Package 'tcplfit2'

August 17, 2023

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Title Concentration-Response Modeling of HTS or Transcriptomics Data
Version 0.1.5
Description Performs the basic concentration response curve fitting used in the 'tcpl' package. It is a substitute for the original tcplFit() function (and sub-functions) and allows a wider variety of concentration-response models. All of the models included in the 'BMDExpress' package are now part of this package, and the output includes a calculation of the bmd (Benchmark Dose or concentration) value.
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R topics documented:
acgnlsobj acy bmdbounds bmdobj cnst concRespCore concRespPlot exp2 exp3 exp4
• <u>• </u>

2 acgnlsobj

```
15
16
22
 23
loggnls.....
 25
poly1
poly2
28
34
 36
```

acgnlsobj

AC GNLS Objective Function

Description

GNLS objective function set to y for gnls solver.

Usage

Index

```
acgnlsobj(x, y, tp, ga, p, la, q)
```

Arguments

X	Concentration.
у	Desired activity level.
tp	Top.
ga	Gain AC50.
р	Gain power.
la	Loss AC50.
q	Loss power.

acy 3

Value

Difference between GNLS model repsone at x and y.

acy Activity Concentration y

Description

Returns concentration at which model equals y.

Usage

```
acy(
   y,
   modpars,
   type = "hill",
   returntop = FALSE,
   returntoploc = FALSE,
   getloss = FALSE,
   verbose = FALSE
)
```

Arguments

У	Activity value at which the concentration is desired. y should be less than the model's top, if there is one, and greater than zero.
modpars	List of named model parameters. Model parameters can include: "a", "b", "ga", "la", "p", "q", "tp". ga and la should NOT be in log units.
type	Model type; must be one of: "exp1", "exp2", "exp3", "exp4", "gnls", "hill", "poly1", "poly2", "pow".
returntop	When TRUE, returns actual top value for gnls. Has no effect for other models.
returntoploc	When TRUE, returns concentration of top for gnls. Has no effect for other models. If top location can't be found, NA is returned.
getloss	When TRUE, returns value on loss side of curve for gnls. Has no effect for other models.
verbose	When TRUE, shows warnings.

Details

Mathematically inverts model functions of the given type, except for gnls, which is numerically inverted. gnls returns NA when y > tp. Other options return the actual top (as opposed to theoretical tp) and top location for gnls model. gnls model defaults to giving concentration on gain side. Only one of getloss, returntop, and returntoploc should be TRUE at a time. If top location solution fails for gnls, top is set to tp. Returns NA if gnls numerical solver fails. Returns NA if model was not successfully fit.

Value

Ouputs concentration at activity y, or gnls top or top concentration, when applicable.

4 bmdbounds

Examples

```
acy(1, list(ga = 10, tp = 2, p = 3), type = "hill")
acy(1, list(ga = .1, tp = 2, p = 3, q = 3,la = 10), type = "gnls")
acy(1, list(ga = .1, tp = 2, p = 3, q = 3,la = 10), type = "gnls", getloss = TRUE)
acy(1, list(ga = .1, tp = 2, p = 3, q = 3,la = 10), type = "gnls", returntop = TRUE)
acy(1, list(ga = .1, tp = 2, p = 3, q = 3,la = 10), type = "gnls", returntoploc = TRUE)
```

bmdbounds

BMD Bounds

Description

Uses maximum likelihood method to tune the upper and lower bounds on the BMD (BMDU, BMDL)

Usage

```
bmdbounds(
   fit_method,
   bmr,
   pars,
   conc,
   resp,
   onesidedp = 0.05,
   bmd = NULL,
   which.bound = "lower"
)
```

Arguments

fit_method	Fit method: "exp2", "exp3", "exp4", "exp5", "hill", "gnls", "poly1", "poly2", or "pow".
bmr	Benchmark response.
pars	Named vector of model parameters: a,b,tp,ga,p,la,q,er output by httrfit, and in that order.
conc	Vector of concentrations (NOT in log units).
resp	Vector of responses corresponding to given concentrations.
onesidedp	The one-sided p-value. Default of .05 corresponds to 5 percentile BMDL, 95 percentile BMDU, and 90 percent CI.
bmd	Can optionally input the bmd when already known to avoid unnecessary calculation.

Details

which.bound

"lower".

Takes in concentration response fit details and outputs a bmdu or bmdl, as desired. If bmd is not finite, returns NA. If the objective function doesn't change sign or the root finding otherwise fails, it returns NA. These failures are not uncommon since some curves just don't reach the desired confidence level.

Returns BMDU if which.bound = "upper"; returns BMDL if which.bound =

bmdobj 5

Value

Returns either the BMDU or BMDL.

