# Package 'NonCompart'

July 15, 2022

Title Noncompartmental Analysis for Pharmacokinetic Data

**Description** Conduct a noncompartmental analysis with industrial strength.

**Version** 0.6.0 **Date** 2022-07-15

1) Use of CD		
2) Automatic		
	or manual slope selection	
	both 'linear-up linear-down' and 'linear-up log-down' method	
	rtial) AUCs with 'linear' or 'log' interpolation method	4
	Gabrielsson J, Weiner D. Pharmacokinetic and Pharmacodynamic Da Applications. 5th ed. 2016. (ISBN:9198299107).	ta Anaiysis -
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•	Seop Bae <k@acr.kr></k@acr.kr>	
Copyright 2016-, 1	Kyun-Seop Bae	
License GPL-3		
NeedsCompilation	no	
LazyLoad yes		
Repository CRAN		
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R topics docu	imented:	
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# Description

It conducts a noncompartmental analysis(NCA) with industrial strength.

## **Details**

The main functions are

```
tblNCA to perform NCA for many subjects.

sNCA to perform NCA for one subject.
```

#### Author(s)

Kyun-Seop Bae <k@acr.kr>

#### References

- 1. Gabrielsson J, Weiner D. Pharmacokinetic and Pharmacodynamic Data Analysis Concepts and Applications. 5th ed. 2016.
- 2. Shargel L, Yu A. Applied Biopharmaceutics and Pharmacokinetics. 7th ed. 2015.
- 3. Rowland M, Tozer TN. Clinical Pharmacokinetics and Pharmacodynamics Concepts and Applications. 4th ed. 2011.
- 4. Gibaldi M, Perrier D. Pharmacokinetics. 2nd ed. revised and expanded. 1982.

AUC 3

AUC

Calculate Area Under the Curve (AUC) and Area Under the first Moment Curve (AUMC) in a table format

## **Description**

Calculate Area Under the Curve(AUC) and the first Moment Curve(AUMC) in two ways; 'linear trapezoidal method' or 'linear-up and log-down' method. Return a table of cumulative values.

## Usage

```
AUC(x, y, down = "Linear")
```

#### **Arguments**

x vector values of independent variable, usually time

y vector values of dependent variable, usually concentration

down either of "Linear" or "Log" to indicate the way to calculate AUC and AUMC

#### **Details**

down="Linear" means linear trapezoidal rule with linear interpolation. down="Log" means linear-up and log-down method.

#### Value

Table with two columns, AUC and AUMC; the first column values are cumulative AUCs and the second column values cumulative AUMCs.

#### Author(s)

Kyun-Seop Bae <k@acr.kr>

#### References

Rowland M, Tozer TN. Clinical Pharmacokinetics and Pharmacodynamics - Concepts and Applications. 4th ed. pp687-689. 2011.

#### See Also

```
LinAUC, LogAUC
```

```
 AUC(Theoph[Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"]) \\ AUC(Theoph[Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"], down="Log")
```

4 BestSlope

BestSlope Choose the best-fit slope for the $log(y)$ and x regression by the criteria of adjusted R-square.	BestSlope	Choose the best-fit slope for the $log(y)$ and $x$ regression by the criteria of adjusted $R$ -square.
---	-----------	--

## **Description**

It sequentially fits  $(\log(y) \sim x)$  from the last point of x to the previous points with at least 3 points. It chooses a slope the highest adjusted R-square. If the difference is less then 1e-4, it pickes longer slope.

## Usage

```
BestSlope(x, y, adm = "Extravascular", TOL=1e-4, excludeDelta = 1)
```

#### **Arguments**

x vector values of x-axis, usually time

y vector values of y-axis, usually concentration

adm one of "Bolus" or "Infusion" or "Extravascular" to indicate drug adminis-

tration mode

TOL tolerance. See Phoneix WinNonlin 6.4 User's Guide p33 for the detail.

excludeDelta Improvement of R2ADJ larger than this value could exclude the last point. De-

fault value 1 is for the compatibility with other software.

