma concentration-time data, **ncappc** also performs simulation-based posterior predictive checks (ppc) using NCA metrics for the corresponding population PK (PopPK) model used to generate the simulated data. The diagnostic analysis is performed at the population as well as the individual level. The distribution of the simulated population means of each NCA metric is compared with the corresponding observed population mean. The individual level comparison is performed based on the deviation of the mean of any NCA metric based on simulations for an individual from the corresponding NCA metric obtained from the observed data. Additionally, **ncappc** reports the normalized prediction distribution error (NPDE) of the simulated NCA metrics for each individual and their distribution within a population. **ncappc** produces two default outputs depending on the type of analysis performed, i.e., traditional NCA and PopPK diagnosis. The PopPK diagnosis feature of **ncappc** produces 7 sets of graphical outputs to assess the ability of a population model to simulate the concentration-time profile of a drug and thereby identify model misspecification. In addition, tabular outputs are generated showing the values of the NCA metrics estimated from the observed and the simulated data, along with the deviation, NPDE, regression parameters used to estimate the elimination rate constant and the related population statistics. The default values of the arguments used in **ncappc** are shown in the **Usage** section of this document and/or in **bold** in the **Arguments** section.

## Value

NCA results and diagnostic test results

## Examples

```
out <- ncappc(obsFile=system.file("extdata", "pkdata.csv", package="ncappc"),
  onlyNCA = TRUE,
  extrapolate = TRUE,
  printOut = FALSE,
  evid = FALSE,
  psnOut=FALSE)
```

```
data_1 <- data.frame(
  ID=1,
  TIME = c(0,0.25,0.5,1,1.5,2,3,4,6,8,12,16,24),
  DV=c(0, 0.07, 0.14, 0.21, 0.24, 0.27, 0.26, 0.25, 0.22, 0.19, 0.13, 0.081, 0.033)
)
out_1 <- ncappc(obsFile=data_1,
  onlyNCA = TRUE,
  extrapolate = TRUE,
  printOut = FALSE,
  evid = FALSE,
  timing=TRUE)

data_2 <- dplyr::filter(data_1,TIME>17|TIME<3)
out_2 <- ncappc(obsFile=data_2,
  onlyNCA = TRUE,
  extrapolate = TRUE,
  printOut = FALSE,
  evid = FALSE,
  force_extrapolate=TRUE)
```

---

nca\_ind\_data

*Prepare individual PK data*

---

## Description

Extracts time-conc data for a given individual.

## Usage

```
nca_ind_data(
  pkData,
  dvLog = FALSE,
  dataType = "obs",
  idNm = "ID",
  timeNm = "TIME",
  concNm = "DV",
  adminType = "extravascular",
  TI = NULL,
  dateColNm = NULL,
  dateFormat = NULL,
  timeFormat = "number"
)
```

**Arguments**

pkData	PK concentration-time data.
dvLog	If TRUE concentration is in logarithmic scale. Default is FALSE
dataType	Indicates if the data is observed ("obs") or simulated ("sim"). Since the simulated data is assumed to be obtained from NONMEM output, DATE and clock time (dateColNm, dateFormat, timeFormat) are not used for time data. Default is " <b>obs</b> ".
idNm	Column name for ID in PK data. Default is " <b>ID</b> "
timeNm	Column name for time in PK data. Default is " <b>TIME</b> "
concNm	Column name for concentration in PK data. Default is " <b>DV</b> "
adminType	Route of administration. Allowed options are iv-bolus, iv-infusion or extravascular. Default is " <b>extravascular</b> "
TI	Infusion duration. If TI is a single numeric value, TI is the same for all individuals. If TI is the name of a column with numeric data present in the data set, TI is set to the unique value of the column for a given individual. Default is NULL
dateColNm	column name for date if used (e.g. "Date", "DATE"). Default is NULL
dateFormat	date format (D-M-Y, D/M/Y or any other combination of D,M,Y). Default is NULL
timeFormat	time format (number, H:M, H:M:S). Default is " <b>number</b> "

**Details**

Extracts time-conc data for a given individual.

**Value**

A list of objects with time-conc data and individual infusion duration for iv-infusion data

---

out.digits	<i>output value with correct digits and trailing zero</i>
------------	---

---

**Description**

Function to present a value with correct digits and trailing zero

**Usage**

```
out.digits(x, dig = 3)
```

**Arguments**

x	is the value
dig	is the number of significant digits

**Details**

This is a function to present a value with correct digits and trailing zero. Numbers  $\geq 10000$ , or  $\leq 0.0001$  will be presented in scientific format

**Examples**

```
## Not run:
out.digits(1234)

## End(Not run)
```

---

read_nm_table	<i>Read NONMEM table files produced.</i>
---------------	--

---

**Description**

The function reads in NONMEM table files. The files can be created from the \$EST line or from the \$SIM line in a NONMEM model file.

**Usage**

```
read_nm_table(
  nm_table,
  only_obs = FALSE,
  method = "default",
  quiet = TRUE,
  sim_num = FALSE,
  sim_name = "NSIM",
  verbose = FALSE
)
```

**Arguments**

nm_table	The NONMEM table file to read. A text string. If <code>dplyr</code> and <code>readr</code> are available and <code>method="default"</code> or <code>method="readr_*</code> then <code>nm_table</code> can either a path to a file, a connection, or literal data (either a single string or a raw vector). Files ending in <code>.gz</code> , <code>.bz2</code> , <code>.xz</code> , or <code>.zip</code> will be automatically uncompressed. Files starting with <code>http://</code> , <code>https://</code> , <code>ftp://</code> , or <code>ftps://</code> will be automatically downloaded. Remote <code>gz</code> files can also be automatically downloaded & decompressed.
only_obs	Should the non-observation lines in the data set be removed? Currently filtered using the expected MDV column. TRUE or FALSE.
method	Can be one of <code>default</code> , <code>readr_1</code> , <code>readr_2</code> , <code>readr_3</code> , <code>slow</code> . <code>readr_1</code> should be fastest. All but <code>slow</code> require <code>dplyr</code> and <code>readr</code> (version $\geq 0.2.2$ ).
quiet	Should there be verbose messages about what the function is doing?

sim_num	Should the function add a column to the returned data frame that identifies the simulation number (if present)?
sim_name	The name of the resulting column in the returned data frame if sim_num is true.
verbose	Should the output be more verbose for error checking?

### Details

Currently the function searches the \$TABLE for multiple header lines, and uses that to identify multiple simulations. The function expects at least one header line (NOHEADER option is not allowed in NONMEM table files).

### Value

Returns a data frame of the simulated table with an added column for the simulation number. The data frame is given class `c("tbl_df", "tbl", "data.frame")` for easy use with [dplyr](#).

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