Examples

bmdobj

BMD Objective Function

Description

Utility function for bmdbounds

Usage

```
bmdobj(bmd, fname, bmr, conc, resp, ps, mll, onesp, partype = 2)
```

Arguments

bmd	Benchmark dose.
fname	Function name: "exp2", "exp3", "exp4", "exp5", "hillfn", "gnls", "poly1", "poly2", or "pow".
bmr	Benchmark response.
conc	Vector of concentrations NOT in log units.
resp	Vector of corresponding responses.
ps	Named list of parameters.
mll	Maximum log-likelihood of winning model.
onesp	One-sided p-value.
partype	Number for parameter type. Type 1 is y-scaling: a or tp. Type 2 is x-scaling: b or ga, when available, a otherwise. Type 3 is power scaling: p when available,

then b or ga, then a if no others. Since bmd is linked to the x-scale, type 2 should always be used. Other types can also be vulnerable to underflow/overflow.

Value

Objective function value to find the zero of.

6 concRespCore

cnst

Constant Model

Description

```
f(x) = 0
```

Usage

```
cnst(ps, x)
```

Arguments

ps Vector of parameters (ignored)

x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

```
cnst(1,1)
```

concRespCore

Concentration Response Core

Description

Core of concentration response curve fitting for pvalue based cutoff. This function calls tcplfit2_core to get curve fits, and then tcplhit2_core to perform the hitcalling.

Usage

```
concRespCore(
  row,
  fitmodels = c("cnst", "hill", "gnls", "poly1", "poly2", "pow", "exp2", "exp3", "exp4",
        "exp5"),
  conthits = TRUE,
  aicc = FALSE,
  force.fit = FALSE,
  bidirectional = TRUE,
  verbose = FALSE,
  do.plot = FALSE,
  return.details = FALSE,
  bmr_scale = 1.349,
  bmd_low_bnd = NULL,
  bmd_up_bnd = NULL
)
```

concRespCore 7

Arguments

row A named list that must include:

• conc - list of concentrations (not in log units)

• resp - list of corresponding responses

• bmed - median of noise estimate.

· cutoff - noise cutoff

• onesd - 1 standard deviation of the noise (for bmd calculation)

Other elements (usually identifiers, like casrn) of row will be attached to the final output.

fitmodels Vector of model names to use.

conthits = TRUE uses continuous hitcalls, otherwise they're discrete.

aicc aicc = TRUE uses corrected AIC to choose winning method; otherwise regular

AIC.

force.fit If TRUE force the fitting to proceed even if there are no points outside of the

bounds (default FALSE)

bidirectional If TRUE allow fitting to happen in both directions (default TRUE)

verbose If TRUE, write extra output from tcplfit2_core (default FALSE)

do.plot If TRUE, create a plot in the tcplfit2_core function (default FALSE)

return.details If TRUE, return the hitcalling details and the summary, if FALSE (default), just

return the summary

bmr_scale - bmr scaling factor (for bmd calculation) default = 1.349

bmd_low_bnd Multiplier for bmd lower bound. A value of .1 would require the bmd to be no

lower than 1/10th of the lowest concentration tested.

bmd_up_bnd Multiplier for the bmd upper bound. A value of 10 would require the bmd to be

no lower than 10 times the highest concentration tested.

Value

A list of two elements. The first (summary) is the output from tcplhit2_core. The second, params is the output from tcplfit2_core a dataframe of one row containing

8 concRespPlot

concRespPlot

Concentration Response Plot

Description

Plots a concentration response curve for one sample/endpoint combination. This is a generic function and it is expected that users will make their own versions

Usage

```
concRespPlot(row, ymin = -120, ymax = 120, draw.error.arrows = FALSE)
```

Arguments

row

Named list containing:

- conc conc string separated by I's
- resp response string separated by I's
- method scoring method determines plot bounds
- name chemical name for plot title
- cutoff noise cutoff
- bmr baseline median response; level at which bmd is calculated
- er fitted error term for plotting error bars
- a, tp, b, ga, p, la, q other model parameters for fit curve
- fit_method curve fit method
- bmd, bmdl, bmdu bmd, bmd lower bound, and bmd upper bound
- ac50, acc curve value at 50% of top, curve value at cutoff
- top curve top
- name name of the chemical
- · assay name of the assay, signature, or other endpoint
- other identifiers

Other elements are ignored.

ymin

Minimum value of response for the plot

ymax

Maximum value of response for the plot

draw.error.arrows

If TRUE, draw lines representing the uncertainty in the response estimate, instead of the actual response points

Details

row is one row of data from concRespCore

Value

No output.