## Details

Choosing the best terminal slope (y in log scale) in pharmacokinetic analysis is somewhat challenging, and it could vary by analysis performer. Pheonix WinNonlin chooses a slope with highest adjusted R-squared and the longest one. The difference of adjusted R-Squared less than TOL considered to be 0. This function uses ordinary least square method (OLS). Author recommends to use excludeDelta option with about 0.3.

#### Value

R2	R-squared
1\4	ix-squarcu

R2ADJ adjusted R-squared

LAMZNPT number of points used for slope
LAMZ negative of the slope, lambda\_z
b0 intercept of the regression line
CORRXY correlation of log(y) and x
LAMZLL earliest x for lambda\_z
LAMZUL last x for lambda\_z

CLSTP predicted y value at the last point, predicted concentration for the last time point

#### Author(s)

DetSlope 5

#### See Also

Slope

## **Examples**

DetSlope

Determine slope for the log(y) and x regression manually

## **Description**

You choose a slope for terminal half-life.

#### Usage

```
DetSlope(x, y, SubTitle="", sel.1=0, sel.2=0)
```

## **Arguments**

X	vector values of x-axis, usually time
У	vector values of y-axis, usually concentration
SubTitle	subtitle to be shown on the plot
sel.1	default index of the first element to use
sel.2	default index of the last element to use

#### **Details**

Sometimes BestSlope cannot find terminal slope satisfactorily. Then you can use this function to choose manually. It returns the same format result with BestSlope with an attribute indicating used points.

#### Value

R2	R-squared
R2ADJ	adjusted R-squared
LAMZNPT	number of points used for the slope
LAMZ	negative of the slope, lambda_z
b0	intercept of the regression line
CORRXY	correlation of log(y) and x
LAMZLL	earliest x for lambda_z
LAMZUL	last x for lambda_z
CLSTP	predicted y value at the last point, predicted concentration for the last time point

#### Author(s)

6 gAUC

#### See Also

Slope

#### **Examples**

```
DetSlope(Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"])
DetSlope(Indometh[Indometh$Subject==2, "time"], Indometh[Indometh$Subject==2, "conc"])
```

gAUC

General Area Under the Curve

## **Description**

General AUC function for Emax, TEmax and AUCs

## Usage

```
gAUC(x, y, Ymax = "Emax", XofYmax = "TEmax", AUCname = "AUEClast", iAUC = "",
    Outer = "NEAREST")
```

#### **Arguments**

x usually time

y usually concentration or effect. This can be negative/

Ymax usually Cmax or Emax
XofYmax usually Tmax or TEmax
AUCname usually AUClast or AUEClast

iAUC a data.frame to calculate interval AUCs

Outer indicates how to do the out of x range point

#### **Details**

This is a general purpose AUC function. It calculates only Cmax(Emax), Tmax(TEmax) and AUCs(AUECs). This can be used for effect(pharmacodynamic) data which has negative values. For concentration data, use IntAUC.

## Value

Column names can vary according to the options.

Emax maximum y value

TEmax x value at the maximum y value

AUEClast Area under the y versus x curve

iAUCs Columns from iAUC input

#### Author(s)

gIntAUC 7

#### **Examples**

```
# For one subject
x = Theoph[Theoph$Subject=="1", "Time"]
y = Theoph[Theoph$Subject=="1", "conc"]
gAUC(x, y)

iAUC = data.frame(Name=c("AUC[0-12h]", "AUC[0-24h]"), Start=c(0,0), End=c(12,24))
gAUC(x, y, iAUC=iAUC)
```

gIntAUC

Calculate interval AUC of general form

## **Description**

It calculates interval AUC of general form. This is useful for pharmacodynamic data.

