

Package ‘nplr’

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Type Package

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Depends methods

Imports stats,graphics,utils

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VignetteBuilder knitr

Description Performing drug response analyses and IC50 estimations using n-Parameter logistic regression. Can also be applied to proliferation analyses.

License GPL

URL <https://github.com/mini-pw/nplr>

NeedsCompilation no

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`convertToProp`*Function to Convert a Vector Into Proportions.*

Description

Convert a vector of values to proportions, given a minimum and a maximum value (optional). See Details and Examples.

Usage

```
convertToProp(y, T0 = NULL, Ctrl = NULL)
```

Arguments

`y` : a vector of values (responses to `x`).
`T0` : the minimal value to consider. If `NULL` (default), `min(y, na.rm=TRUE)` will be used. See Details and Warning.
`Ctrl` : the maximal value to consider. If `NULL` (default), `max(y, na.rm=TRUE)` will be used. See Details and Warning.

Details

In typical cell viability experiments, responses to drug concentrations (inhibition rate) may be estimated with respect to a time zero (`T0`) and an untreated condition values (`Ctrl`), as described in [1]:

If none of the `T0` and `Ctrl` values are provided, `min(y, na.rm=TRUE)` and `max(y, na.rm=TRUE)` will be used, respectively. See Warning.

Value

a vector of values.

Warning

Note that, for drug response analyses, rescaling the responses between 0 to 1 using to the min and max of `y`, would lead to estimate a `EC50` (the half effect between the maximum and the minimum of the observed effects), rather than a `IC50`.

Note

The data used as examples come from the NCI-60 Growth Inhibition Data: <https://wiki.nci.nih.gov/display/NCIDTPdata/NCI-60+Growth+Inhibition+Data>, except for `multicell.tsv` which are simulated data.

Author(s)

Frederic Commo, Brian M. Bot

References

1 - <https://dtp.nci.nih.gov/branches/btb/ivclsp.html>

See Also

[nplr](#)

Examples

```
## Using the MDA-N data
op <- par(no.readonly=TRUE)          # save default parameters

require(nplr)
path <- system.file("extdata", "mdan.txt", package = "nplr")
mdan <- read.delim(path)

# fit a model on the original responses (proportions of control):
conc <- mdan$CONC
y0 <- mdan$GIPROP
model0 <- nplr(conc, y0)

# Adjust the data between 0 to 1, then fit a new model:
y1 <- convertToProp(y0)
model1 <- nplr(conc, y1)

par(mfrow=c(1, 2))
plot(model0, ylim = range(0, 1), main = "Original y values")
plot(model1, ylim = range(0, 1), main = "Rescaled y values")
par(op)
```

getEstimates

Function to Estimate x Given y.

Description

This function takes as its first argument a model returned by `nplr()`. By inverting the logistic model, it estimates the x values corresponding to one (or a vector of) y target(s) provided. The standard error of the model, defined as the mean squared error on the fitted values, is used to estimate a confidence interval on the predicted x values, according to the specified `conf.level`. See Details.

Usage

```
## S4 method for signature 'nplr'
getEstimates(object, targets = seq(.9, .1, by = -.1), B = 1e4, conf.level = .95)
```

Arguments

<code>object</code>	an object of class <code>nplr</code> .
<code>targets</code>	one, or a vector of, numerical value(s) for which the corresponding <code>x</code> has to be estimated. Default are target values from .9 to .1.
<code>B</code>	the length of the <code>y</code> distribution from which the <code>x</code> confidence interval is estimated.
<code>conf.level</code>	the estimated <code>x</code> confidence interval, bounded by $(1-\text{conf.level})/2$ and $1 - (1-\text{conf.level})/2$ (by default .95, which gives <code>x.025</code> and <code>x.975</code>).

Details

In `n`-parameter logistic regressions, none of the parameters follow any particular distribution from which confidence intervals can be estimated. To overcome this issue, the standard error is used to generate a normal distribution of the target(s) passed to the function. The quantiles of that distribution are used in order to provide estimated bounds for the corresponding `x` value, with respect to `conf.level`. See also **Warning**.

Value

A data set containing:

`y` the target value.

x.05 the lower bound of the estimated 95% confidence interval (default). If another value is passed to `conf.level`, `x` will be labelled as `x.(1-conf.level)/2`.

`x` the estimated value.

x.95 the upper bound of the estimated 95% confidence interval (default). If another value is passed to `conf.level`, `x` will be labelled as `x.1-(1-conf.level)/2`.

Warning

Notice that, if any $target \leq B$ or $target \geq T$, in other words outside the 2 asymptotes, the maximal (or minimal) possible value the model can estimate is returned.

Note

The data used in the examples are samples from the NCI-60 Growth Inhibition Data: <https://wiki.nci.nih.gov/display/NCIDTPdata/NCI-60+Growth+Inhibition+Data>, except for `multicell.tsv` which are simulated data.