exp2 9

Examples

exp2

Exponential 2 Model

Description

$$f(x) = a * (e^{(x/b)} - 1)$$

Usage

Arguments

ps Vector of parameters: a,b,er

x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

```
exp2(c(1,2),1)
```

exp3

Exponential 3 Model

Description

$$f(x) = a * (e^{(x/b)^p} - 1)$$

Usage

```
exp3(ps, x)
```

10 exp4

Arguments

ps Vector of parameters: a,b,p,er

x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

```
exp3(c(1,2,2),1)
```

exp4

Exponential 4 Model

Description

$$f(x) = tp * (1 - 2^{(-x/ga)})$$

Usage

Arguments

ps Vector of parameters: tp,ga,er

x Vector of concentrations (regular units)

Value

Vector of model responses

```
exp4(c(1,2),1)
```

exp5

exp5

Exponential 5 Model

Description

$$f(x) = tp * (1 - 2^{(-(x/ga)^p)})$$

Usage

```
exp5(ps, x)
```

Arguments

ps Vector of parameters: tp,ga,p,er
x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

```
exp5(c(1,2,3),1)
```

fitcnst

Constant Model Fit

Description

Function that fits a constant line f(x) = 0 and returns generic model outputs.

Usage

```
fitcnst(conc, resp, nofit = FALSE, ...)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

nofit If nofit = TRUE, returns formatted output filled with missing values.

... Space for parameters so fitchest can be called similar to other fitting functions

(currently unused)

Details

success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. aic, rme, and er are set to NA in case of nofit or failure. pars always equals "er".

12 fitexp2

Value

List of five elements: success, aic (Akaike Information Criteria), rme (root mean square error), er (error parameter), pars (parameter names).

Examples

```
fitcnst(c(.1,1,10,100), c(1,2,0,-1))
fitcnst(c(.1,1,10,100), c(1,2,0,-1), nofit = TRUE)
```

fitexp2

Exponential 2 Model Fit

Description

Function that fits to $f(x) = a * (e^{(x/b)} - 1)$ and returns generic model outputs.

Usage

```
fitexp2(conc, resp, bidirectional = TRUE, verbose = FALSE, nofit = FALSE)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

Details

Zero background and increasing absolute response are assumed. Parameters are "a" (y scale), "b" (x scale), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

```
fitexp2(c(.1,1,10,100), c(0,.1,1,10))
```

fitexp3 13

fitexp3

Exponential 3 Model Fit

Description

Function that fits to $f(x) = a * (e^{(x/b)^p} - 1)$ and returns generic model outputs.

Usage

```
fitexp3(
  conc,
  resp,
  bidirectional = TRUE,
  verbose = FALSE,
  nofit = FALSE,
  dmin = 0.3
)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

dmin Minimum allowed value of p.

Details

Zero background and increasing absolute response are assumed. Parameters are "a" (y scale), "b" (x scale), "p" (power), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

```
fitexp3(c(.03,.1,.3,1,3,10,30,100), c(0,0,.1, .2, .4, 1, 4, 50))
```

14 fitexp5

f	i	t	۹	x	n	4

Exponential 4 Model Fit

Description

Function that fits to $f(x) = tp * (1 - 2^{(-x/ga)})$ and returns generic model outputs.

Usage

```
fitexp4(conc, resp, bidirectional = TRUE, verbose = FALSE, nofit = FALSE)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

Details

Zero background and increasing absolute response are assumed. Parameters are "tp" (top), "ga" (AC50), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

Examples

```
fitexp4(c(.03,.1,.3,1,3,10,30,100), c(0,0,.1, .2, .5, 1, 1.5, 2))
```

fitexp5

Exponential 5 Model Fit

Description

Function that fits to $f(x) = tp * (1 - 2^{(-(x/ga)^p)})$ and returns generic model outputs.

fitgnls 15

Usage

```
fitexp5(
  conc,
  resp,
  bidirectional = TRUE,
  verbose = FALSE,
  nofit = FALSE,
  dmin = 0.3
)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

dmin Minimum allowed value of p.

Details

Zero background and increasing absolute response are assumed. Parameters are "tp" (top), "ga" (AC50), "p" (power), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

Examples

```
fitexp5(c(.03,.1,.3,1,3,10,30,100), c(0,0,.1, .2, .5, 1, 1.5, 2))
```

fitgnls

Gain-Loss Model Fit

Description

```
Function that fits to f(x) = \frac{tp}{[(1+(ga/x)^p)(1+(x/la)^q)]} and returns generic model outputs.
```

16 fithill

Usage

```
fitgnls(
  conc,
  resp,
  bidirectional = TRUE,
  verbose = FALSE,
  nofit = FALSE,
  minwidth = 1.5
)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

minwidth Minimum allowed distance between gain ac50 and loss ac50 (in log10 units).