## Usage

```
gIntAUC(x, y, t1, t2, Outer = "NEAREST")
```

## **Arguments**

Х	vector values of independent variable, usually time
у	vector values of dependent variable, usually concentration
t1	start time for AUC
t2	end time for AUC

Outer indicates how to do the out of x range point

## **Details**

This calculates an interval (partial) AUC (from t1 to t2) with the given series of x and y. If t1 and/or t2 cannot be found within x vector, it interpolates. If t1 and/or t2 are out of x range, it uses the nearest value. For concentration data, use IntAUC.

#### Value

```
return interval AUC value (scalar)
```

#### Author(s)

```
Kyun-Seop Bae <k@acr.kr>
```

# See Also

```
gAUC, gInterpol, tblAUC
```

```
gIntAUC (The oph [The oph \$Subject == 1, "Time"], The oph [The oph \$Subject == 1, "conc"], t1 = 0.5, t2 = 11)
```

8 gInterpol

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Interpolate y value for general y value not for concentration

## **Description**

It interpolates y value when a corresponding x value (xnew) does not exist within x vector

## Usage

```
gInterpol(x, y, xnew, Outer="NEAREST")
```

## **Arguments**

x vector values of x-axis, usually time

y vector values of y-axis, usually concentration

xnew new x point to be interpolated, usually new time point

Outer indicates how to do the out of x range point

#### **Details**

This function interpolate y value, if xnew is not in x vector. If xnew is in the x vector, it just returns the given x and y vector. This function usually is called by gIntAUC function Returned vector is sorted in the order of increasing x values.

#### Value

new x and y vector containing xnew and ynew point

#### Author(s)

Kyun-Seop Bae <k@acr.kr>

#### See Also

```
gIntAUC
```

```
x = 1:10 + 0.1

y = -2*x + 40.2

gInterpol(x, y, 1.5)

gInterpol(x, y, 0.5) # Out of range, Left

gInterpol(x, y, 11) # Out of range, Left
```

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IntAUC

Calculate interval AUC

### **Description**

It calculates interval AUC

#### Usage

```
IntAUC(x, y, t1, t2, Res, down = "Linear")
```

#### **Arguments**

X	vector values of independent variable, usually time
У	vector values of dependent variable, usually concentration
t1	start time for AUC
t2	end time for AUC
Res	result from sNCA function
down	either of "Linear" or "Log" to indicate the way to calculate AUC

#### **Details**

This calculates an interval (partial) AUC (from t1 to t2) with the given series of x and y. If t1 and/or t2 cannot be found within x vector, it interpolates according to the down option.

## Value

return interval AUC value (scalar)

# Author(s)

Kyun-Seop Bae <k@acr.kr>

#### References

- 1. Gabrielsson J, Weiner D. Pharmacokinetic and Pharmacodynamic Data Analysis Concepts and Applications. 5th ed. 2016.
- 2. Shargel L, Yu A. Applied Biopharmaceutics and Pharmacokinetics. 7th ed. 2015.
- 3. Rowland M, Tozer TN. Clinical Pharmacokinetics and Pharmacodynamics Concepts and Applications. 4th ed. 2011.
- 4. Gibaldi M, Perrier D. Pharmacokinetics. 2nd ed. revised and expanded. 1982.

## See Also

```
AUC, Interpol
```

10 Interpol

Interpolate y value
• •

## Description

It interpolates y value when a corresponding x value (xnew) does not exist within x vector

# Usage

```
Interpol(x, y, xnew, Slope, b0, down = "Linear")
```

## **Arguments**

X	vector values of x-axis, usually time
у	vector values of y-axis, usually concentration
xnew	new x point to be interpolated, usually new time point
Slope	slope of regression $log(y) \sim x$
b0	y value of just left point of xnew
down	either of "Linear" or "Log" to indicate the way to interpolate

#### **Details**

This function interpolate y value, if xnew is not in x vector. If xnew is in x vector, it just returns the given x and y vector. This function usually is called by IntAUC function Returned vector is sorted in the order of increasing x values.