Author(s)

Frederic Commo, Brian M. Bot

See Also

[nplr](#), [plot.nplr](#), [nplrAccessors](#)

Examples

```
# Using the PC-3 data
require(nplr)
path <- system.file("extdata", "pc3.txt", package="nplr")
pc3 <- read.delim(path)
model <- nplr(x = pc3$CONC, y = pc3$GIPROP)
getEstimates(model)
getEstimates(model, c(.3, .6), conf.level = .9)
```

nplr

Function to Fit n-Parameter Logistic Regressions.

Description

This function computes a weighted n-parameters logistic regression, given x (typically compound concentrations) and y values (responses: optic densities, fluorescence, cell counts,...). See Details.

Usage

```
nplr(x, y, useLog = TRUE, LPweight = 0.25, npars = "all",
      method = c("res", "sdw", "gw"), silent = FALSE)
```

Arguments

x	a vector of numeric values, e.g., a vector of drug concentrations.
y	a vector of numeric values, e.g., a vector of responses, typically provided as proportions of control.
useLog	Logical. Should x-values be Log10-transformed? Default is TRUE; set to FALSE if x is already in Log10.
LPweight	a coefficient to adjust the weights. <i>LPweight</i> = 0 will compute a non-weighted np-logistic regression.
npars	a numeric value (or "all") to specify the number of parameters to use in the model. If "all", the logistic model will be tested with 2 to 5 parameters, and the best option will be returned. See Details.
method	a character string to specify which weight method to use. Options are "res" (Default), "sdw", "gw". See Details.
silent	Logical. Specify whether warnings and/or messages should be silenced. Default is FALSE.

Details

The 5-parameter logistic regression is of the form:

$$y = B + (T - B) / [1 + 10^{b * (x_{mid} - x)}]^s$$

where B and T are the bottom and top asymptotes, respectively, b and xmid are the Hill slope and the x-coordinate at the inflection point, respectively, and s is an asymmetric coefficient. This equation is sometimes referred to as the Richards' equation [1,2].

When specifying npars = 4, the s parameter is forced to be 1, and the corresponding model is a 4-parameter logistic regression, symmetrical around its inflection point. When specifying npars = 3 or npars = 2, two more constraints are added, forcing B and T to be 0 and 1, respectively.

Weight methods:

The model parameters are optimized, simultaneously, using **nlm**, given a sum of squared errors function, $sse(Y)$, to minimize:

$$sse(Y) = \sum [W.(Y_{obs} - Y_{fit})^2]$$

where Yobs, Yfit, and W are the vectors of observed values, fitted values, and weights, respectively.

In order to reduce the effect of possible outliers, the weights can be computed in different ways, specified in nplr:

residual weights, "res":

$$W = (1/residuals)^{LPweight}$$

where residuals and LPweight are the squared error between the observed and fitted values, and a tuning parameter, respectively. Best results are generally obtained by setting $LPweight = 0.25$ (default value), while setting $LPweight = 0$ results in computing a non-weighted sum of squared errors.

standard weights, "sdw":

$$W = 1/Var(Y_{obs_r})$$

where Var(Yobs_r) is the vector of the within-replicates variances.

general weights, "gw":

$$W = 1/Y_{fit}^{LPweight}$$

where Yfit are the fitted values. As for the residuals-weights method, setting $LPweight = 0$ results in computing a non-weighted sum of squared errors.

The standard weights and general weights methods are described in [3].

Value

An object of class nplr.

slots

x the x values as they are used in the model. It can be $\text{Log}_{10}(x)$ if useLog was set to TRUE.

y the y values.

useLog logical.

npars the best number of parameters if npars="all", or the specified number of parameters otherwise.

LPweight the weights tuning parameter.

yFit the y fitted values.

xCurve the x values generated to draw the curve. 200 points between the min and max of x.

yCurve the fitted values used to draw the curve. These correspond to xCurve.

inflPoint the inflection point x and y coordinates.

goodness the goodness-of-fit. The correlation between the fitted and the observed y values.

stdErr the mean squared error between the fitted and the observed y values.

pars the model parameters.

AUC the area under the curve estimated using both the trapezoid method and Simpson's rule.

Note

The data used in the examples are samples from the NCI-60 Growth Inhibition Data: <https://wiki.nci.nih.gov/display/NCIDTPdata/NCI-60+Growth+Inhibition+Data>, except for multicell.tsv which are simulated data.

Author(s)

Frederic Commo, Brian M. Bot

References

- 1- Richards, F. J. (1959). A flexible growth function for empirical use. J Exp Bot 10, 290-300.
- 2- Giraldo J, Vivas NM, Vila E, Badia A. Assessing the (a)symmetry of concentration-effect curves: empirical versus mechanistic models. Pharmacol Ther. 2002 Jul;95(1):21-45.
- 3- Motulsky HJ, Brown RE. Detecting outliers when fitting data with nonlinear regression - a new method based on robust nonlinear regression and the false discovery rate. BMC Bioinformatics. 2006 Mar 9;7:123.