Details

Concentrations are converted internally to $\log 10$ units and optimized with $f(x) = \frac{tp}{[(1+10^{(p*(ga-x))})(1+10^{(q*(x-la))})]}$, then ga, la, ga_sd, and la_sd are converted back to regular units before returning. Zero background and increasing initial absolute response are assumed. Parameters are "tp" (top), "ga" (gain AC50), "p" (gain power), "la" (loss AC50), "q" (loss power) and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

Examples

```
fitgnls(c(.03,.1,.3,1,3,10,30,100), c(0,.3,1, 2, 2.1, 1.5, .8, .2))
```

fithill

Hill Model Fit

Description

```
Function that fits to f(x) = \frac{tp}{[(1+(ga/x)^p)]} and returns generic model outputs.
```

Usage

```
fithill(conc, resp, bidirectional = TRUE, verbose = FALSE, nofit = FALSE)
```

fitpoly1

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

Details

Concentrations are converted internally to $\log 10$ units and optimized with $f(x) = \frac{tp}{(1+10(p*(ga-x)))}$, then ga and ga_sd are converted back to regular units before returning. Zero background and increasing initial absolute response are assumed. Parameters are "tp" (top), "ga" (gain AC50), "p" (gain power), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

Examples

```
fithill(c(.03,.1,.3,1,3,10,30,100), c(0,0,.1,.2,.5,1,1.5,2))
```

fitpoly1

Polynomial 1 (Linear) Model Fit

Description

Function that fits to f(x) = a * x and returns generic model outputs.

Usage

```
fitpoly1(conc, resp, bidirectional = TRUE, verbose = FALSE, nofit = FALSE)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

18 fitpoly2

Details

Zero background and increasing absolute response are assumed. Parameters are "a" (y scale) and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

Examples

```
fitpoly1(c(.03,.1,.3,1,3,10,30,100), c(0,.01,.1,.1,.2,.5,2,5))
```

fitpoly2

Polynomial 2 (Quadratic) Model Fit

Description

Function that fits to $f(x) = a * (\frac{x}{b} + \frac{x^2}{b^2})$ and returns generic model outputs.

Usage

```
fitpoly2(conc, resp, bidirectional = TRUE, verbose = FALSE, nofit = FALSE)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

Details

Zero background and monotonically increasing absolute response are assumed. Parameters are "a" (y scale), "b" (x scale), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

fitpow 19

Examples

```
fitpoly2(c(.03,.1,.3,1,3,10,30,100), c(0,.01,.1, .1, .2, .5, 2, 8))
```

fitpow

Power Model Fit

Description

Function that fits to $f(x) = a * x^p$ and returns generic model outputs.

Usage

```
fitpow(
  conc,
  resp,
  bidirectional = TRUE,
  verbose = FALSE,
  nofit = FALSE,
  nmin = 0.3
)
```

Arguments

conc Vector of concentration values NOT in log units.

resp Vector of corresponding responses.

bidirectional If TRUE, model can be positive or negative; if FALSE, it will be positive only.

verbose If TRUE, gives optimization and hessian inversion details.

nofit If nofit = TRUE, returns formatted output filled with missing values.

nmin Minimum allowed value of p.

Details

Zero background and monotonically increasing absolute response are assumed. Parameters are "a" (y scale), "p" (power), and error term "er". success = 1 for a successful fit, 0 if optimization failed, and NA if nofit = TRUE. cov = 1 for a successful hessian inversion, 0 if it fails, and NA if nofit = TRUE. aic, rme, modl, parameters, and parameter sds are set to NA in case of nofit or failure.

Value

Named list containing: success, aic (Akaike Information Criteria), cov (success of covariance calculation), rme (root mean square error), modl (vector of model values at given concentrations), parameters values, parameter sd (standard deviation) estimates, pars (vector of parameter names), sds (vector of parameter sd names).

```
fitpow(c(.03,.1,.3,1,3,10,30,100), c(0,.01,.1, .1, .2, .5, 2, 8))
```

20 gnlsderivobj

gnls

Gain-Loss Model

Description

$$f(x) = \frac{tp}{[(1+(ga/x)^p)(1+(x/la)^q)]}$$

Usage

```
gnls(ps, x)
```

Arguments

ps Vector of parameters: tp,ga,p,la,q,er x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

```
gnls(c(1,2,1,2,2),1)
```

gnlsderivobj

GNLS Derivative Objective Function

Description

Derivative of the gnls function set to zero for top location solver.

Usage

```
gnlsderivobj(x, tp, ga, p, la, q)
```

Arguments

X	Concentration
tp	Top.
ga	Gain AC50.
p	Gain power.
la	Loss AC50.
q	Loss power.

Value

Value of gnls derivative at x.

hillfn 21

hillfn

Hill Model

Description

$$f(x) = \frac{tp}{[(1 + (ga/x)^p)]}$$

Usage

```
hillfn(ps, x)
```

Arguments

ps Vector of parameters: tp,ga,p,er

x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

```
hillfn(c(1,2,3),1)
```

hitcont

Continuous Hitcalls

Description

Wrapper that computes continuous hitcalls for a provided concRespCore input row.