# Value

new x and y vector containing xnew and ynew point

## Author(s)

Kyun-Seop Bae <k@acr.kr>

# See Also

IntAUC

```
x = 10:1 + 0.1

y = -2*x + 40.2

Interpol(x, y, 1.5)

Interpol(x, y, 1.5, down="Log")
```

LinAUC 11

LinAUC

Area Under the Curve(AUC) and Area Under the first Moment Curve(AUMC) by linear trapezoidal method

## **Description**

It calculates AUC and AUMC using the linear trapezoidal method

## Usage

```
LinAUC(x, y)
```

## **Arguments**

x vector values of the independent variable, usually time

y vector values of the dependent variable, usually concentration

#### **Details**

This function returns AUC and AUMC by the linear trapezoidal method.

#### Value

AUC area under the curve

AUMC area under the first moment curve

#### Author(s)

Kyun-Seop Bae <k@acr.kr>

#### References

- 1. Gabrielsson J, Weiner D. Pharmacokinetic and Pharmacodynamic Data Analysis Concepts and Applications. 5th ed. 2016.
- 2. Shargel L, Yu A. Applied Biopharmaceutics and Pharmacokinetics. 7th ed. 2015.
- 3. Rowland M, Tozer TN. Clinical Pharmacokinetics and Pharmacodynamics Concepts and Applications. 4th ed. 2011.
- 4. Gibaldi M, Perrier D. Pharmacokinetics. 2nd ed. revised and expanded. 1982.

#### See Also

```
LogAUC, AUC
```

```
LinAUC(Theoph[Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"])
AUC(Theoph[Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"]) # compare the last line
```

12 LogAUC

LogAUC

Area Under the Curve(AUC) and Area Under the first Moment Curve(AUMC) by linear-up log-down method

## **Description**

It calculates AUC and AUMC using the linear-up log-down method

## Usage

```
LogAUC(x, y)
```

## **Arguments**

x vector values of the independent variable, usually time

y vector values of the dependent variable, usually concentration

#### **Details**

This function returns AUC and AUMC by the linear-up log-down method.

#### Value

AUC area under the curve

AUMC area under the first moment curve

## Author(s)

Kyun-Seop Bae <k@acr.kr>

#### References

- 1. Gabrielsson J, Weiner D. Pharmacokinetic and Pharmacodynamic Data Analysis Concepts and Applications. 5th ed. 2016.
- 2. Shargel L, Yu A. Applied Biopharmaceutics and Pharmacokinetics. 7th ed. 2015.
- 3. Rowland M, Tozer TN. Clinical Pharmacokinetics and Pharmacodynamics Concepts and Applications. 4th ed. 2011.
- 4. Gibaldi M, Perrier D. Pharmacokinetics. 2nd ed. revised and expanded. 1982.

#### See Also

```
LinAUC,AUC
```

```
LogAUC(Theoph[Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"])
# Compare the last line with the above
AUC(Theoph[Theoph$Subject==1, "Time"], Theoph[Theoph$Subject==1, "conc"], down="Log")
```

Slope 13

Slope Get the Slope of regression $log(y) \sim x$	
---	--

## **Description**

It calculates the slope with linear regression of  $log(y) \sim x$ 

# Usage

```
Slope(x, y)
```

## Arguments

vector values of the independent variable, usually time Χ У vector values of the dependent variable, usually concentration

## **Details**

With time-concentration curve, you frequently need to estimate slope in log(concentration) ~ time. This function is usually called by BestSlope function, and you seldom need to call this function directly.