See Also

[convertToProp](#), [getEstimates](#), [plot.nplr](#), [nplrAccessors](#)

Examples

```
# Using the PC-3 data
require(nplr)
path <- system.file("extdata", "pc3.txt", package = "nplr")
pc3 <- read.delim(path)
model <- nlplr(x = pc3$CONC, y = pc3$GIPROP)
plot(model)
```

nplrAccessors	nplr accessor functions
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Description

Methods for extracting information from an object of class `nplr`. Each of the below methods are simply convenience functions which extract the corresponding slots (as the name of each method suggests) from the object of class `nplr`.

Methods

```
signature(object = "nplr") • getX(object)
  • getY(object)
  • getXcurve(object)
  • getYcurve(object)
  • getFitValues(object)
  • getInflexion(object)
  • getPar(object)
  • getAUC(object)
  • getGoodness(object)
  • getStdErr(object)
  • getWeights(object)
```

See Also

[nplr](#), [getEstimates](#)

overlay	<i>Plotting Multiple nplr Objects</i>
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Description

To superimpose multiple logistic models fitted using `nplr`.

Usage

```
overlay(modellist = NULL, showLegend = TRUE, cols = NULL, ...)
```

Arguments

<code>modellist</code>	: list. A list of objects of class <code>nplr</code> .
<code>showLegend</code>	: logical. Whether the legend has to be displayed.
<code>cols</code>	: character. A vector of colors to use. If NULL (default), greys will be used.
<code>...</code>	: Other graphical parameters. See par .

Details

None

Source

None

References

None

See Also[plot.nplr](#)**Examples**

```

path <- system.file("extdata", "multicell.tsv", package="nplr")
multicell <- read.delim(path)

# Computing models (to store in a list)
cellsList <- split(multicell, multicell$cell)
Models <- lapply(cellsList, function(tmp){
  nplr(tmp$conc, tmp$resp, silent = TRUE)
})

# Visualizing
overlay(Models, xlab = expression(Log[10](Conc.)), ylab = "Resp.",
  main="Superimposing multiple curves", cex.main=1.5)

```

plot.nplr

*Plotting nplr Objects***Description**

This function allows visualizing logistic models fitted using [nplr](#).

Usage

```

## S3 method for class 'nplr'
plot(x, pcol = "aquamarine1", lcol = "red3",
  showEstim = FALSE, showCI = TRUE, showGOF = TRUE, showInfl = FALSE,
  showPoints = TRUE, showSDerr = FALSE, B = 1e4, conf.level = .95, unit = "", ...)

```

Arguments

x	: an object of class <code>nplr</code>
pcol	: the points color.
lcol	: the line color.
showEstim	: logical/numeric. If a numerical value is passed (a y value to reach), the estimated x value , and interval, is displayed on the plot. Default is FALSE
showCI	: logical. show the estimated confidence interval
showGOF	: logical. show the estimated goodness-of-fit.
showInfl	: logical. add the inflexion point on the plot.
showPoints	: logical. add the points on the plot.
showSDerr	: logical. add the standard errors on the plot (maybe useful in case of experiment with replicates).
B	: the length of simulated y values. Used to estimate the confidence interval
conf.level	: the confidence level. See <code>getEstimates</code>
unit	: the unit to specify when showEstim is TRUE. Default is an empty string.
...	: other graphical parameters. See <code>par</code> .

Details

None

Note

The data used in the examples are samples from the NCI-60 Growth Inhibition Data:
<https://wiki.nci.nih.gov/display/NCIDTPdata/NCI-60+Growth+Inhibition+Data>, except for multicell.tsv which are simulated data.

Source

None

References

None

See Also

`overlay`

Examples

```
# Using the PC-3 data
require(nplr)
path <- system.file("extdata", "pc3.txt", package = "nplr")
pc3 <- read.delim(path)
model <- nplr(x = pc3$CONC, y = pc3$GIPROP)
plot(model, showEstim = 0.5, unit = "nM")
```

summary.nplr	<i>summarizing nplr Objects</i>
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Description

A S3 method to visualize a model summary as a table.

Usage

```
## S3 method for class 'nplr'  
summary(object, ...)
```

Arguments

object : an object of class `nplr`
... : other optional parameters (not used).

Details

None

Note

The data used in the examples are samples from the NCI-60 Growth Inhibition Data:
<https://wiki.nci.nih.gov/display/NCIDTPdata/NCI-60+Growth+Inhibition+Data>, except
for multicell.tsv which are simulated data.

Source

None

References

None

See Also

`plot.nplr`

Examples

```
# Using the PC-3 data  
require(nplr)  
path <- system.file("extdata", "pc3.txt", package = "nplr")  
pc3 <- read.delim(path)  
model <- nplr(x = pc3$CONC, y = pc3$GIPROP)  
summary(model)
```

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