Usage

```
hitcont(indf, xs = NULL, ys = NULL, newcutoff)
```

Arguments

indf	Dataframe similar to concRespCore output. Must contain "conc" and "resp"
	columns if xs and ys are not provided. Must contain "top", "ac50", "er", "fit_method",

"caikwt", and "mll" columns as well as columns for each model parameter.

xs List of concentration vectors that can be provided for speed.
ys List of response vectors that can be provided for speed.

newcutoff Vector of new cutoff values to use. Length should be equal to rows in indf.

Details

indf parameter columns should be NA when not required by fit method. "conc" and "resp" entries should be a single string with values separated by I. Details on indf columns can be found in concRespCore.

22 hitcontinner

Value

Vector of hitcalls between 0 and 1 with length equal to indf row number.

Examples

```
conc <- list(.03, .1, .3, 1, 3, 10, 30, 100)
resp <- list(0, .2, .1, .4, .7, .9, .6, 1.2)
row <- list(
   conc = conc,
   resp = resp,
   bmed = 0,
   cutoff = 1,
   onesd = .5,
   name = "some chemical",
   assay = "some assay"
)
res <- concRespCore(row, conthits = TRUE)
hitcont(res, newcutoff = 0.2)</pre>
```

hitcontinner

Continuous Hitcalls Inner

Description

Calculates continuous hitcall using 3 statistical metrics.

Usage

```
hitcontinner(conc, resp, top, cutoff, er, ps, fit_method, caikwt, mll)
```

Arguments

```
Vector of concentrations.
conc
resp
                   Vector of responses.
                  Model top.
top
cutoff
                  Desired cutoff.
                  Model error parameter.
er
                  Vector of used model parameters in order: a, tp, b, ga, p, la, q, er.
ps
                  Name of winning fit method (should never be constant).
fit\_method
caikwt
                  Akaike weight of constant model relative to winning model.
                  Maximum log-likelihood of winning model.
mll
```

Details

This function is called either directly from concRespCore or via hitcont. Details of how to compute function input are in concRespCore.

Value

Continuous hitcall between 0 and 1.

hitlogic 23

Examples

```
conc = c(.03,.1,.3,1,3,10,30,100)
resp = c(0,.1,0,.2,.6,.9,1.1,1)
top = 1.023239
er = -3.295307
ps = c(1.033239, 2.453014, 1.592714, er = -3.295307) #tp,ga,p,er
fit_method = "hill"
caikwt = 1.446966e-08
mll = 12.71495
hitcontinner(conc,resp,top,cutoff = 0.8, er,ps,fit_method, caikwt, mll)
hitcontinner(conc,resp,top,cutoff = 1, er,ps,fit_method, caikwt, mll)
hitcontinner(conc,resp,top,cutoff = 1.2, er,ps,fit_method, caikwt, mll)
```

hitlogic

Hit Logic (Discrete)

Description

Wrapper that computes discrete hitcalls for a provided concRespCore dataframe.

Usage

```
hitlogic(indf, newbmad = NULL, xs = NULL, ys = NULL, newcutoff = NULL)
```

Arguments

indf	Dataframe similar to concRespCore input Must contain "conc" and "resp" columns if xs and ys are not provided. Must contain "cutoff" and "bmad_factor" columns if newbmad is not NULL. Must contain "top" and "ac50" columns. "conc" and "resp" entries should be a single string with values separated by l.
newbmad	(Deprecated) New number of bmads to use for the cutoff.
xs	List of concentration vectors that can be provided for speed.
ys	List of response vectors that can be provided for speed.
newcutoff	Vector of new cutoff values to use. Length should be equal to rows in indf.

Value

Vector of hitcalls with length equal to number of rows in indf.

```
conc = rep(".03|.1|.3|1|3|10|30|100",2)
resp = rep("0|0|.1|.1|.5|.5|1|1",2)
indf = data.frame(top = c(1,1), ac50 = c(3,4), conc = conc, resp = resp,
    stringsAsFactors = FALSE)
hitlogic(indf, newcutoff = c(.8,1.2))
```

24 loggnls

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Hit Logic Inner (Discrete)

Description

Contains hit logic, called directly during CR fitting or later through "hitlogic".

Usage

```
hitloginner(conc = NULL, resp, top, cutoff, ac50 = NULL)
```

Arguments

conc	Vector of concentrations (No longer necessary).
resp	Vector of responses.
top	Model top.
cutoff	Desired cutoff.
ac50	Model AC50 (No longer necessary).

Details

The purpose of this function is to keep the actual hit rules in one location so it can be called during CR fitting, and then again after the fact for a variety of cutoffs. Curves fit with constant winning should have top = NA, generating a miss.

Value

Outputs 1 for hit, 0 for miss.