## Value

R2	R-squared
R2ADJ	adjusted R-squared
LAMZNPT	number of points used for slope
LAMZ	negative of the slope, lambda_z
b0	intercept of the regression line
CORRXY	correlation of log(y) and x
LAMZLL	earliest x for lambda_z
LAMZUL	last x for lambda_z

## Author(s)

Kyun-Seop Bae <k@acr.kr>

### See Also

```
BestSlope
```

```
Slope(Indometh[Indometh$Subject==1, "time"], Indometh[Indometh$Subject==1, "conc"])
```

14 sNCA

sNCA Simplest NCA
-------------------

## Description

This is the work-horse function for NCA.

## Usage

```
sNCA(x, y, dose = 0, adm = "Extravascular", dur = 0, doseUnit = "mg", timeUnit = "h",
    concUnit = "ug/L", iAUC = "", down = "Linear", R2ADJ = 0.7, MW = 0, Keystring="",
    excludeDelta = 1)
```

## **Arguments**

X	usually time
У	usually concentration
dose	given amount, not amount per body weight
adm	one of "Bolus" or "Infusion" or "Extravascular" to indicate drug administration mode $% \left( 1\right) =\left( 1\right) \left( $
dur	duration of infusion
doseUnit	unit of dose
timeUnit	unit of time
concUnit	unit of concentration
iAUC	interval AUCs to calculate
down	either of "Linear" or "Log" to indicate the way to calculate AUC and AUMC
R2ADJ	Minimum adjusted R-square value to determine terminal slope automatically
MW	molecular weight of the drug
Keystring	a text string to be shown at the plot in case of manual selection of terminal slope
excludeDelta	Improvement of R2ADJ larger than this value could exclude the last point. Default value 1 is for the compatibility with other software.

## **Details**

This replaced previous IndiNCA. Author recommends to use excludeDelta option with about 0.3.

## Value

CMAX	maximum concentration, Cmax
CMAXD	dose normalized Cmax, CMAX / Dose, Cmax / Dose
TMAX	time of maximum concentration, Tmax
TLAG	time to observe the first non-zero concentration, for extravascular administration only
CLST	last positive concentration observed, Clast
CLSTP	last positive concentration predicted, Clast_pred

sNCA 15

TLST time of last positive concentration, Tlast half-life by lambda z, ln(2)/LAMZ

LAMZ lambda\_z negative of the best-fit terminal slope

LAMZLL earliest time for LAMZ
LAMZUL last time for LAMZ

LAMZNPT number of points for LAMZ

CORRXY correlation of log(concentration) and time

R2 R-squared

R2ADJ R-squared adjusted

C0 back extrapolated concentration at time 0, for intravascular bolus administration

only

AUCLST AUC from 0 to TLST

AUC using all the given points, including trailing zero concentrations

AUCIFO AUC infinity observed

AUCIFOD AUCIFO / Dose

AUCIFP AUC infinity predicted using CLSTP instead of CLST

AUCIFPD AUCIFP / Dose

AUCPEO AUC % extrapolation observed AUCPEP AUC % extrapolated for AUCIFP

AUCPBEO AUC % back extrapolation observed, for bolus IV administration only

AUCPBEP AUC % back extrapolation predicted with AUCIFP, for bolus IV administration

only

AUMCLST AUMC to the TLST

AUMCIFO AUMC infinity observed using CLST

AUMCIFP AUMC infinity determined by CLSTP

AUMCPEO AUMC % extrapolated observed

AUMCPEP AUMC % extrapolated predicted

MRTIVLST mean residence time (MRT) to TLST, for intravascular administration

MRTIVIFO mean residence time (MRT) infinity using CLST, for intravascular administra-

tion

MRTIVIFP mean residence time (MRT) infinity using CLSTP, for intravascular administra-

tion

MRTEVLST mean residence time (MRT) to TLST, for extravascular administration

MRTEVIFO mean residence time (MRT) infinity using CLST, for extravascular administra-

tion

MRTEVIFP mean residence time (MRT) infinity using CLSTP, for extravascular administra-

tion

VZO volume of distribution determined by LAMZ and AUCIFO, for intravascular

administration

VZP volume of distribution determined by LAMZ and AUCIFP, for intravascular ad-

ministration

VZFO VZO for extravascular administration, VZO/F, F is bioavailability

sNCA

VZFP	VZP for extravascular administration, VZP/F, F is bioavailability
CLO	clearance using AUCIFO, for intravascular administration
CLP	clearance using AUCIFP, for intravascular administration
CLF0	CLO for extravascular administration, CLO/F, F is bioavailability
CLFP	CLP for extravascular administration, CLP/F, F is bioavailability
VSS0	volume of distribution at steady state using CLST, for intravascular administration only
VSSP	volume of distribution at steady state using CLSTP, for intravascular administration only

#### Author(s)

Kyun-Seop Bae <k@acr.kr>

#### References

Gabrielsson J, Weiner D. Pharmacokinetic and Pharmacodynamic Data Analysis - Concepts and Applications. 5th ed. 2016.

#### See Also

help, tblNCA

```
# For one subject
x = Theoph[Theoph$Subject=="1","Time"]
y = Theoph[Theoph$Subject=="1","conc"]
sNCA(x, y, dose=320, doseUnit="mg", concUnit="mg/L", timeUnit="h")
sNCA(x, y, dose=320, concUnit="mg/L")
iAUC = data.frame(Name=c("AUC[0-12h]", "AUC[0-24h]"), Start=c(0,0), End=c(12,24))
sNCA(x, y, dose=320, doseUnit="mg", concUnit="mg/L", timeUnit="h", iAUC=iAUC)
MW = 180.164 # Molecular weight of theophylline
sNCA(x, y/MW, dose=320, doseUnit="mg", concUnit="mmol/L", timeUnit="h")
sNCA(x, y/MW, dose=320, doseUnit="mg", concUnit="mmol/L", timeUnit="h", MW=MW)
sNCA(x, y, dose=320/MW, doseUnit="mmol", concUnit="mg/L", timeUnit="h", MW=MW)
sNCA(x, y/MW, dose=320/MW, doseUnit="mmol", concUnit="mmol/L", timeUnit="h", MW=MW)
sNCA(x, y/MW, dose=320/MW, doseUnit="mmol", concUnit="mmol/L", timeUnit="h", MW=MW)\\
{\tt sNCA(x, y/MW, doseUnit="mmol", concUnit="mmol/L", timeUnit="h", MW=MW)}\\
sNCA(x, y/MW, dose=as.numeric(NA), doseUnit="mmol", concUnit="mmol/L", timeUnit="h",
     MW=MW)
sNCA(x, y, dose=320, concUnit="mg/L", timeUnit="hr")
sNCA(x*60, y, dose=320, concUnit="mg/L", timeUnit="min")
```

tblAUC 17

41.7	4110	
Thi	AUC	

Table output of gAUCs

## **Description**

Do multiple AUCs and returns a result table. See gNCA for more detail i.e. iAUC

## Usage

## **Arguments**

Data	data table name
key	column names of Data to be shown in the output table
colX	column name for x axis
colY	column name for y axis
iAUC	a data.frame to calculate interval AUCs
Ymax	usually Cmax or Emax
XofYmax	usually Tmax or TEmax
AUCname	usually AUClast or AUEClast
Outer	indicates how to do the out of x range point

#### **Details**

Tabular output of AUC with many subjects. This calculates only Cmax(Emax), Tmax(TEmax), AUCs

## Value

Basically same with gAUC

## Author(s)

Kyun-Seop Bae <k@acr.kr>

#### See Also

help, gAUC

```
tblAUC(Theoph, key="Subject", colX="Time", colY="conc")
iAUC = data.frame(Name=c("AUC[0-12h]","AUC[0-24h]"), Start=c(0,0), End=c(12,24))
tblAUC(Indometh, key="Subject", colX="time", colY="conc", iAUC=iAUC)
```