Examples

```
hitloginner(resp = 1:8, top = 7, cutoff = 5) #hit
hitloginner(resp = 1:8, top = 7, cutoff = 7.5) #miss: top too low
hitloginner(resp = 1:8, top = 9, cutoff = 8.5) #miss: no response> cutoff
hitloginner(resp = 1:8, top = NA, cutoff = 5) #miss: no top (constant)
```

loggnls

Log Gain-Loss Model

Description

$$f(x) = \frac{tp}{[(1+10^{(p*(ga-x))})(1+10^{(q*(x-la))})]}$$

Usage

```
loggnls(ps, x)
```

loghill 25

Arguments

ps Vector of parameters: tp,ga,p,la,q,er

x Vector of concentrations (log10 units)

Value

Vector of model responses

Examples

loghill

Log Hill Model

Description

$$f(x) = \frac{tp}{(1 + 10^{(p*(ga - x))})}$$

Usage

Arguments

ps Vector of parameters: tp,ga,p,er

x Vector of concentrations (log10 units)

Value

Vector of model responses

```
loghill(c(1,2,3),1)
```

26 nestselect

mc3

Sample concentration-response data set from invitrodb

Description

A data set containing 100 chemicals worth of data for the Tox21 assay TOX21_ERa_BLA_Agonist_ratio, which measures response to estrogen receptor agonists. The data can be accessed further through the Comptox Chemicals Dashboard (https://comptox.epa.gov/dashboard).

Usage

mc3

Format

An object of class data. frame with 32175 rows and 7 columns.

Details

This data is extracted from the released version of the ToxCast database, invitrodb, at level 3 (mc3) and contains the concentration-response information.

A data frame with 32175 rows and 7 variables:

- dtxsid DSSTox generic substance ID
- casrn Chemical Abstracts Registry Number (CASRN)
- name chemical name
- spid sample ID there can be multiple samples per chemical
- logc log10(concentration), micromolar (uM)
- resp response in %
- assay name of the assay / assay component endpoint name

Source

doi:10.23645/epacomptox.6062623.v5

nestselect

Nest Select

Description

Chooses between nested models.

Usage

```
nestselect(aics, mod1, mod2, dfdiff, pval = 0.05)
```

poly1 27

Arguments

aics	Named vector of model aics (can include extra models).
mod1	Name of model 1, the model with fewer degrees of freedom.
mod2	Name of model 2, the model with more degrees of freedom.
dfdiff	Absolute difference in number of degrees of freedom (i.e. the difference in parameters). $ \\$
pval	P-value for nested model test.

Value

Named aic vector with losing model removed.

Examples

```
aics = c(-5,-6,-3)
names(aics) = c("poly1", "poly2", "hill")
nestselect(aics, "poly1", "poly2", 1)

aics = c(-5,-7,-3)
names(aics) = c("poly1", "poly2", "hill")
nestselect(aics, "poly1", "poly2", 1)
```

poly1

Polynomial 1 Model

Description

$$f(x) = a * x$$

Usage

```
poly1(ps, x)
```

Arguments

ps Vector of parameters: a,erx Vector of concentrations (regular units)

Value

Vector of model responses

```
poly1(1,1)
```

28 pow

poly2

Polynomial 2 Model

Description

$$f(x) = a * \left(\frac{x}{b} + \frac{x^2}{b^2}\right)$$

Usage

Arguments

ps Vector of parameters: a,b,er

x Vector of concentrations (regular units)

Value

Vector of model responses

Examples

pow

Power Model

Description

$$f(x) = a * x^p$$

Usage

Arguments

ps Vector of parameters: a,p,er

x Vector of concentrations (regular units)

Value

Vector of model responses

signatures 29

signatures

Sample concentration-response data set from HTTR

Description

A data set containing 6 of the active transcriptional signatures after perturbation of MCF7 cells with Clomiphene citrate (1:1).

Usage

signatures

Format

An object of class data. frame with 6 rows and 8 columns.

Details

A data frame with 6 rows and 8 variables:

- sample_id experimental sample ID
- dtxsid DSSTox generic substance ID
- name chemical name
- · signature transcriptional signature name
- cutoff the 95% confidence interval from the baseline response (2 lowest concentrations)
- onesd one standard deviation of the baseline response
- conc experimental concentrations, micromolar (uM)
- resp transcriptional signature response for each experimental concentrations, ssGSEA score

Source

doi:10.1093/toxsci/kfab009

References

Joshua A. Harrill, Logan J. Everett, Derik E. Haggard, Thomas Sheffield, Joseph L. Bundy, Clinton M. Willis, Russell S. Thomas, Imran Shah, Richard S. Judson, High-Throughput Transcriptomics Platform for Screening Environmental Chemicals, Toxicological Sciences, Volume 181, Issue 1, May 2021, Pages 68 - 89, https://doi.org/10.1093/toxsci/kfab009.

30 tcplfit2_core

tcplfit2_core	Concentration-response curve fitting	
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Description

Concentration response curve fitting using the methods from BMDExpress

Usage

Arguments

conc	Vector of concentrations (NOT in log units).
resp	Vector of responses.
cutoff	Desired cutoff. If no absolute responses > cutoff and force.fit = FALSE, will only fit constant model.
force.fit	If force.fit = TRUE, will fit all models regardless of cutoff.
bidirectional	If bidirectional = FALSE, will only give positive fits.
verbose	If verbose = TRUE, will print optimization details and aics.
do.plot	If do.plot = TRUE, will generate a plot comparing model curves.
fitmodels	Vector of model names to try fitting. Missing models still return a skeleton output filled with NAs.
	Other fitting parameters (deprecated).

Details

All models are equal to 0 at 0 concentration (zero background). To add more models in the future, write a fit_____ function, and add the model name to the fitmodels and modelnames vectors.

Value

List of N(models) elements, one for each of the models run (up to 10), followed by a last element "modelnames", which is a vector of model names so other functions can easily cycle through the output. For a full list, see the documentation for the individual fitting method functions. For each model there is a sublist with elements including:

· success - was the model successfully fit

tcplhit2_core 31

- aic the AIC value
- cov success of the the covariance matrix calculation
- rme root mean error of the data around the curve
- modl vector of model values at the given concentrations
- tp the top of the curve fit
- ga the AC50 or Hill paramters
- er the error term
- ... other paramters specific to the model (see the documentation for the specific models)
- tp_sd, ga_sd, p_sd, etc., the values of the standard deviations of the paramters for the models
- er_sd standard deviation of the error term
- pars the names of the parameters
- sds the names of the standard deviations of the paramters

Examples

```
conc <- c(.03, .1, .3, 1, 3, 10, 30, 100)
resp <- c(0, .1, 0, .2, .6, .9, 1.1, 1)
output <- tcplfit2_core(conc, resp, .8,
  fitmodels = c("cnst", "hill"), verbose = TRUE,
  do.plot = TRUE
)</pre>
```

tcplhit2_core

Hitcalling Function

Description

Core of hitcalling function. This method chooses the winning model from tcplfit2_core, extracts the top and ac50, computes the hitcall, and calculates bmd/bmdl/bmdu among other statistics. Nested model selection is used to choose between poly1/poly2, then the model with the lowest AIC (or AICc) is declared the winner. Continuous hitcalls requires tcplfit2_core to be run with force.fit = TRUE and "cnst" never to be chosen as the winner.

Usage

```
tcplhit2_core(
  params,
  conc,
  resp,
  cutoff,
  onesd,
  bmr_scale = 1.349,
  bmed = 0,
  conthits = TRUE,
  aicc = FALSE,
  identifiers = NULL,
  bmd_low_bnd = NULL,
  bmd_up_bnd = NULL)
```

32 tcplhit2_core

Arguments

params The output from tcplfit2_core

conc list of concentrations (not in log units)

resp list of corresponding responses

cutoff noise cutoff

onesd 1 standard deviation of the noise (for bmd calculation)

bmr_scale bmr scaling factor. Default = 1.349 bmed median of noise estimate. Default 0

conthits = TRUE uses continuous hitcalls, otherwise they're discrete. Default

TRUE

aicc aicc = TRUE uses corrected AIC to choose winning method; otherwise regular

AIC. Default FALSE

identifiers A one-row data frame containing identifiers of the concentration-response pro-

file, such as the chemical name or other identifiers, and any assay identifiers. The column names identify the type of value. This can be NULL. The values

will be included in the output summary data frame

bmd_low_bnd Multiplier for bmd lower bound. A value of .1 would require the bmd to be no

lower than 1/10th of the lowest concentration tested.

bmd_up_bnd Multiplier for the bmd upper bound. A value of 10 would require the bmd to be

no lower than 10 times the highest concentration tested.

Value

A list of with the detailed results from all of the different model fits. The elements of summary are:

- any elements of the identifiers input
- n_gt_cutoff number of data points above the cutoff
- · cutoff noise cutoff
- · fit method curve fit method
- top_over_cutoff top divided by cutoff
- rmse RMSE of the data points around the best model curve
- a fitting parameter methods: exp2, exp3, poly1, poly2, pow
- b fitting parameter methods: exp2, exp3, ploy2
- p fitting parameter methods: exp3, exp5, gnls, hill, pow
- q fitting parameter methods: gnls,
- tp top of the curve
- ga ac50 for the rising curve in a gnls model or the Hill model
- la ac50 for the falling curve in a gnls model
- er fitted error term for plotting error bars
- bmr benchmark response; level at which bmd is calculated = onesd*bmr_scale default bmr_scale is 1.349
- bmd benchmark dose, curve value at bmr
- bmdl lower limit on the bmd
- bmdu upper limit on the bmd

tcplObj 33

• caikwt - one factor used in calculating the continuous hitcall. It is calculated from the formula = exp(-aic(cnst)/2)/(exp(-aic(cnst)/2) + exp(-aic(fit_method)/2)) and measures how much lower the selected method AIC is than that for the constant model