18 tblNCA

tblNCA Taba	le output NCA
-------------	---------------

### **Description**

Do multiple NCA and returns a result table. See sNCA for more detail i.e. iAUC

#### Usage

```
tblNCA(concData, key = "Subject", colTime = "Time", colConc = "conc", dose = 0,
    adm = "Extravascular", dur = 0, doseUnit = "mg", timeUnit = "h",
    concUnit = "ug/L", down = "Linear", R2ADJ = 0, MW = 0, iAUC="",
    excludeDelta = 1)
```

## **Arguments**

concData concentration data table

key column names of concData to be shown in the output table

colTime column name for time

colConc column name for concentration

dose administered dose

adm one of "Bolus" or "Infusion" or "Extravascular" to indicate drug adminis-

tration mode

duration of infusion

doseUnit unit of dose timeUnit unit of time

concUnit unit of concentration

down method to calculate AUC, "Linear" or "Log"

R2ADJ Lowest threshold of adjusted R-square value to do manual slope determination

MW molecular weight of drug iAUC data.frame for interval AUC

excludeDelta Improvement of R2ADJ larger than this value could exclude the last point. De-

fault value 1 is for the compatibility with other software.

#### **Details**

Tabular output of NCA with many subjects. Author recommends to use excludeDelta option with about 0.3.

## Value

Basically same with sNCA

#### Author(s)

Unit 19

#### See Also

```
help, sNCA
```

#### **Examples**

Unit

Display CDISC standard units and multiplied factor of NCA results

# Description

It displays CDISC PP output units and multiplication factor for them.

## Usage

```
Unit(code = "", timeUnit = "h", concUnit = "ng/mL", doseUnit = "mg", MW = 0)
```

# Arguments

code vector of PPTESTCD

timeUnit unit of time

concUnit unit of concentration

doseUnit unit of dose

MW molecular weight of drug

## Value

row names PPTESTCD

Unit unit

Factor internal mulitplication factor

#### Author(s)

Kyun-Seop Bae <k@acr.kr>

```
Unit(concUnit="ug/L", doseUnit="mg")
Unit(concUnit="ng/L", doseUnit="mg")
Unit(concUnit="umol/L", doseUnit="mmol")
Unit(concUnit="nmol/L", doseUnit="mmol")
Unit(concUnit="mmol/L", doseUnit="mg", MW=500)
Unit(concUnit="umol/L", doseUnit="mg", MW=500)
Unit(concUnit="nmol/L", doseUnit="mg", MW=500)
Unit(concUnit="nmol/mL", doseUnit="mg", MW=500)
```

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```
Unit(concUnit="ug/L", doseUnit="mmol", MW=500)
Unit(concUnit="ug/L", doseUnit="mol", MW=500)
Unit(concUnit="ng/L", doseUnit="mmol", MW=500)
Unit(concUnit="ng/mL", doseUnit="mmol", MW=500)
Unit(concUnit="nmol/L", doseUnit="mg")
Unit(concUnit="ug/L", doseUnit="mmol")
```

UnitUrine

Retuns a conversion factor for the amount calculation from urine concentration and volume

## Description

You can get a conversion factor for the multiplication: conc \* vol \* factor = amount in the given unit.

#### Usage

```
UnitUrine(conU = "ng/mL", volU = "mL", amtU = "mg", MW = 0)
```

#### **Arguments**

conU concentration unit
volU volume unit
amtU amount unit
MW molecular weight

## Value

Factor conversion factor for multiplication with the unit in name

# Author(s)

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```
UnitUrine()
UnitUrine("ng/mL", "mL", "mg")
UnitUrine("ug/L", "mL", "mg")
UnitUrine("ug/L", "L", "mg")

UnitUrine("ng/mL", "mL", "g")

UnitUrine("ng/mL", "mL", "mol", MW=500)
UnitUrine("ng/mL", "mL", "mmol", MW=500)
UnitUrine("ng/mL", "mL", "umol", MW=500)
```

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