- mll another factor used in calcualting the continuous hitcall = length(modpars) aic(fit_method)/2
- hitcall the final hitcall, a value ranging from 0 to 1
- top curve top
- ac50 curve value at 50% of top, curve value at cutoff
- lc50 curve value at 50% of top corresponding to the loss side of the gain-loss curve
- ac5 curve value at 5% of top
- ac10 curve value at 10% of top
- ac20 curve value at 20% of top
- · acc curve value at cutoff
- ac1sd curve value at 1 standard deviation
- conc conc string separated by I's
- resp response string separated by I's

tcpl0bj

Concentration Response Objective Function

Description

Log-likelihood to be maximized during CR fitting.

Usage

```
tcplObj(p, conc, resp, fname, errfun = "dt4", err = NULL)
```

Arguments

p	Vector of parameters, must be in order: a, tp, b, ga, p, la, q, er. Does not require names.
conc	Vector of concentrations in log10 units for loghill/loggnls, in regular units otherwise.
resp	Vector of corresponding responses.
fname	Name of model function.
errfun	Which error distribution to assume for each point. "dt4" is the original 4 degrees of freedom t-distribution. "dnorm" is the normal distribution.
err	An optional estimation of error for the given fit.

Details

This function is a generalized version of the log-likelihood estimation functions used in the ToxCast Pipeline (TCPL). Hill model uses fname "loghill" and gnls uses fname "loggnls". Other model functions have the same fname as their model name; i.e. exp2 uses "exp2", etc. errfun = "dnorm" may be better suited to gsva pathway scores than "dt4". Setting err could be used to fix error based on the null data noise distribution instead of fitting the error when maximizing log-likelihood.

34 toplikelihood

Value

Log-likelihood.

Examples

```
conc = c(.03,.1 , .3 , 1 , 3 , 10 , 30 , 100)
resp = c(0 , 0 , .1 ,.2 , .5 , 1 , 1.5 , 2 )
p = c(tp = 2, ga = 3, p = 4, er = .5)
tcplObj(p,conc,resp,"exp5")
lconc = log10(conc)
tcplObj(p,lconc,resp,"loghill")
```

toplikelihood

Top Likelihood

Description

Probability of top being above cutoff.

Usage

```
toplikelihood(fname, cutoff, conc, resp, ps, top, mll)
```

Arguments

fname	Model function name (equal to model name except hill which uses "hillfn")
cutoff	Desired cutoff.
conc	Vector of concentrations.
resp	Vector of responses.
ps	Vector of parameters, must be in order: a, tp, b, ga, p, la, q, er
top	Model top.
mll	Winning model maximum log-likelihood.

Details

Should only be called by hitcontinner. Uses profile likelihood, similar to bmdbounds. Here, the y-scale type parameter is substituted in such a way that the top equals the cutoff. Then the log-likelihood is compared to the maximum log-likelihood using chisq function to retrieve probability.

Value

Probability of top being above cutoff.

toplikelihood 35

```
fname = "hillfn"
conc = c(.03,.1,.3,1,3,10,30,100)
resp = c(0,.1,0,.2,.6,.9,1.1,1)
ps = c(1.033239, 2.453014, 1.592714, er = -3.295307)
top = 1.023239
mll = 12.71495
toplikelihood(fname, cutoff = .8, conc, resp, ps, top, mll)
toplikelihood(fname, cutoff = 1, conc, resp, ps, top, mll)
toplikelihood(fname, cutoff = 1.2, conc, resp, ps, top, mll)
```

Index

 ${\tt nestselect}, \textcolor{red}{26}$

* datasets mc3, 26 signatures, 29	poly1, 27 poly2, 28 pow, 28
acgnlsobj,2 acy,3	signatures, 29
bmdbounds, 4 bmdobj, 5	tcplfit2_core, 30 tcplhit2_core, 31 tcpl0bj, 33 toplikelihood, 34
cnst, 6 concRespCore, 6 concRespPlot, 8	, ,
exp2, 9 exp3, 9 exp4, 10 exp5, 11	
fitcnst, 11 fitexp2, 12 fitexp3, 13 fitexp4, 14 fitexp5, 14 fitgnls, 15 fithill, 16 fitpoly1, 17 fitpoly2, 18 fitpow, 19	
gnls,20 gnlsderivobj,20	
hillfn, 21 hitcont, 21 hitcontinner, 22 hitlogic, 23 hitloginner, 24	
loggnls, 24 loghill, 25	
mc3, 26	